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Immobilization of heavy metals in sewage sludge by using subcritical water technology

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Abstract

Heavy metals (HMs) immobilization in sewage sludge was investigated by using subcritical water technology (SCWT) in this study. The characteristics of sludge and toxicity of HMs were analyzed after SCWT process. The results showed that besides large reduction in sludge volume, SCWT had some positive effect on HMs dissolution into liquid phase, while the majority of HMs was still accumulated in solid phase. The direct toxicity and bioavailability of HMs in sludge was greatly decreased with no toxicity fractions of HMs highly increased. Pb was always at low risk level and the risk of other HMs was greatly reduced from low risk to no risk after SCWT treatment. Moreover, the leaching toxicity of HMs declined after SCWT and the best result was obtained at 280°C with the metal concentrations in leachate decreased by 97.46%, 93.91%, 86.14%, 73.67%, 71.93% and 10.71% for Cu, Cd, Zn, Cr, Ni and Pb, respectively.

Keywords: Sewage sludge; subcritical water technology; heavy metals; immobilization
1. Introduction

Land use is one of the most economic ways for sewage sludge disposal because the sludge can provide many easily available nutrient sources like N, P, K and organic matters. However, accumulated heavy metals (HMs) and their bioavailability are often the limiting factors (Fuentes et al., 2004; Wang et al., 2005). In order to reduce the toxicity of HMs to the environment, two main approaches can be applied, i.e. HMs removal from the sludge or HMs immobilization in the sludge.

After being removed from sludge the total concentrations of HMs can be reduced, resulting in the decrease of potential environmental contamination. This approach includes various extraction methods such as chemical extraction by organic or inorganic acid (Deng et al., 2009), ionic liquids extraction (Fuerhacker et al., 2012), and solid-liquid-solid extraction (Sprynskyy, 2009), bioleaching (Liu et al., 2012) and electrokinetic remediation (Hanay et al., 2009), etc. However, these methods are not practical and cost-effective due to their requirement of long contact time and difficulties in removal efficiency control.

HMs immobilization has been widely utilized in soil remediation because of its simplicity and cost-effective management. Some commonly used materials for HMs immobilization are biochar, phosphate, carbonate, silicate and phosphate-, carbonate-, silicate-based minerals (McGowen et al., 2001; Mendez et al., 2012). The immobilization mechanisms are different for specific heavy metals, including precipitation with complex anions, surface adsorption, ion exchange, etc (Basta et al., 2001; Theodoratos et al., 2002). After this process the total HMs concentration in sludge cannot be reduced and its potential environmental contamination sometimes increases. However, the direct toxicity or leachable fractions of some HMs may be reduced, which is much more important for the decrease of environmental risk. For the immobilization of HMs, such methods as forming ceramsite by incineration (Xu et al., 2009), mixing with biochar produced from agricultural residues (Uchimiya et al., 2011; Debela et al., 2012), natural mineral adsorption (Kosobucki et al., 2008), and thermal treatment (Obrador et al., 2001) have been tried and tested.
Thermal treatment, especially sub- and super-critical water technology, is regarded as environmentally friendly and has been used for sewage sludge treatment. It was reported that the toxicity and leaching characteristics of HMs could be greatly changed after thermal treatment at 300-400°C (Li et al., 2012; Yuan et al., 2011). Up to now, the related research works were conducted at temperatures higher than 300°C resulting in higher energy consumption and lower organic matter content left in the sludge. In the present study, subcritical water technology (SCWT) was applied under lower temperatures, and the objective of this study was to: (1) characterize the variation of sludge after SCWT process; (2) analyze the HMs (Cr, Ni, Cu, Zn, Cd and Pb) speciation and fractional transformation; (3) evaluate the toxicity and bioavailability of HMs in sludge; and (4) examine the leachability and leaching toxicity of HMs after SCWT.

2. Materials and methods

2.1. Sewage sludge

Dewatered anaerobically digested sewage sludge was used in this study and obtained from a wastewater treatment center in Ibaraki prefecture, Japan. The sludge was kept in 4°C after sampling. Before SCWT treatment, the sludge was diluted by adding deionized water, and mixed with an appropriate volume of HMs mixture including Cr (VI), Ni (II), Cu (II), Zn (II), Cd (II) and Pb (II) because of very low levels of HMs contained in the original sewage sludge according to the designed HMs concentrations. The HMs mixture was prepared by using the standard HMs solutions (1000 mg/L in 0.1 M HNO₃, respectively), and then adjusted its pH value to 7.0 with 2 M NaOH solution. The sludge was homogeneously agitated for 2 h and then kept in a glass beaker (enclosed by plastic wrap) in refrigerator at 4°C. During the following week the sludge was stirred manually for 5 min once every day in order to let the supplemented HMs completely dispersed or distributed into the sludge. Table 1 lists the characteristics of the sludge used in this study.

2.2. SCWT procedure

The SCWT experiments were conducted at 170°C (SCWT170), 200°C (SCWT200) and 280°C (SCWT280), respectively for 1.0 h in an enclosed stainless steel reactor with a volume of 200ml. In order to keep the same condition, the added sludge volume was 100ml. After SCWT process, the sludge sample was cooled to ambient temperature, and
then the liquid phase was collected after separation by vacuum filtration at constant pressure (0.02 MPa) for 10 min and labeled as liquid sample (LS) 170, LS200 and LS280, respectively. The solid residue (SR) was dried at 60 °C for 48 h, and ground into particle size smaller than 0.15 mm which was sufficiently fine to be homogeneously mixed, and then stored in enclosed plastic bags as samples of SR170, SR200 and SR280, respectively.

The solid residue (CSR) samples from control experiment (CK, without subcritical treatment) were obtained by directly using the sewage sludge prepared in section 2.1.

2.3. Fractionation procedure of HMs

Speciation and extraction of HMs in SR was conducted by using the sequential extraction procedure proposed by Tessier et al. (1979) with 1.0 g sample. The five fractions were exchangeable metals (F1), metals bound to carbonate (F2), metals bound to iron and manganese oxides (F3), metals bound to organic and sulfide (F4) and metals in residue (F5). For F1 to F4, after every step, the supernatant was collected by centrifuging at 10,000 rpm for 10 min, and diluted to a fixed volume with 2% HNO3. The supernatant samples were filtered to remove fine particles and digested with H2O2 and concentrated HNO3 to remove dissolved organics. The HMs in F5 and the total concentration of HMs in SR was extracted by digestion with acid mixture (VHCl/VHNO3/VHF = 3/1/1) (Bettinelli et al., 2000) in PTFE beakers and heated on a hot plate. All the samples were kept in 4 °C before analysis.

2.4. Treatment efficiency assessment

2.4.1. Overall performance of SCWT on sewage sludge

After SCWT treatment, the following parameters were used to demonstrate the performance of SCWT on sewage sludge including moisture, decomposition rate (Eq. 1), organic matter and total HMs content. Low moisture and high decomposition rate can reduce the sludge volume and weight adding benefits to its ultimate disposal. As for agricultural use, a high organic matter content and low total HMs concentration in sludge is of great importance.

Decomposition rate (%) = 100×(W₀-W)/W₀                     (1)

where W₀ (mg) and W (mg) are the dry weights of sludge before and after SCWT treatment, respectively.
2.4.2. Eco-toxicity and bioavailability

It is widely recognized that the mobility, eco-toxicity and bioavailability of HMs in sludge depends not only on the total concentration but also on their existing forms (Xian, 1989; Angelova et al., 2004). Based on Tessier’s method, the eco-toxicity and bioavailability of HMs can be described as: (F1+F2) concentration of HMs represents direct toxicity and bioavailability \( (C_{bio}) \), (F3+F4) concentration relates to potential toxicity and bioavailability \( (C_{Pbio}) \), while HMs in F5 possess no toxicity \( (C_{Nbio}) \) (Vela et al., 1993; Li et al., 2012).

More attention should be paid to the weakly bonded fractions (F1+F2) as they can equilibrate with the aqueous phase and thus become more rapidly bioavailable (Sundaray et al., 2011). According to this consideration, specific quantity evaluation of HMs in sludge can be carried out by using a risk assessment code (RAC) based on the proportion of these two fractions (F1+F2) to the total amount of each metal extracted from the sludge, namely the risk index (RI) (Huang et al., 2011; Singh et al., 2005). The HMs in sludge can be classified by using RAC as no risk, low risk, medium risk, high risk and very high risk when the RI value ranges < 1, 1–10, 11-30, 31-50 and > 50, respectively. Furthermore, the ratio of (F1+F2)/F5 can be used to describe the potential lability of HMs in sludge (Obrador et al., 2001).

2.4.3. Leachability

The leachability of HMs in SR directly relates to their availability and toxicity, and can be evaluated by the standard Toxicity Characteristic Leaching Procedure (TCLP). TCLP simulates landfill conditions by using acetic acid solution (pH 2.88, liquid/solid ratio= 20:1). The leaching tests were carried out in polyethylene tubes and shaken at 200 rpm for 18 h. The supernatant was harvested by centrifugation, and then digested with \( H_2O_2 \) and concentrated HNO\(_3\) for HMs analysis.

2.5. Analysis

The moisture or water content in sludge was analyzed by weight loss when heating at 105 °C for 48 h. The concentration of each HM was determined by using ICP-MS (ELAN DRC-e, PerkinElmer) after filtration with 0.2µm membrane filter. Phosphorus was
measured with molybdate ascorbic acid method by a Shimadzu UV1800 spectrophotometer. Ammonia nitrogen was analyzed with Nessler's reagent spectrophotometry method. The organic matter was determined by burning at 873 K in a muffle. Before the determination of the content of HMs and phosphorus in LS, the samples of LS should be digested with H₂O₂ and concentrated HNO₃ to remove the dissolved organics. Each test was conducted in triplicate. The results presented in this study were the mean values ± standard deviation.

3. Results and discussion

3.1. Physicochemical property changes

Table 2 shows the changes of some physicochemical properties of sludge before and after SCWT process. It can be seen that the decomposition rate of sludge increased with temperature resulting in the decrease of organic matter from 46.09% to 21.35% (SR280) and the increase of total phosphorus in SR from 17.15 g kg⁻¹ to 25.40 g kg⁻¹ (SR280). The pH value and ammonia concentration in LS increased with the increase of temperature, possibly brought about by the decomposition of dead microbes. The color of SR was changed from grey to black, attributable to some portions of organics being converted into biochar (Hu et al., 2010). The dehydration performance of sludge after SCWT treatment was obtained by vacuum filtration followed by moisture measurement. The water content in the sludge decreased from 85.47% to 74.74%, 62.37% and 44.68% after SWCT treatment at 170 °C, 200 °C and 280 °C, resulting in sludge volume reduction approximately by 42%, 61% and 74%, respectively according to the relationship between sludge volume and water content. The results show that stabilization of sludge could be achieved along with large reduction in sludge volume after SCWT process. Based on the results obtained in this study, the higher temperature SCWT process (such as 280°C) seems to be more preferable in view of the ultimate disposal of sludge. The phenomenon, i.e. sludge stabilization and volume reduction after SCWT treatment may have some contribution to the HMs dissolution and distribution in the treated sludge, which is explored further in the following experiments.

3.2. HMs dissolution and distribution in SR and LS
The redistributions of HMs in LS and SR are shown in Figs. 1 and 2, respectively. The result indicates that HMs could be dissolved from solid sludge particles into liquid phase resulting in increased concentrations in liquid phase after the SCWT process. This observation might be brought about by the degradation of extracellular polymeric substances and thereafter some HMs binding to them were released into the liquid phase (Appels et al., 2010). The dissolution of HMs into liquid seems to be coupled with the dissolution of total phosphorus (Table 2), providing a chance for HMs adsorption (Eq. 2), precipitation (Eqs. 3 and 4) and thus immobilization in the sludge (McGowen et al., 2001).

\[
S - HPO_4^{2-} + M^{2+} \leftrightarrow S - HPO_4^{2-} + M^{2+} \quad (2)
\]

\[
HPO_4^{2-} + M^{2+} \leftrightarrow MHPO_4 \quad (3)
\]

\[
PO_4^{3-} + M^{2+} \leftrightarrow M_2(PO_4)_2 \quad (4)
\]

where S represents sludge particle, and M represents heavy metal.

As for total content, different metals behaved differently. The released amount into liquid increased with the increase of temperature for Zn, Cd or Pb, while the highest release was obtained at 200 °C for Cr, Ni or Cu. Although SCWT process seems to have some positive effect on HMs release from sludge into liquid phase, the amount of HMs in LS was very low (< 1.3% of the total) with the majority being remained (> 98.7%) in SR. Figure 2 shows that the HMs contents increase in SR with the increase of SCWT temperature, i.e. HMs tend to be accumulated in SR after SCWT treatment. This observation signals an increased environmental risk of sludge with the increase of SCWT treatment temperature if only using the total HMs content as an assessment index.
3.3. Fraction of HMs and environmental risk analysis

3.3.1. Fraction and fractional transformation of HMs

By using the Tessier’s method, different fractions of the six HMs (Cr, Ni, Cu, Zn, Cd and Pb) in the sludge were extracted and analyzed before and after SCWT treatment (Fig. 3).

The main existing forms in CSR were found to be different for the six HMs. 64.69% of Cu was bound to organic and sulfide substances (F4), which may be attributed to its high stability of complexion with organic matter (Staelens et al., 2000; Fuentes et al., 2008) and sulfide. Pb was found to be mainly existed in F4 (63.90%), and the majority of Ni, about 60.20% was in the residual fraction (F5). Zn was found mainly in three forms in the sludge, namely F5 (39.48%), F4 (31.63%) and F3 (26.95%). Two forms, F5 (47.33%) and F4 (45.07%) were the main existing ones for Cr in the sludge. About half of Cd was bound to iron and manganese oxides (F3, 53.51%). It should be stated that all the six HMs are very low at exchangeable (F1, < 5%) and carbonate bounded (F2, <2%) fractions in CSR (except for Cd in F2, 13.30%).

Figure 3 also illustrates the effect of SCWT treatment on the redistribution of HMs in SR. The contents of Cu and Pb were all sharply decreased in the first 3 fractions while increased in F5 after SCWT, especially for Cu in F1 and Pb in F2 with the contents decreased negatively with treatment temperature and for Pb in F5 increased positively with temperature. In addition, SCWT exerted different effect on both Cu and Pb in F4, and the F4 of Cu decreased after treatment at 170°C and 280°C while Pb content in F4 was reduced at temperatures higher than 200°C.

In the case of Zn, the contents decreased in F1 (except in SR170), F2 and F3 while increased in F4 and F5. A considerable reduction of Ni content was observed in F1 (except treatment at 200°C), F2, F3 and F4, and the corresponding content of F5 was significantly increased.

As for Cr and Cd, SCWT showed some positive effect on the reduction of F1 and F2 fractions in the sludge. After treatment at 170°C and 280°C, Cr content was observed to decrease in F4 and increase in F5 (Fig. 3). The percentages of F3 (except for 280°C) and F4 of Cd in the sludge increased after SCWT treatment, while the F5 fraction firstly increased and then decreased with temperature. Still, the major form of Cd was F4 after treated at 280°C, mostly contributed by the conversion from its F3 fraction.
From the above results, it can be concluded that the weakly bounded forms (F1 and F2) of HMs are all decreased in sludge after SCWT treatment except for F1 fractions of Zn and Ni after treatment at 170°C and 200°C. The F5 fractions of HMs are all increased after SCWT with the exception of Zn and Cr in SR200 and Cd in both SR200 and SR280. Most importantly, the proportions of (F1+F2) for the six HMs are at the lowest levels in all the SR280 samples (Fig. 3); and the corresponding first four fractions of HMs are all decreased with F5 increased accordingly after SCWT treatment at 280 °C (except for Cd). The results indicate that SCWT treatment at 280 °C is the best condition with respect to the toxicity and bioavailability of HWs (i.e. existing forms).

From above results, it can be inferred that HMs in sludge could be transformed from weakly bounded fractions (especially F1 and F2) to a more stable state (F5) during SCWT process, implying that some complexation might occur between HMs and the crystal lattices of the residual solid phase. The above fractional transformation and redistribution of HMs can possibly realize the immobilization of HMs in sludge, and some anions such as phosphate (Table 2) and silicate contained in the sludge may recombine with these HMs to become stable precipitate with very low solubility (Chen et al., 2007; Geebelen et al., 2006). The detailed mechanisms are still under investigation.

3.3.2. Eco-toxicity and bioavailability assessment

Table 3 shows the variation of C_{bio}, C_{phio} and C_{Nhio} of HMs in the sludge after SCWT treatment, which can be used to assess the eco-toxicity and bioavailability. The significance of temperature on the fraction distribution of HMs is indicated by F- and p- values obtained by one-way analysis of variance (ANOVA) with the experimental data (Table 3). The statistical results show that treatment temperature significantly influences the concentrations of the three portions of HMs during SCWT process except for Pb in C_{phio}.

From Table 3 and Fig. 3, the proportions of each metal existed in the first two fractions (F1+F2) are very low (less than 5%, except for Cd, 16.67%) in CSR. The concentrations of the HMs in these two parts (except for Zn in SR170) were all decreased after SCWT treatment, indicating its positive effect on reducing the direct toxicity and bioavailability of these HMs in sludge (Li et al., 2012; Huang et al., 2011). The concentration of Cr in SR200 was the same as that in CSR denoting that
the direct toxicity of Cr was not reduced after SCWT at 200 °C. The SCWT treatment at 280 °C exhibited the most significant effect on reducing this toxicity due to the lowest C_{bio} of HMs obtained in SR280, and the HMs in the bioavailable fraction was decreased by 53.75 %, 69.32 %, 81.15 %, 64.75 %, 98.37 % and 61.18% for Cr, Ni, Cu, Zn, Cd and Pb, respectively.

Except that Ni was mainly existed in the non-toxicity fraction, the other five metals were mainly found in the forms with potential toxicity in CSR (Table 3). After SCWT process, the C_{Pbio} values of Cu and Cd increased with temperature while C_{Pbio} of Pb remained relatively stable; the C_{Pbio} values of Cr and Zn slightly decreased at 170 °C and then increased at higher temperatures. Although having an increase trend, the C_{Pbio} values of Ni in SR were less than those in CSR. And the potential toxicity of Cr, Cu, Zn and Cd in sludge was increased after SCWT treatment at temperatures higher than 200°C. This increase was more significant for Cd as its concentration even doubled in SR280. The potential toxicity of Ni was reduced while that of Pb kept almost unchanged. Restated, the total concentrations of Ni and Pb increased after SCWT (Fig. 2), but the proportion of these two metals relating to potential toxicity was sharply decreased (corresponding to large increase in F3+F4, Fig. 3). This observation indicates that the SCWT process seems to have more significant effect on reducing toxicity and enhancing immobilization of Pb and Ni in the sludge.

All the C_{Nbio} concentrations of HMs increased (except Cd in SR200) after SCWT treatment (Table 3), which would be more meaningful for HWs immobilization if the proportions of these metals in no toxicity state also increased. From Fig.3, Pb and Ni associated with no toxicity were both increased with temperature, and other HMs in this fraction were increased after being treated at 170 °C and 280 °C. Besides, the percentage increase in fraction F5 together with the decrease in the easily available fractions (F1 + F2) (Fig. 3), thus a lower ratio of (F1+F2)/F5, implies a reduced potential lability of HMs in sludge after the SCWT treatment (Obrador et al., 2001). Taken the reduction effect on the direct toxicity of HWs into consideration, the SCWT process under 280°C is proposed for the best HMs immobilization in sewage sludge in this study.

3.3.3. RAC analysis

The environmental risk of these HMs in CSR and SR can also be evaluated by RAC using (F1+F2) ratios because these weakly bonded fractions are easily affected by
ionic strength and vulnerable to pH changes in soil environment resulting in a very high mobility (Lasheen and Ammar, 2009; Yuan et al., 2011). Based on the results of Fig. 3, all the metals showed low risk in CSR, and the risk of the six HMs was greatly lowered after SCWT, especially for Cu, Zn, Ni and Cd with their risk changed from low risk to no risk. The easily bioavailable HMs (F1 and F2) were mainly transformed into the relatively stable heavy metal fractions (F3, F4 and F5) after SCWT process, resulting in the decrease of direct toxicity and environmental risk of HMs. In addition, no environment risk of HMs in SR was attained after being treated at 280 °C except for Pb. Still, the risk of Pb remained at a very low level (RI=RI<1-10).

3.4. Leaching test and leaching toxicity

The leaching toxicity of HMs in CSR and SR was determined with the standard TCLP test shown in Table 4 and Fig. 4.

The metal concentrations in the leachate extracted from CSR were 0.30, 11.75, 5.05, 2.25, 0.72 and 0.03 mg kg\(^{-1}\) for Cr, Ni, Cu, Zn, Cd and Pb, respectively. The concentrations of leachable HMs were lower than the USEPA permissible limits (SW-846) expect for that of Ni in CSR which exceeded the limit value by about two times. The concentrations of leached Cu, Zn and Cd from sludge were declined after SCWT process, while Cr and Ni behaved slightly different. The concentration of Cr and Ni in leachate from SR170 and SR280 decreased but increased from SR200. In the case of Pb, although the leachable contents increased in SR170 and SR200, the concentrations were very low (0.045 and 0.081 mg kg\(^{-1}\), respectively) showing no risk, and the leachable content was decreased in SR280. Briefly, all the lowest concentrations of leachable HMs were found in SR280, and the concentrations decreased by 97.46%, 93.91%, 86.14%, 73.67%, 71.93% and 10.71% for Cu, Cd, Zn, Cr, Ni and Pb, respectively, indicating the release of HMs (except for Pb) was significantly decreased. The SCWT process at 280 °C exhibited the most significant effect on reducing the leaching risk of HMs to the environment.

The leaching rates of the six HMs are shown in Fig. 4. This indicator is defined as the ratio of individual metal concentration in the leachate to the total concentration of this metal (Huang et al., 2011), which can be used to reflect the proportion of HMs in the leachable state. The decreased leaching rates of Ni, Cu, Zn and Cd indicate the decrease in the leachable fraction of these metals, which is in contrast to the leachability increase of Cr from SR200 and Pb from SR170 and SR200. From Fig. 4
and Table 4, it’s obvious that the HMs related to leachable fraction were at the lowest proportions in the sludge after SCWT treatment at 280 °C, clearly showing the reduced leaching toxicity and leachable fraction of HMs in sludge after SCWT process (especially 280 °C). This observation coincides with the declination of weakly bounded fractions (F1+F2) shown in Fig 3.

3.5. Implication of this study to practice

In this study, the risk of HMs in sludge is mainly assessed both by their total concentrations and existing forms. The experimental data shows that the tested HMs can be transformed into more stable existing forms in the sludge after SCWT process, although the total metal concentrations tend to increase due to the large decrease of sludge volume. This phenomenon is also in agreement with the decreased bioavailability and leachability tests of the treated sludge. Restated, the total concentrations of HMs remained in the solid phase are still lower than the USEPA permissible limits (SW-846). The results up to present suggest that SCWT could be a promising sludge treatment method for HMs immobilization, after which the treated sludge can be more appropriate for agricultural application due to much less or no risk of HMs to the environment. Our followed-up research will focus on the mechanisms involved in this process and the enhancement effect of some additives on HMs immobilization in the sludge.

4. Conclusions

The present study indicates that SCWT is a promising method for sewage sludge disposal. After SCWT treatment, organics stabilization and volumetric reduction of sludge can be achieved. Although the majority of HMs is still accumulated in the solid phase, most of the HMs can be transformed from easily available fractions to relatively stable states resulting in large decrease in direct toxicity and environmental risk of the treated sludge. Moreover, the leachable fraction and the leaching toxicity of HMs are also largely decreased after SCWT. Among the tested conditions, SCWT at 280°C is preferable for the final disposal of sludge.

References


Table 1. Physicochemical characteristics of the sewage sludge.

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<tr>
<th>Parameters</th>
<th>Value $^a$</th>
</tr>
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<tbody>
<tr>
<td>Solid content (%)</td>
<td>14.53 ± 0.39</td>
</tr>
<tr>
<td>Organic matter (%)</td>
<td>46.09 ± 0.16</td>
</tr>
<tr>
<td>Total phosphorus (g kg$^{-1}$ dry weight)</td>
<td>17.15 ± 1.84</td>
</tr>
<tr>
<td>pH</td>
<td>6.43 ± 0.21</td>
</tr>
<tr>
<td>HMs (mg kg$^{-1}$ dry weight)</td>
<td></td>
</tr>
<tr>
<td>Cr</td>
<td>150.18 ± 1.22</td>
</tr>
<tr>
<td>Ni</td>
<td>638.56 ± 96.15</td>
</tr>
<tr>
<td>Cu</td>
<td>415.00 ± 29.46</td>
</tr>
<tr>
<td>Zn</td>
<td>750.65 ± 59.59</td>
</tr>
<tr>
<td>Cd</td>
<td>73.02 ± 0.63</td>
</tr>
<tr>
<td>Pb</td>
<td>122.14 ± 9.06</td>
</tr>
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</table>

$^a$ Mean ± standard deviation.
Table 2. Physicochemical property changes of sewage sludge before and after SCWT treatment.

<table>
<thead>
<tr>
<th></th>
<th>CK  a</th>
<th>SCWT170 b</th>
<th>SCWT200 b</th>
<th>SCWT280 b</th>
</tr>
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<tbody>
<tr>
<td>Decomposition rate (%)</td>
<td>0</td>
<td>14.32± 0.22</td>
<td>18.99± 0.19</td>
<td>32.47±0.12</td>
</tr>
<tr>
<td>Moisture (%) (SR c)</td>
<td>85.47± 0.39</td>
<td>74.74± 0.25</td>
<td>62.37± 0.31</td>
<td>44.68± 0.46</td>
</tr>
<tr>
<td>Organic matter (%) (SR c)</td>
<td>46.09 ± 0.16</td>
<td>34.98± 0.21</td>
<td>29.27± 0.14</td>
<td>21.35± 0.10</td>
</tr>
<tr>
<td>pH (LS d)</td>
<td>N.D.</td>
<td>7.61± 0.18</td>
<td>8.52± 0.13</td>
<td>9.17± 0.07</td>
</tr>
<tr>
<td>NH₄⁺-N (LS, mg l⁻¹)</td>
<td>N.D.</td>
<td>2357.24±17.91</td>
<td>2586.24±8.99</td>
<td>3566.17±7.37</td>
</tr>
<tr>
<td>Total phosphorus (mg l⁻¹) (LS d)</td>
<td>N.D.</td>
<td>12.48±1.04</td>
<td>15.81±0.83</td>
<td>30.43±2.17</td>
</tr>
<tr>
<td>Total phosphorus (g kg⁻¹) (SR c)</td>
<td>17.15±1.84</td>
<td>20.02±0.94</td>
<td>21.17±2.01</td>
<td>25.40±1.29</td>
</tr>
</tbody>
</table>

a CK-control experiment (without SCWT treatment). b SCWT170, SCWT200, and SCWT280 are the sludges after SCWT treatment under 170, 200, and 280 °C, respectively. c SR-solid residue. d LS-liquid sample after vacuum filtration. e N.D., No determination because of little liquid (LS) could be obtained from control samples due to the high solids content and high viscosity of sludge used in this study.
Table 3. Concentrations of HMs in bioavailable (C_{bio}), potential bioavailable (C_{Pbio}) and non-bioavailable (C_{Nbio}) fractions in sludge before and after subcritical water treatment (unit: mg kg^{-1}) (n=3, \( \alpha < 0.05 \)).

<table>
<thead>
<tr>
<th></th>
<th>Cr</th>
<th>Ni</th>
<th>Cu</th>
<th>Zn</th>
<th>Cd</th>
<th>Pb</th>
</tr>
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<tbody>
<tr>
<td>C_{bio} CSR a</td>
<td>1.60±0.35</td>
<td>24.38±0.82</td>
<td>12.20±0.26</td>
<td>14.98±3.10</td>
<td>12.25±0.44</td>
<td>5.95±0.92</td>
</tr>
<tr>
<td>SR170 b</td>
<td>1.34±0.18</td>
<td>14.79±2.36</td>
<td>6.15±0.75</td>
<td>22.65±1.91</td>
<td>2.12±0.37</td>
<td>2.00±0.05</td>
</tr>
<tr>
<td>SR200 b</td>
<td>1.59±0.24</td>
<td>19.67±0.55</td>
<td>4.36±1.18</td>
<td>10.90±0.51</td>
<td>1.36±0.02</td>
<td>2.82±0.88</td>
</tr>
<tr>
<td>SR280 b</td>
<td>0.74±0.11</td>
<td>7.48±0.65</td>
<td>2.30±1.27</td>
<td>5.28±1.19</td>
<td>0.20±0.01</td>
<td>2.31±0.34</td>
</tr>
<tr>
<td>ANOVA</td>
<td>F</td>
<td>p-value</td>
<td>&lt;0.001</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>C_{Pbio} CSR a</td>
<td>77.50±1.71</td>
<td>229.74±1.30</td>
<td>266.61±1.41</td>
<td>442.90±32.30</td>
<td>56.88±2.50</td>
<td>88.06±3.09</td>
</tr>
<tr>
<td>SR170 b</td>
<td>70.86±6.08</td>
<td>165.24±18.75</td>
<td>274.50±33.04</td>
<td>432.41±55.27</td>
<td>79.59±1.78</td>
<td>96.17±18.16</td>
</tr>
<tr>
<td>SR200 b</td>
<td>95.82±7.67</td>
<td>187.8±13.94</td>
<td>334.81±27.39</td>
<td>533.73±25.41</td>
<td>89.78±0.80</td>
<td>86.53±7.99</td>
</tr>
<tr>
<td>SR280 b</td>
<td>102.44±10.55</td>
<td>210.04±23.06</td>
<td>384.5±34.30</td>
<td>633.3±76.07</td>
<td>108.92±1.35</td>
<td>91.65±6.57</td>
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<tr>
<td>ANOVA</td>
<td>F</td>
<td>p-value</td>
<td>0.0002</td>
<td>0.004</td>
<td>0.004</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>C_{Nbio} CSR a</td>
<td>71.08±2.41</td>
<td>384.44±0.48</td>
<td>137.28±1.93</td>
<td>321.11±10.17</td>
<td>4.58±1.82</td>
<td>29.05±0.33</td>
</tr>
<tr>
<td>SR170 b</td>
<td>92.98±6.44</td>
<td>507.41±21.11</td>
<td>188.90±31.92</td>
<td>381.67±26.01</td>
<td>7.04±1.85</td>
<td>43.04±9.52</td>
</tr>
<tr>
<td>SR200 b</td>
<td>81.80±7.91</td>
<td>539.38±13.39</td>
<td>177.22±27.86</td>
<td>377.82±59.45</td>
<td>2.05±0.27</td>
<td>65.78±8.60</td>
</tr>
<tr>
<td>SR280 b</td>
<td>116.67±10.65</td>
<td>705.93±23.70</td>
<td>244.46±33.36</td>
<td>487.93±116.47</td>
<td>5.22±0.41</td>
<td>90.26±5.95</td>
</tr>
<tr>
<td>ANOVA</td>
<td>F</td>
<td>p-value</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>0.018</td>
<td>0.04</td>
</tr>
</tbody>
</table>

\(^a\) CSR—Solid residue in control experiment. \(^b\) SR170, SR200, and SR280 denote the solid residue conditions after SCWT treatment under 170, 200, and 280 °C, respectively.
Table 4. Concentrations of heavy metals released in TCLP tests (unit: mg kg\(^{-1}\)).

<table>
<thead>
<tr>
<th>HMs</th>
<th>CSR (^a)</th>
<th>SR170 (^b)</th>
<th>SR200 (^b)</th>
<th>SR280 (^b)</th>
<th>Permissible limits (^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cr</td>
<td>0.300 ± 0.061</td>
<td>0.141 ± 0.006</td>
<td>0.424 ± 0.053</td>
<td>0.079 ± 0.018</td>
<td>5.0</td>
</tr>
<tr>
<td>Ni</td>
<td>11.751 ± 1.62</td>
<td>3.881 ± 0.357</td>
<td>12.165 ± 1.596</td>
<td>3.298 ± 0.672</td>
<td>5.0</td>
</tr>
<tr>
<td>Cu</td>
<td>5.047 ± 0.709</td>
<td>0.600 ± 0.035</td>
<td>1.387 ± 0.393</td>
<td>0.128 ± 0.042</td>
<td>Not enlisted</td>
</tr>
<tr>
<td>Zn</td>
<td>2.251 ± 0.432</td>
<td>0.391 ± 0.047</td>
<td>1.429 ± 0.297</td>
<td>0.312 ± 0.049</td>
<td>5.0</td>
</tr>
<tr>
<td>Cd</td>
<td>0.722 ± 0.092</td>
<td>0.070 ± 0.001</td>
<td>0.171 ± 0.050</td>
<td>0.044 ± 0.011</td>
<td>1.0</td>
</tr>
<tr>
<td>Pb</td>
<td>0.028 ± 0.001</td>
<td>0.045 ± 0.002</td>
<td>0.081 ± 0.005</td>
<td>0.025 ± 0.002</td>
<td>5.0</td>
</tr>
</tbody>
</table>

\(^a\) CSR-Solid residue in control experiment.

\(^b\) SR170, SR200, and SR280 denote the solid residue conditions after SCWT treatment under 170, 200, and 280 °C, respectively.

Figure captions

Fig. 1. The total contents of heavy metals in liquid phase before and after SCWT treatment. LS-liquid sample after vacuum filtration, LS170, LS200, and LS280 denote the total amount of specific heavy metal in the liquid phase after SCWT treatment under 170, 200, and 280°C, respectively.

Fig. 2. The total contents of heavy metals in solid residue before and after SCWT treatment. CSR-solid residue in control experiment (without SCWT treatment), SR-solid residue. SR170, SR200, and SR280 indicate the total amount of specific heavy metal in the solid residue after SCWT treatment under 170, 200, and 280°C, respectively.

Fig. 3. Fractional distribution changes of each metal in sewage sludge before and after SCWT treatment. CSR-solid residue in control experiment (without SCWT treatment), SR-solid residue. SR170, SR200, and SR280 indicate the change of corresponding heavy metal fraction in the solid residue after SCWT treatment under 170, 200, and 280°C, respectively.

Fig. 4. Leaching rate of heavy metals in TCLP tests. CSR-solid residue in control experiment (without SCWT treatment), SR-solid residue. SR170, SR200, and SR280 indicate the leachability change of heavy metals in the solid residue after SCWT treatment under 170, 200, and 280°C, respectively.
Fig. 1 Shi et al.
Fig. 2 Shi et al.
Fig. 3 Shi et al.
Fig. 4 Shi et al.