Original Research

Distribution and Fractionation of Potentially Toxic Metals under Different Land-Use Patterns in Suburban Areas

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Abstract

To explore the geochemical effects of land-use patterns on potentially toxic metals in agricultural soils in tropical areas, soil samples were collected in Hainan Island, China. Total concentrations of chromium, copper, lead and zinc were analyzed by inductively coupled plasma mass spectrometry. The sequential extraction procedure was applied to determine the fractions of these potentially toxic metals. The results showed that the concentrations of potentially toxic metals differed among different land-use types. The concentrations of Cr, Pb and Zn revealed the following order: abandoned cropland > paddy field > vegetable land, while Cu was ordered as abandoned cropland > vegetable land > paddy field. Fractionation analysis showed that the carbonate-bound fraction was present in high proportion in abandoned croplands, the Fe-Mn oxide-bound fraction and the organic matter-bound fraction were present in high proportion in paddy fields, and the exchangeable fraction and residual fraction were present in high proportion in vegetable lands. The chemical fractionations of potentially toxic metals were correlated with pH and organic matter, resulting in the transformation of different fractions. The results of this study contribute to a better understanding of the effects of land-use patterns on potentially toxic metals in tropical areas.

Keywords: chemical fraction, heavy metals, land-use type, soil property, suburban soil

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**Introduction**

Soil is an important part of the ecological environment and the material basis for human survival and sustainable development. However, with sewage irrigation and the irrational use of pesticides and fertilizers, the problem of potentially toxic metal pollution in soil has become increasingly serious [1-2]. In addition, potentially toxic metals have limited self-purification capacity; they do not migrate readily and cannot be degraded by microorganisms [3]. Potentially toxic metals can be enriched in living organisms, e.g., they can enter the human body through water, plants and other media, endangering human health [4-5]. When potentially toxic metals accumulate to a certain extent in the soil, they will also lead to the reduction of soil productivity and crop yield, thus affecting the sustainable development of society and the economy [6]. Since soil properties such as soil pH, Eh and SOM (soil organic matter) content are highly dependent on land use and cropping systems, it is important to assess potentially toxic metal distribution and fractionation under contrasting land type (e.g., paddy land, abandoned land and vegetable land) [7]. The land type can significantly change the properties and conditions of soils and further affect the bioavailability of potentially toxic metals [8]. Through dissolution, adsorption, complexation, precipitation, condensation and other reactions, potentially toxic metals exist in various forms in soil, including the exchangeable fraction, carbonate-bound fraction, Fe-Mn oxide-bound fraction, organic matter-bound fraction and residual fraction [9-10]. Even if the total concentration is the same, the biological effects and environmental impacts are different due to the variation in fractionation. By exploring the available potentially toxic metals in soil, which are easily absorbed and utilized by organisms and then transferred through the food chain, the soil pollution status of potentially toxic metals and the harm caused to the environment can be better evaluated [4]. The pollution mechanism of potentially toxic metals in the soil system and organisms can be further understood.

In recent years, the distribution characteristics, influential factors, pollution assessment and remediation techniques of potentially toxic metals in soil under different land-use patterns have been reported by some researchers [11-13]. However, research on certain topics should be further studied, for example, the effect of land-use patterns on the fractionation distribution of potentially toxic metals on a typical tropical region; why the fraction concentrations differ under different land-use patterns; and the influence of the unique tropical climate of Hainan Island on fraction concentrations. For this, the following hypotheses were tested: 1) soil depths and land-use patterns (paddy field (PF), abandoned cropland (AC), vegetable land (VL)) influence the total concentrations of potentially toxic metals (Cr, Cu, Pb and Zn); 2) land-use types influence the chemical fractionation of potentially toxic metals; and 3) soil chemical properties (pH and SOM) influence the fractionation of potentially toxic metals. An attempt was made to identify the effects of land-use patterns on selected soil properties and potentially toxic metals in tropical areas.

**Materials and Methods**

**Study Area**

Dingan County (19°13′-19°44′N, 110°07′-110°31′E) is located in northeastern Hainan Island, China. The study area is classified as a tropical monsoon climate zone, with high annual precipitation and frequent typhoons [2]. It has an annual temperature of 24°C and an annual average precipitation of 1953 mm, with an uneven temporal distribution and most precipitation falling between May and October [14]. The maximum precipitation in a day is more than 200 mm. The study area is located in the Latosol zone of the monsoon forest in tropical temperate and humid areas. Paddy soil, lateritic soil and tidal sand soil are the main representative soil types.

**Sample Sources and Pretreatment**

Based on the understanding of the distribution of agricultural land in the study area, the soil of PF, AC and VL was selected as the research objects according to the principles of typicality and representativeness. The basic situation of each land-use pattern is as follows:

1) PF: Rice has been cultivated for more than 5 years; the planting method is based on crop rotation.

2) AC: Rice, peanuts and other crops were planted before the cropland was abandoned. Weeds and small trees grew on the surface after the cropland was abandoned for 3-5 years, with good coverage.

3) VL: Mainly planted cowpea, peanuts, *Pachyrhizus, Luffa* and other crops.

Five sampling sites with superior agricultural production conditions, complex land-use patterns and management measures were selected in the research area, as illustrated in Fig. 1. At each sampling site, adjacent PF, AC and VL were selected as sample plots, and there was a sample plot from PF, a sample plot from AC and a sample plot from VL. A total of 15 sample plots were selected. For each plot, three samples were collected from 0-10, 10-20 and 20-30 cm layer, respectively. A total of 45 samples were collected in the same time during soil sampling (5 sampling sites × 3 land-use patterns × 3 depths). A composite soil sample of 10 subsamples was collected within an area of about 50 m × 50 m. A total of 2 kg of fresh soil was collected from the mixed samples. At the same time, we recorded the cultivation and fertilization situation of the sample land areas in recent years through investigation.
The collected soil samples were stored in polyethylene bags for transport to the laboratory. Samples were placed in plastic trays. Impurities such as plant roots and gravel were removed. Then, clods were air-dried (20-25°C), crushed into small particles with a wooden hammer and sieved through a 0.15-mm nylon sieve. These samples were used for soil pH, SOM and elemental analyses.

Analytical Methods

The soil samples under different land-use patterns were analyzed after acid microwave digestion (ETHOS A, Milestone, Sorisole, BG, Italy). Soil samples (0.1 g dry weight) were digested in a pre-cleaned Teflon digestion tank with a solution of concentrated nitric acid (65% HNO₃; 6 mL) and hydrofluoric acid (65% HF; 3 mL). The samples were subjected to acid microwave digestion for 20 min after heating up to 200°C. After digestion and cooling, the digest solution was placed on a hot plate, heated at 120°C to near dryness with 0.5 mL of H₂O₂, and washed and dissolved with 0.02 mol of HNO₃ after cooling. Each sample was quantitatively transferred to a volumetric flask, which was then filled up to 50 mL with ultrapure water. Guaranteed reagent-grade chemicals and ultra-pure water were used for all procedures, unless stated otherwise. The concentrations of potentially toxic metals were measured using inductively coupled plasma-mass spectrometry (ICP-MS, Agilent 7900, Agilent Technologies, Santa Clara, CA, USA). An environmental mixed calibration standard of Cr, Cu, Pb and Zn (10 μg/mL, Agilent, Part#5183-4688) was diluted to produce a standard solution series in stages with 5% nitric acid (Merck, Germany). A mixed internal standard stock solution of ⁴⁰Li, ⁴⁰Sc, ⁷⁰Ge, ⁸⁰Y, ¹¹⁵In, ¹⁵⁹Tb, and ²⁰⁹Bi (10 μg/mL, Agilent, part#5183-4680) was diluted to 1 μg/mL with 5% nitric acid (Merck, Germany). Under optimum conditions, blank and standard solution series were measured, and the standard curves were automatically drawn by the instrument (r>0.9999). The main operating parameters of the instrument were as follows: RF power 1550 W, cooling gas 15.0 L/min, auxiliary gas 1.0 L/min, carrier gas 1.06 L/min, oxide formation (CeO/Ce<0.5%), doubly charged (Ce²⁺/Ce⁺<2%). The isotopes used were ⁵³Cr, ⁶⁶Zn, ⁶⁴Cu and ²⁰⁸Pb. The recoveries were between 85% and 125%, and the relative standard deviation was less than 5%. Precision and accuracy for potentially toxic metals analysis (Cr, Cu, Pb and Zn) are verified using standard reference materials from the National Research Center for Geo analysis of China [sediment, GBW07407].

Fig. 1. The location of the study area and distribution of sampling sites.
The potentially toxic metal concentrations of each sample and blank samples were analyzed. A sequential extraction scheme was applied to determine the fraction of potentially toxic metals in soil samples. The Cr, Cu, Pb and Zn concentrations of different fractions were determined by the sequential extraction procedure of Tessier et al. [15]. The chemical reagents, extraction conditions and corresponding fractions are listed in Table 1. Accepted recoveries range from 93% to 112%. Differences in potentially toxic metal concentrations between the determined and certified values are less than 10%.

Soil pH was determined in a 1:5 (soil: water ratio, w/v) suspension with a pH meter [16]. Soil organic carbon was first measured by wet oxidation, and then was calculated as SOM using the transfer factor 1.724 [17].

The significant differences of potentially toxic metal distributions were determined by one-way analysis of variance (ANOVA), followed by Duncan’s multiple comparisons [18]. Correlations between fraction concentrations of potentially toxic metals and selected soil properties were tested using Pearson’s correlation test. All statistical analyses in this study were performed with SPSS 19.0 and Origin 2017 software.

### Results and Discussion

#### Distribution Characteristics of Soil Properties and Potentially Toxic Metals in Soil under Different Land-Use Patterns

The properties of soil samples with three land-use patterns were determined. The results showed that the soil pH was acidic in PFs (5.15-6.06), ACs (5.83-7.22) and VLs (4.88-7.62). The average SOM contents in PF, AC and VL were 3.57, 1.57 and 1.02 g/kg, respectively. Soil physicochemical characteristics and microbial properties will change under different land-use patterns, which will further affect the migration and distribution of potentially toxic metals in soil.

Potentially toxic metal concentrations of soil samples from PFs, ACs and VLs were determined, and variance analysis and multiple comparisons were conducted. The results are shown in Fig. 2.

### Table 1. Sequential extraction procedure and the corresponding fractions.

| Step | Fraction          | Extraction procedure                                                                 |
|------|-------------------|--------------------------------------------------------------------------------------|
| 1    | Exchangeable      | 1 g of soil sample, 8 mL 1 mol/L MgCl₂, pH 7.0, shake 1 h, room temperature          |
| 2    | Carbonate-bound   | 8 mL 1 mol/L CH₃COONa, adjusted pH to 5.0 with CH₃COOH, shake 5 h, room temperature |
| 3    | Fe-Mn oxide-bound | 20 mL 0.04 mol/L NH₄OH·HCl in 25% CH₃COOH, pH 2.0, water bath, 96°C, 6 h, occasional shaking |
| 4    | Organic matter-bound | 3 mL 0.02 mol/L HNO₃, 30% H₂O₂ (adjusted to pH 2.0), water bath, 85°C, 5 h, 3.2 mol/L CH₃COONH₄ in 20% (v/v) HNO₃, shake 30 min |
| 5    | Residual          | 3 mL HNO₃+HClO₄+HF under high pressure, 170°C                                       |
as well [2]. In addition, the ACs are not cultivated or managed. Potentially toxic metals cannot be degraded by soil microorganisms. Therefore, potentially toxic metals are mostly concentrated in topsoil. The soil was relatively loose in the PFs and VLs. Continuous ploughing of PFs and leaching from VLs are conducive to the migration of potentially toxic metals to lower layers. In addition, crops also absorbed potentially toxic metals in the soil [20], resulting in potentially toxic metals in PFs and VLs being lower than those in ACs. This is consistent with the results of Inyang et al. [21], who reported the effect of fallow duration on the concentration of Cr. The concentrations of Cr, Pb and Zn in PFs were slightly higher than those in VLs, while the concentration of Cu in VLs was higher (Fig. 2). The main reason is that compound fertilizers are commonly used in VLs. The concentrations of Cu and Zn in compound fertilizers are relatively high [19], while the concentrations of Cr and Pb are relatively low. Zn shows more potential mobility and bioavailability than Cu [19] and is more readily migrates downward under the leaching action of rainwater, while Cu does not readily migrate. Therefore, the Cu concentration in VLs was higher.

The distribution of potentially toxic metals in the soil profile under different land-use patterns can reflect the trend of migration and transformation in the vertical direction, as shown in Fig. 3. In AC, the concentrations of Cr (42.26 mg/kg), Cu (42.30 mg/kg) and Zn (19.43 mg/kg) were the highest at a depth of 0-10 cm and decreased with increasing soil depth, while the concentration of Pb (19.99 mg/kg) was the highest at a depth of 10-20 cm. In PF, the concentrations of potentially toxic metals generally increased first and then decreased, mainly concentrated at a depth of 10-20 cm. In VL, the concentrations of potentially toxic metals were mainly concentrated in the surface layer (0-10 cm). The differences in the vertical distribution of the PF and VL may be related to their different farming methods. Before rice planting, PF soil needs to be turned over and basal fertilizer needs to be applied, which mainly affects the 10-20-cm soil layer. The application of chemical fertilizers and pesticides to VL depends on spraying, which mainly affects the soil surface.

Fractionation Distribution of Potentially Toxic Metals in Soil under Different Land-Use Patterns

The individual chemical fractions of Cr, Cu, Pb and Zn in the soil from the three land-use patterns were expressed as a percentage of the total metal concentration, as shown in Fig. 4. The proportions of the exchangeable fractions differed slightly. The exchangeable fraction is sensitive to the environment, is easily transferred, transformed and bioabsorbed, and has the strongest toxicity [22].

The proportion of carbonate-bound fractions under the three land-use patterns declined in the following order: AC>VL>PF. Driven by economic benefits,
farmers applied a large number of pesticides and fertilizers to increase crop yields, which led to the acidification of PF and the decrease of pH [23-24]. The soil microenvironment is relatively humid, and the base cations are easily leached. As a result, $H^+$ accumulates on the soil colloid, and the soil pH is low [25]. The carbonate-bound fraction is sensitive to soil pH and is easily released when the pH decreases [7]. The PF had the lowest soil pH (pH = 5.68) and the lowest proportion of the carbonate-bound fraction.

Metals in the Fe-Mn oxide-bound fraction were relatively high in PFs. This fraction is easily released when paddy soil is flooded and hypoxic. The Fe-Mn oxide-bound fraction can cause potential harm and is susceptible to anthropogenic activities.

The proportions of organic-bound fractions in the three land-use patterns were as follows: PF>AC>VL. Organic matter-bound fractions of potentially toxic metals exist in a variety of organics. Under aerobic conditions, the organics degrade and release the bound metals, and more SOM in the soil has a greater effect on the absorption of potentially toxic metals [26]. This result is consistent with the order of the SOM concentration: PF (3.57 g/kg)>AC (1.57 g/kg)>VL (1.02 g/kg). The SOM in PF was higher than that in AC. Anthropogenic activities such as the application of organic fertilizers supply SOM in PF, which is conducive to the accumulation of SOM [23, 27]. VL, like PF, is affected by fertilization, but there is no water as a protective layer to inhibit the decomposition of SOM. The decomposition and release of SOM are accelerated by high temperature, sufficient sunshine and strong biological effects in tropical areas [28]. Therefore, the decomposition of SOM content is higher than the supplementary content. The consistent of SOM and organic-bound fraction indicates that SOM can promote the formation of organic-bound fractions.

The residual fractions in VL and PF were higher than those in AC, which is consistent with the conclusion of Yang et al. [29]. They reported that the residual fraction of potentially toxic metals (e.g., Zn and Cu) increases after agricultural activities. This finding may suggest that active biological and cultivation behaviors promoted the mobility of metals, resulting in the relative enrichment of residual metals.

Correlation Analysis between the Fraction Concentrations of Potentially Toxic Metals and Selected Soil Properties

Effects of Soil pH on the Metal Fractions

The soil in the study area was slightly acidic, with a pH ranging from 4.88 to 7.62, with an average value of 6.10. The Pearson correlation analysis results between the fractions of potentially toxic metals and soil pH
are presented in Table 2. The soil pH showed negative correlations with the exchangeable fractions of Cu, Pb and Zn and positive correlations with the carbonate-bound fractions of Cr, Cu and Zn, among which there was a significant positive correlation with the carbonate-bound fraction of Zn (P < 0.05). This is because the soil pH will affect the concentration of soil carbonate. When the pH decreases, potentially toxic metals will be released due to the dissolution of carbonate. With an increase in soil pH, the concentration of carbonate increases, which readily complexes with potentially toxic metal ions to form carbonate-bound fractions [30], thus reducing the exchangeable fraction of potentially toxic metals in soil. Soil pH exhibited positive relationships with Fe-Mn oxide-bound fractions of Cr, Cu and Zn and potentially even Zn. The specific adsorption of Zn on the oxide surface is related to soil pH [31]. When the pH increases, most of the adsorbed potentially toxic metal ions become specifically adsorbed; therefore, the Fe-Mn oxide-bound fractions of Zn increase. There were positive correlations between soil pH and the organic matter-bound fractions of Cr, Cu and Zn and potentially even Zn. The residual fractions of Cu, Pb and Zn showed positive correlations with pH and potentially even Pb.

**Effects of SOM on the Metal Fractions**

The correlation between different fractions of potentially toxic metals in soil and SOM is shown in Table 3. The organic matter-bound fractions of the four potentially toxic metals showed positive relationships with SOM, among which Cr and Cu were significantly correlated at P<0.01 and Zn at P<0.05, indicating that an increase in SOM can increase the organic-bound states of potentially toxic metals. This may occur because increased SOM enhances the adsorption and complexation ability of potentially toxic metal ions. Therefore, the organic matter-bound fraction concentration of potentially toxic metals increases. In other words, an increase in the SOM concentration reduces the exchangeable sites and metal ions under acidic conditions, transforming them into organic matter-bound fractions [32]. There were both positive and negative correlations between the exchangeable fractions of potentially toxic metals and SOM, indicating that SOM has a dual effect on exchangeable fractions, i.e., it can reduce the concentration of the exchangeable fraction by adsorbing free potentially toxic metals and also plays the role of “ligand” and “migration carrier” by complexing and chelating with potentially toxic metals to improve the solubility of potentially toxic metals. Most of the carbonate-bound fractions exhibited negative relationships with SOM. However, the correlations were not significant. The Fe-Mn oxide-bound fractions of Cr, Cu and Zn showed positive correlations with SOM, among which Cr was significantly correlated at P < 0.05, indicating that an increase in SOM may enhance the complexation between potentially toxic metals and Fe-Mn oxides and increase the concentration of Fe-Mn oxide-bound potentially toxic metals. There was no significant correlation between the residual fractions of the four potentially toxic metals and SOM, which is consistent with the results reported by Honma et al. [33].

**Table 2. Correlation coefficient matrix between fractionation of potentially toxic metals and pH in surface soils.**

| Element | Exchangeable fraction | Carbonate-bound fraction | Fe-Mn oxide-bound fraction | Organic matter-bound fraction | Residual fraction |
|---------|-----------------------|--------------------------|---------------------------|-------------------------------|-------------------|
| Cr      | 0.190                 | 0.111                    | 0.409                     | 0.170                         | 0.014             |
| Cu      | -0.028                | 0.030                    | 0.082                     | 0.201                         | -0.149            |
| Pb      | -0.489                | -0.153                   | -0.395                    | -0.299                        | -0.475*           |
| Zn      | -0.052                | 0.737*                   | 0.555*                    | 0.136                         | -0.253            |

**, * Significant at P<0.01 and P<0.05, respectively.

**Table 3. Correlation coefficient matrix between fractionation of potentially toxic metals and SOM in surface soils.**

| Element | Exchangeable fraction | Carbonate-bound fraction | Fe-Mn oxide-bound fraction | Organic matter-bound fraction | Residual fraction |
|---------|-----------------------|--------------------------|---------------------------|-------------------------------|-------------------|
| Cr      | -0.118                | -0.064                   | 0.712*                    | 0.925*                        | 0.279             |
| Cu      | -0.306                | -0.363                   | 0.108                     | 0.900*                        | 0.380             |
| Pb      | -0.080                | 0.432                    | -0.080                    | 0.317                         | -0.078            |
| Zn      | 0.673**               | -0.293                   | 0.196                     | 0.540*                        | 0.107             |

**, * Significant at P<0.01 and P<0.05, respectively.
Conclusions

The concentrations of potentially toxic metals in the three land-use patterns were ranked as AC>PF>VL. Cr, Cu and Pb in VL were concentrated in the 0-10-cm soil layer, whereas Cr, Pb and Zn in PF were concentrated in the 10-20-cm soil layer, mainly due to different tillage and fertilization methods. These results showed that potentially toxic metals in soil are enriched to the tillage layer with the intensification of anthropogenic agricultural activities. Metals showed the highest proportion of the residual fraction in the three land-use types. The highest residual fraction and the relatively low labile fraction showed low mobility and bioavailability. However, the carbonate-bound fraction is sensitive to soil pH and is easily released when pH decreases, and the Fe-Mn oxide-bound fraction is easily released when soil is flooded and hypoxic. Because the pH of PF is low and crops are planted frequently in tropical areas, more attention should be paid to reducing the risk of carbonate-bound fraction and Fe-Mn oxide-bound fraction. In agricultural activities, more suitable cultivation and fertilization methods should be monitored with great concern, which can reduce the hazards of potentially toxic metals.

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Conflict of Interest

No potential conflict of interest was reported by the authors.

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