Changes in metal availability, desorption kinetics and speciation in contaminated soils during repeated phytoextraction with the Zn/Cd hyperaccumulator Sedum plumbizincicola

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ABSTRACT

Phytoextraction is one of the most promising technologies for the remediation of metal contaminated soils. Changes in soil metal availability during phytoremediation have direct effects on removal efficiency and can also illustrate the interactive mechanisms between hyperaccumulators and metal contaminated soils. In the present study the changes in metal availability, desorption kinetics and speciation in four metal-contaminated soils during repeated phytoextraction by the zinc/cadmium hyperaccumulator Sedum plumbizincicola (S. plumbizincicola) over three years were investigated by chemical extraction and the DGT-induced fluxes in soils (DIFS) model. The available metal fractions (i.e. metal in the soil solution extracted by CaCl₂ and by EDTA) decreased greatly by >84% after phytoextraction in acid soils and the decreases were dramatic at the initial stages of phytoextraction. However, the decreases in metal extractable by CaCl₂ and EDTA in calcareous soils were not significant or quite low. Large decreases in metal desorption rate constants evaluated by DIFS were found in calcareous soils. Sequential extraction indicated that the acid-soluble metal fraction was easily removed by S. plumbizincicola from acid soils but not from calcareous soils. Reducible and oxidisable metal fractions showed discernible decreases in acid and calcareous soils, indicating that S. plumbizincicola can mobilize non-labile metal for uptake but the residual metal cannot be removed. The results indicate that phytoextraction significantly decreases metal availability by reducing metal pool sizes and/or desorption rates and that S. plumbizincicola plays an important role in the mobilization of less active metal fractions during repeated phytoextraction.

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1. Introduction

Metal pollution of some soils has become a serious environmental concern and a potential threat to human health. The technique of phytoextraction using hyperaccumulators has been shown to be successful in remediating soils polluted with metals (McGrath et al., 2006; Japenga et al., 2007; Li et al., 2012). However, the technique requires long periods of remediation effort and successive or numerous crops are needed to remove adequate metal from contaminated soils to achieve safe levels (Koopmans et al., 2008; Li et al., 2014). During the process of phytoextraction soil metal availability will change as prolonging the remediation time increases interactions between soil and plant. Both short-term phytoextraction and rhizosphere experiments on hyper-accumulators show clear decreases in soil metal availability in contaminated soils after phytoextraction and in rhizosphere soils as compared to bulk soils (Keller and Hammer, 2004; Liu et al., 2011). However, after short-term phytoextraction the decreased available metal may be replenished through soil metal re-equilibration (Keller and Hammer, 2004). Investigation of metal availability changes over relatively long time periods may help to elucidate metal changes and to alleviate the environmental risk of metals due to phytoextraction. In addition, compared to short-term phytoextraction, the enhanced plant–soil interactions and more marked decrease in soil metals during long periods of phytoextraction can give more information on the mechanisms of phytoextraction. Moreover, the efficiency of remediation can decrease during repeated phytoextraction as shown by Li et al. (2014), and the further discrimination of changes in soil metals during phytoextraction might help to elucidate the factors involved.
Numerous methods have been used to investigate the behavior of metals in soils. Chemical extraction tests are among the most commonly and widely used methods for predicitcating or estimating the bioavailability of various forms of metals (D’Amore et al., 2005). Choice of chemical agent depends on the specific points of interest of the study. Concentrated acids are usually used as to determine total amounts of metals present and estimate the total potentially harmful metals. Soil solution metal or weaker agents extractable metal fractions are relatively soluble and bioavailable. The water-extractable or soil solution metals are commonly considered to be the most readily available fraction to biota (Seguin et al., 2004). Weak chemical agents, e.g. CaCl₂, Ca(NO₃)₂ and BaCl₂, extract fractions that have been used to denote metal availability to plants as shown in some studies (Peijnenburg et al., 2007; Meers et al., 2007a). Extraction with 0.01 M CaCl₂ has been suggested for estimating metal availability (Novozamsky et al., 1993; Houba et al., 2000) and gives a good indication of metal availability to plants (Meers et al., 2007b). The metal fraction extracted by EDTA is used to denote potentially available and mobile metal due to the strong metal complexing ability of EDTA (Anju and Banerjee, 2011). To denote potentially available and mobile metal due to the strong metal complexing ability of EDTA (Anju and Banerjee, 2011). EDTA) and the BCR test were used to investigate changes in metal availability and combined forms during long-term phytoextraction.

Soil metal mobility and availability depend largely on metal desorption from the soil solid phase and on metal concentration in the soil solution. If metal desorption from the solid phase is kinetically limited then this may be a factor limiting plant metal uptake (Zhang et al., 2006). The technique of diffusive gradients in thin films (DGT) is based on kinetic principles and can reflect the soil metal desorption process when metals in the soil solution are depleted at the interface between the roots and the soil (Davison et al., 2000). The kinetic information can be obtained by the model of DGT induced fluxes in soils (DIFS). The DIFS model has been used to study the metal desorption kinetics of contaminated soils (Ernstberger et al., 2005; Zhang et al., 2006) and changes in metal desorption kinetics induced by hyperaccumulators (Fitz et al., 2003). In the present study the changes in soil metal desorption kinetics during the repeated phytoextraction of four metal-contaminated soils with contrasting soil properties were investigated using the DIFS model.

The plant metal uptake and remediation efficiency of four metal-contaminated soils in long-term phytoextraction has been investigated in a previous study (Li et al., 2014). In the present study soil metal changes during the long-term phytoextraction were further investigated by single and sequential extractions and using the DIFS model. The aim was to elucidate changes in metal availability, associated forms and desorption kinetics in the different soils and to understand the possible soil metal processes induced by interactions between soil and plant during long-term phytoextraction.

2. Methods and materials

2.1. Soil characterization

Four types of metal contaminated soil were collected from the arable layer (top 15 cm) of agricultural fields in different parts of China. Two acid soils were collected from Dabaoshan in Guangdong Province, south China, and Huludao in Liaoning Province, north China, and are referred to as DBS and HLD, respectively. The soil from DBS is a Ferralic Anthrosols that has been polluted by metal mining activities and the soil from HLD is a Haplic Lithoxols whose main pollution source is a zinc factory that has operated for several decades. The other two soils are calcareous soils which were collected from Zhujiau and Shuanglingdong in Zhejiang Province, east China, and are labeled ZJW and SLD, respectively. They are from an area polluted by a copper smelter and the soil type is Hydragric Anthrosols. Each of the four contaminated soils was first air-dried and then passed through a 2-mm nylon sieve and mixed thoroughly. The metal contaminated soils were continuously phytoextracted using Sedum plumbizincicola for three years over seven successive crops in a glasshouse located in Nanjing. Soil samples were collected after the third harvest (P3) and the seventh harvest (P7) during long-term phytoextraction. Together with the soils without remediation (NP) there were a total of three types of soil samples with different cropping phytoextraction collected from the pot experiment during the experiment. After phytoextraction by seven crops the total Zn concentrations in the DBS, HLD, SLD and ZJW soils (P7) decreased by 37, 52, 12, and 19%, respectively, and Cd concentrations decreased by 64, 89, 37, and 38% compared to the NP soil (Table 1). All these soil samples were firstly air-dried, then passed through a 2-mm nylon sieve and were used to evaluate the soil metal changes induced by phytoextraction.

Portions of the sieved soils were used for determination of selected soil chemical properties. Soil pH, soil total organic carbon (TOC), soil cation exchange capacity (CEC), and soil total metal were analyzed by previously published methods (Li et al., 2014). Free iron/aluminum oxides (Fe₂O₃/Al₂O₃) were extracted with Na₂S₂O₄—Na₃C₆H₅O₇—NaHCO₃ and determined by atomic absorption spectrophotometry (Varian SpectraAA 220FS, Varian, Palo Alto, CA). The particle size distribution was determined with a Beckman LS230 laser diffraction apparatus. Selected chemicals properties of the contaminated soils with or without phytoextraction are shown in Table 1.

2.2. Calculation of DGT concentration and DIFS

Information on the standard piston DGT devices and the method of calculation of DGT concentration were described in a previous study (Li et al., 2014). In brief, the concentration of metal at the interface of the DGT device and the soil (C_DGT) was calculated as follows,

\[
C_{DGT} = C_E(V_{HNO_3} + V_{gel}) \Delta g / t AD_f e
\]

Where \( C_E \) is the concentration of metal in the 1 M HNO₃ elution solution (µg l⁻¹), \( V_{HNO_3} \) is the volume of HNO₃ added to the resin gel (1 ml), \( V_{gel} \) is the volume of the resin gel, typically 0.15 ml, \( t \) is the elution factor for each metal, typically 0.8, \( 1 \) is the deployment time (in sec), \( A \) is the exposure area, typically 2.54 cm², \( \Delta g \) is the thickness of the diffusive gel (0.08 cm) plus the thickness of the filter membrane (0.014 cm), and \( D \) is the diffusion coefficient of the metal in the gel.

Comparing the concentration of \( C_{DGT} \) to the measured soil solution (gained by centrifugation as described in Li et al. (2014)) concentration (\( C_{soil} \)) provides a ratio, \( R \), which is dimensionless and gives an indication of the extent of the depletion of the soil solution concentration at the interface with the DGT:

\[
R = C_{DGT}/C_{soil}
\]
2.3.2. Soil metal extraction

Soil samples NP, P3 and P7 were used for metal extraction with CaCl₂ and EDTA. (1) Air-dried soil (2 g) was extracted with 0.01 M CaCl₂ (soil: extractant ratio 1:10) by shaking at 19 °C for 2 h, then centrifuging (10 min at 3000 rpm) and filtering. Cadmium and Zn concentrations in the supernatant were measured by ICP-MS (Varian Ultramass) within various calibration ranges using Rh as internal standard. Air-dried soil (2 g) was extracted with 0.05 EDTA (soil: extractant ratio 1:10) by shaking at 25 °C for 1 h, then centrifuging (3 min at 3000 rpm) and filtering. Extractable Cd and Zn in the filtrate were then determined by atomic absorption spectrophotometry (Varian SpectrAA 220FS; Varian, Palo Alto, CA).

Air-dried samples of soils NP, P3 and P7 (2.00 g) were used for extraction using the modified BCR (Community Bureau of Reference) sequential extraction procedure following Luo and Christie (1998). The certified reference material BCR701 was also extracted along with the soil samples for quality control. All sample recoveries (sum of four fractions/total in soil × 100) were between 85 and 115%.

2.4. Statistical analysis

Statistical analysis was performed by one-way analysis of variance (ANOVA) with Duncan’s multiple range test at the 5% level to compare the mean values using the SPSS version 16.0 for Windows software package. Data are presented as mean ± standard error of the mean (SEM, n = 3).

3. Results and discussion

3.1. Changes in soil parameters after repeated phytoextraction

The changes in selected parameters during repeated phytoextraction were determined (Table 1). Compared to the soils without phytoextraction (NP), in the soils with seven crops of S. plumbizincicola there was an apparent decrease in soil organic carbon. This may result from activities during the repeated phytoextraction, e.g. continuous cultivation combined with plant shoot removal and chemical fertilizer application causing the rapid mineralization of soil organic carbon. Soil pH changes were different for soils after phytoextraction. Increases in soil pH were observed in soil ZJW but no significant changes were found in DBS, HLD or SLD after repeated phytoextraction. This suggests that there was no unambiguous trend of changing soil pH induced by the process of repeated phytoextraction in this study. The content of calcium carbonate in calcareous soils SLD and ZJW showed an apparent decline (by 9.68 and 27.0%, respectively) after phytoextraction. This might result from plant root and soil microbial activities involving the release of H⁺. In addition, the amounts of free iron oxides (Fe₄Al₃) in all soils increased with the sole exception of...
Table 2
Zinc and Cd concentrations in the soil solution and extracted by CaCl₂ and EDTA in the contaminated soils without (NP) and with three (P3) or seven (P7) croppings of phytoextraction by the hyperaccumulator Sedum plumbezincicola; DBS, HLD, SLD, ZJW are four metal contaminated soils. Values are mean ± standard error of the mean (n = 3).

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Soil</th>
<th>Zn (mg L⁻¹/mg kg⁻¹)</th>
<th>Cd (µg L⁻¹/mg kg⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>NP</td>
<td>P3</td>
</tr>
<tr>
<td>Soil solution</td>
<td>DBS</td>
<td>21.4 ± 0.3</td>
<td>3.33 ± 0.4</td>
</tr>
<tr>
<td></td>
<td>HLD</td>
<td>15.9 ± 0.5</td>
<td>0.45 ± 0.02</td>
</tr>
<tr>
<td></td>
<td>SLD</td>
<td>0.09 ± 0.01</td>
<td>0.03 ± 0.01</td>
</tr>
<tr>
<td></td>
<td>ZJW</td>
<td>0.13 ± 0.01</td>
<td>0.03 ± 0.00</td>
</tr>
<tr>
<td>CaCl₂*</td>
<td>DBS</td>
<td>29.8 ± 0.5</td>
<td>7.63 ± 0.43</td>
</tr>
<tr>
<td></td>
<td>HLD</td>
<td>62.7 ± 0.4</td>
<td>12.6 ± 0.8</td>
</tr>
<tr>
<td></td>
<td>SLD</td>
<td>0.12 ± 0.01</td>
<td>0.14 ± 0.01</td>
</tr>
<tr>
<td></td>
<td>ZJW</td>
<td>0.28 ± 0.00</td>
<td>0.20 ± 0.07</td>
</tr>
<tr>
<td>EDTA</td>
<td>DBS</td>
<td>26.8 ± 0.2</td>
<td>5.06 ± 0.62</td>
</tr>
<tr>
<td></td>
<td>HLD</td>
<td>60.4 ± 0.8</td>
<td>15.6 ± 0.52</td>
</tr>
<tr>
<td></td>
<td>SLD</td>
<td>40.8 ± 0.52</td>
<td>40.2 ± 0.8</td>
</tr>
<tr>
<td></td>
<td>ZJW</td>
<td>44.6 ± 7.9</td>
<td>32.0 ± 1.6</td>
</tr>
</tbody>
</table>

* The CaCl₂-extracted metal data are cited from Li et al. (2014).

DBS, and this was more pronounced in the calcareous soils. For example, after phytoextraction the increases in Fed in SLD and ZJW were 41.6 and 53.4%, respectively, compared to the soils without phytoextraction. Plant rhizosphere activities might be responsible for the changes. The release of H⁺ and organic materials by roots may dissolve and complex iron and aluminum (Fujii, 2014), and in the partially anaerobic conditions of the rhizosphere some microbial activities might release iron or manganese by using them as terminal electron acceptors (Hoefer et al., 2015). The above chemical or biochemical activities might be beneficial to the transformation of other iron forms to Fe₃⁺.

3.2. Changes in metal availability during repeated phytoextraction

Metals in the soil solution and extracted by CaCl₂ and EDTA can be used as an indicator of the directly available fractions, more readily bioavailability, and potential availability for biota, respectively (Novozamsky et al., 1993; Houba et al., 2000; Seguin et al., 2004; Meers et al., 2007b; Anju and Banerjee, 2011). These three metal fractions were used here to discriminate changes in metal availability during repeated phytoextraction and the data are shown in Table 2. After phytoextraction with seven crops, soil solution Zn and Cd decreased markedly (by > 74%) and the highest decrease (>99%) was found in the acid soil HLD. Although soil solution metal concentrations decreased with increasing remediation cropping, sharp decreases were found at the first three croppings and the differences between soils with 3 and 7 phytoextraction croppings were small (Table 2). This indicates the slow release of the remaining soil metal at the later stages of repeated phytoextraction which might be related to the decrease in available metal pool sizes and increases in binding forces to the soil solid phase. Compared to DBS soil with lower total Cd and Zn, the severely Cd-contaminated soil HLD had lower soil solution Cd and Zn with repeated phytoextraction (Table 2). This may result from the low soil pH of DBS (3.71 for NP, and 3.87 for P, Table 1) which facilitates the distribution of metals into the soil solution. Similarly, because of high soil pH (over 7, Table 1), soil metals might precipitate as hydroxides or combined with carbonates (Du Laing et al., 2009), which might be responsible for the several magnitudes lower soil solution metal found in the calcareous soils than in the acid soils (Table 2).

When soil solution metal is considered as the fraction directly available to the plant, the amount of metal extracted by CaCl₂ and EDTA reflects the size of two pools with different bioavailability. The acid soils without phytoextraction had higher available metal pool sizes (extracted by CaCl₂) than the calcareous soils but the metal concentrations in potentially available pools (determined by extraction with EDTA) were of the same magnitude (Table 2). During repeated phytoextraction the changes in extractable metals by CaCl₂ and EDTA depended on the soils. The extractable Zn and Cd by CaCl₂ and EDTA decreased greatly (by 84–96%) after seven croppings of phytoextraction in the acid soils and the trends of decline were similar to metals in the soil solution, showing dramatic changes at the initial stages of phytoextraction. This indicates that the metal declines in the soil solution are controlled mainly by the decreases in the available metal pool sizes in acid soils during phytoextraction. Previous studies show that hyperaccumulators can significantly decrease soil metal availability (Keller and Hammer, 2004; Dessureault-Rompre et al., 2010; Liu et al., 2011), but if just short-term phytoextraction is conducted the decreased available metal can be offset as the soil metals rebalance when phytoextraction ceases (Keller and Hammer, 2004). In the long-term repeated phytoextraction of the present study, metal extracted by CaCl₂ and EDTA showed clear decreasing trends during phytoextraction of the acid soils. These results indicate that phytoextraction shows outstanding effects in reducing metal environmental risk. However, in the calcareous soils there was no significant change in either CaCl₂ or EDTA-extractable Zn during phytoextraction. This could be related to the supplementation of Zn from less available pools because of the large buffering capacity for Zn in the calcareous soils. Although the available metal pool sizes did not decline the soil Zn environmental risk was to some extent alleviated. This is due to the soil Zn desorption rates showing large decreases after phytoextraction as discussed below in the next section. Moreover, in long-term phytoextraction the available metal pool size must be decreased with further increase in phytoextraction time and this is confirmed by the absence of significant changes in CaCl₂-extractable Cd after three croppings of S. plumbezincicola but a clear decrease after seven croppings (Table 2). The above results indicate that the target of phytoextraction in acid soils can be decreases in soil metal availability, but in calcareous soils should be based on the decline in the total metal concentration when there are no clear decreases in available fractions because of their high metal replenishment capacity.

3.3. Metal desorption kinetics based on the DIFS model

Knowledge of metal mobility and availability in soils depends on soil solution metals and available metal pool sizes in the solid phase, and metal desorption connects these two aspects. The dynamic DIFS model of the soil-DGT system is used here to obtain information on the changes in metal desorption kinetics during
The metal distribution coefficient between solid and solution, $K_d$, along with the measured $R$ value and other soil parameters (Table 3) were used to calculate the response time ($T_c$), which is the characteristic time taken by the system to approach equilibrium (Degryse et al., 2009) (Fig. 1). Soil metal desorption rate constants ($k/C_0$) were also calculated based on Equation (4) (Fig. 1).

In the soils without phytoextraction, $T_c$ values were much higher in the acid soils (DBS and HLD) than the calcareous (SLD and ZJW), indicating the slow metal supply from the soil solid phase to solution in acid soils. The observed longer response time for acid soils is reasonable. Acid soils have relatively high soil solution metal concentrations but small pool sizes of metals in the solid phase, and this leads to large disturbance induced by removal of metals to the DGT. The long response time in acid soils and short response times in calcareous soils are consistent with previous studies (Zhang et al., 2006; Ernstberger et al., 2005). After repeated phytoextraction the response time of Zn and Cd in all soils changed markedly and showed large decreases in the acid soils but there were clear increases in the calcareous soils (Fig. 1). By definition $T_c$ is the characteristic time taken to approach equilibrium between the soil solid and solution phases and should depend on the labile metal distribution coefficient between solid and solution ($K_d$) and the metal desorption rate constant ($k/C_0$) (Degryse et al., 2009). Theoretically, the metal desorption rate cannot be enhanced after phytoextraction because more easily desorbed metal was removed first by the hyperaccumulator as validated by the decreased desorption rate constants ($k/C_0$) (Fig. 1). Thus, the decreases in

<table>
<thead>
<tr>
<th>Soil</th>
<th>$P_c$</th>
<th>$\Phi$</th>
<th>$R^1$</th>
<th>$K_d$</th>
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<tbody>
<tr>
<td></td>
<td>Zn NP</td>
<td>P3 P7</td>
<td>Cd NP</td>
<td>P3 P7</td>
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<tr>
<td>DBS</td>
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<td></td>
<td></td>
<td>1200</td>
<td>1590</td>
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* Denotes that $R$ value data are cited from Li et al. (2014); $C_{DGT}$ is the concentration of metal measured by DGT; $C_{soln}$ is the concentration of metal measured in the soil solution; $K_d$ is the ratio of total metal concentration to metal concentration in the soil solution; NP, soil without phytoextraction; P3 and P7, soils were phytoextracted over three and seven crops of the hyperaccumulator *Sedum plumbizincicola*, respectively; DBS, HLD, SLD, ZJW are four metal-contaminated soils.
response time in acid soils after phytoextraction should be related to large increases in $K_d$ as a result of the decreases of $>90\%$ in soil solution metal (Table 3). Although in calcareous soils $K_d$ also showed marked increases, the response time still showed an increasing trend during phytoextraction. This result indicates that the effect of the much decreased $k_{-1}$ on the response time was more profound than the effect of the increased $K_d$ in calcareous soils as illustrated by the sharp decreases in $k_{-1}$ between soil without phytoextraction (NP) and with three crops of the hyperaccumulator (P3).

In DBS soil the $k_{-1}$ of Cd during repeated phytoextraction remained constant indicating that the same mechanism controls
metal release from the soil solid phase to the soil solution (Fig. 1). This is supported by the acid soluble Cd being the main available fraction (Fig. 3). With the exception of Cd in DBS soil, the desorption rate constants \((k_{-1})\) of Cd and Zn in all soils decreased greatly after repeated phytoextraction (Fig. 1). This indicates that there were different available metal pools controlling metal desorption during phytoextraction, and the remaining soil metal after phytoextraction had stronger binding forces to soil particles. Especially in calcareous soils, desorption rate constants showed a much more dramatic decline than in acid soils during the initial stages of phytoextraction (Fig. 1). As the target of phytoextraction is based on total metal removal, the combined decreases in soil solution metal concentrations and desorption ability indicate that phytoextraction of calcareous soils can greatly decrease soil metal mobility and the risk of metal transfer to the environment.

3.4. Metal chemical fractions based on BCR sequential extraction

The different extraction steps of BCR chemical extraction represent different forms of association and different mechanisms of availability (Ure et al., 1993; Anju and Banerjee, 2011). In the present study the changes in Zn and Cd chemical fractions in the four contaminated soils during repeated phytoextraction were investigated and the data are shown in Figs. 2 and 3.

In the soils without phytoextraction (NP), Zn distribution in the acid soils DBS and HLD followed the orders residual > acid soluble and oxidisable > reducible fraction and residual > oxidisable and reducible > acid soluble fractions, respectively (Fig. 2). Large residual fractions of Zn indicate that most of the metal is background Zn in soil crystal lattices, a form not easily taken up by plants. This is also indicated by the absence of significant changes in residual fractions of metals after three years of phytoextraction (Figs. 2 and 3). Soil Cd in DBS was distributed mainly in the acid soluble fraction (76.3% of total Cd), suggesting that Cd existed in highly available forms in the soil. This may result from the low soil pH (Table 1) which promotes the release of Cd from the soil solid phase. In HLD soil Cd was mainly partitioned in the oxidisable and acid soluble fractions (Fig. 1). Large amounts of oxidisable Cd in HLD soil may be associated with high pollution with elemental sulfur because of the activities of a zinc factory in this area (Zhu et al., 2001) promoting the binding of Cd mostly to the sulfide phase. During repeated phytoextraction, acid soluble Zn and Cd may have been easily taken up by S. plumbizincicola and were almost removed after three croppings in the acid soils (Figs. 2 and 3). Reducible and oxidisable metals also showed clear decreases during the first three croppings, indicating that these two metal fractions can also be mobilized and subsequently be taken up by S. plumbizincicola. In HLD soil, although soil Cd still remained in the highly available fractions (acid soluble + reducible + oxidisable, 86.4% of total) after the first three croppings of phytoextraction, the Cd removal efficiency was much lower during the subsequent four croppings. This suggests that if no other additional measures are taken to enhance metal phytoavailability, the later stages of phytoextraction will be time consuming and ineffective. Given that the acid soluble metal was the dominating fraction in the acid soils after phytoextraction when excluding the non-available residual fraction, measures to accelerate soil organic matter transformation and changing soil redox conditions might enhance the efficiency of remediation.

Soil Zn and Cd in the calcareous soils showed high potential availability based on BCR extraction. Soil Zn was partitioned mainly in the first three fractions (acid soluble + reducible and oxidisable) which accounted for 86.5 and 92.3% of the total metal in SLD and ZJW soils, respectively (Fig. 2), and soil Cd was distributed mainly in the acid soluble and reducible fractions which accounted for 91.8 and 80.6%, respectively, of the total (Fig. 3). However, the Zn and Cd decreases in the calcareous soils after phytoextraction were less pronounced than those in the acid soils. Only slight decreases in acid soluble and oxidisable Zn were found in both calcareous soils (Fig. 2). Although there were visible declines in acid soluble, reducible and oxidisable Cd, the removal rates were quite low and showed further decline in the low Cd polluted calcareous soil (ZJW) in the later stages of phytoextraction (Fig. 3). Moreover, in contrast to the acid soils, after phytoextraction both calcareous soils had high remaining amounts of acid soluble Zn and Cd as well as reducible and oxidisable fractions (Figs. 2 and 3). This may have resulted from metal precipitation or strong adsorption/complexation of metals to carbonates in the high pH calcareous soils (Du Laing et al., 2009). These results suggest that in calcareous soils acid soluble metal is not easily removed by plants (in contrast to the acid soils) and lowering soil pH might be a necessary measure for increasing remediation efficiency.

In the present study there was no significant change in the residual metals in any of the soils (Figs. 2 and 3). This suggests that the above metal processes during phytoextraction had no effects on residual metals, the fraction unavailable to plants and not to be considered as a target of phytoextraction. This is important for phytoextraction of soils contaminated with large amounts of residual metals. If the residual metals can be ignored it is possible to increase the efficiency and shorten the time required for phytoextraction. For example, after seven croppings of phytoextraction, the soil Zn remaining was mostly distributed in the residual fraction, and the percentages of residual Zn to total were 72.3 and 61.7% for DBS and HLD soils, respectively. The remaining available Zn fractions (acid soluble + reducible + oxidisable) were very low (50.5 and 93.3 mg kg\(^{-1}\)) and were already below the permissible level for agriculture soils in China (200 mg kg\(^{-1}\)). Given that the residual fraction of Zn is unavailable to plants it is not necessary to achieve a target based on total metal phytoextraction. However further biological tests under greenhouse or field conditions are still required to ensure safe crop production.

3.5. Soil metal processes during phytoextraction

The lower metal removal in calcareous soils than in acid soils may be related to the two main soil metal processes during repeated phytoremediation. First, metal releases from the solid phase to the soil solution are based on an equilibrium principle when soil solution metal depletion is induced by plant metal uptake. In this process soil metals with a rapid desorption rate and high concentration in the soil solution might be propitious to phytoextraction. For example, the acid soil HLD showed high amounts of metal removal (Table 1). In the acid soil DBS, although the soil solution metal concentration was as high as that of HLD during repeated phytoextraction, its metal desorption ability was weak as evidenced by large response time \((T_r)\) and low desorption rate constants \((k_{-1})\) (Fig. 1), resulting in lower metal removal by plants from DBS than HLD. In the calcareous soils SLD and ZJW, although metals in NP soils showed rapid supply to solution as estimated by a small \(T_r\) and high \(k_{-1}\) (Fig. 1), the plant metal concentrations in calcareous soils were still several times lower than those in the acid soils (Li et al., 2014). This resulted mainly from soil solution metals being one to two orders of magnitude lower in calcareous soils (Table 2). This may be one of the explanations for the low metal removal efficiency by plants in the calcareous soils. Moreover, the sharp decreases in soil solution metals (Table 2) and visible decline in metal desorption ability (indicated by \(k_{-1}\)) in the acid soils (Fig. 1) might also be the main factor responsible for the lower metal removal at the later stages of repeated phytoextraction compared to the early stages (Li et al., 2014). Interestingly, although
the soil solution metal concentrations, metal availability and metal desorption ability showed large decreases (Table 2 and Fig. 1) during repeated phytoextraction, plant metal uptake in calcareous soil remained constant as in Li et al. (2014). This indicates that in calcareous soils metal releases based on equilibrium principles cannot explain plant metal uptake. As previous studies using isotope dilution techniques indicate, acid-soluble and part of oxidisable metals in soils can be released to the soil solution based on equilibrium principles (Ahnstrom and Parker, 2001; Marzouk et al., 2013). However, in this study soil reducible and oxidisable metal fractions in both acid and calcareous soils showed large decreases after phytoextraction (Figs. 2 and 3). These decreases also cannot be explained by the above metal process based on the equilibrium principle. Furthermore, rhizosphere activities can change plant metal accessibility and availability (Chardot-Jacques et al., 2013). The sequestering of hyperaccumulator roots can enhance metal availability by decreasing soil pH or forming soluble DOM-metal complexes (Desureau-Rompre et al., 2010; Li et al., 2013a,b). In addition, the associated microorganisms and their activities in hyperaccumulator rhizosphere soil also play a role in enhancing metal availability (Li et al., 2010; Wei and Twardowska, 2013). In addition to the microbial activities, soil bound metals can be released with the transformation of Fe/Al/Mn oxides when there is a shift in soil redox conditions (Husson, 2013) or decomposition of soil organic matter. The rhizosphere effect on metal uptake has been considered to be one of the most important mechanisms in phytoextraction using hyperaccumulators and might be responsible for the removal of reducible and oxidisable metals found after repeated phytoextraction. However, the actual mechanism and how it may be enhanced to increase remediation efficiency are still poorly understood.

4. Conclusions

Repeated phytoextraction can significantly decrease soil solution metal concentrations in both acid and calcareous soils; the removal of available metals (extracted by CaCl₂ and EDTA) depends on the soils, with high removal efficiency in acid soils but no significant change or only slight changes in calcareous soils during phytoextraction. Metal desorption rates in all soils (with the sole exception of the low pH soil DBS) showed marked decreases, especially in the calcareous soils which showed a very large decline in desorption rate constant at the initial stages of phytoextraction. These results indicate that phytoextraction can reduce metal availability by decreasing the soil available metal pool size and/or metal desorption rates.

In the present study acid soluble metals were easily removed by S. plumbizincicola from acid soils during the initial stages of phytoextraction, but there was little removal from the calcareous soils. Repeated phytoextraction can remove reducible and oxidisable metals from both acid and calcareous soils but the residual metals cannot be removed. This suggests that S. plumbizincicola can play an important role in the mobilization of the non-labile fraction of metals during repeated phytoextraction.

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