Lead Immobilization in Artificial Contaminated Soil Using Sulfur-Impregnated Carbonaceous Rice Straw

T. Wajima

Abstract—A novel carbonaceous immobilizing agent for heavy metal contaminated soil was prepared from rice straw using sulfur immersion and pyrolysis, and the lead immobilization in artificial contaminated soils using sulfur-impregnated carbonaceous material was estimated. The rice straws were cut to 1 cm pieces, and then immersed in 0.1 - 1 M K₂S solution for 0 - 24 h to prepare sulfur-immersed materials. The immersed-materials were heated at 400 °C for 1 h in nitrogen gas to produce the sulfur-impregnated carbonaceous material by pyrolysis. The abilities of the product to immobilize lead from aqueous solution were examined to obtain the product with high lead immobilization ability. With increasing K₂S concentration, the immobilization ability of the product for lead gradually increases and then above 0.5 M K₂S those are almost constant, while 15 min is sufficient for the immersed time to obtain the product with high lead immobilization ability. The product prepared from material immersed in more than 0.5 M K₂S solution for less than 15 min has a maximum immobilization ability for lead ion. The lead immobilization using the sulfur-impregnated product is sustainable due to the formation of leadbilitite [Pb₃(SO₄)(CO₃)₂(OH)₆] and anglesite [PbSO₄]. The product can immobilize lead ion in various artificial lead contaminated soils. By mixing artificial lead contaminated soil with the sulfur-impregnated product, the eluted solution became neutral, and the eluted concentrations of lead ion dropped below the Japanese elution standard for soil.

Index Terms—Soil contamination, lead immobilization, sulfur-impregnation, rice straw, pyrolysis.

I. INTRODUCTION

Soil contamination with heavy metals is a worldwide problem. Accumulation of heavy metals in soils affect soil ecology, agricultural productivity, quality of agricultural products, water resources, and serious health problems for human and animal [1]. In the United States, for example, approximately 63 % of the sites on the National Priority List (NPL) for the treatment of contaminated soils are approximately 63 % of the sites on the National Priority List (NPL) for the treatment of contaminated soils are contaminated by metals, and lead is the most common metal, found at 15 % of the sites. In Japan, according to a report by the Ministry of Environment, 43 % of the contaminated sites that exceed the environmental quality standards are contaminated by lead compounds. Soil contamination by lead compounds is also prevalent in other developed and developing countries where lead compounds are used extensively in industrial activities without careful contamination management.

Lead is a ubiquitous heavy metal pollutant in soils due to their use widely. The primary sources of Pb contamination include industrial activities such as mining, smelting of metals, and the use of Pb-containing products such as paints, lead-acid batteries, bullets, gasoline and pesticides [2]. It can damage human nervous (especially children) and reproductive systems [3]. The high concentration of Pb in the soil poses risks to human and animal health by the leaching of metals from the soil into water and the consumption of edible plants growth in the contaminated soil. Therefore, proper remediation is necessary to reduce metal availability in soil for protecting human health.

Among available remediation technologies, in situ immobilization of heavy metals using a chemical amendment can be a cost-effective and environmentally sustainable remediation approach for the immobilization of heavy metals by reducing the mobility and availability. This immobilization technique may provide a long-term remediation solution if low solubility minerals and/or stable precipitates are produced in situ [4]. Therefore, the choice of the soil amendments need that the amendments must reduce heavy metals transfers from contaminated soils to the surface water or groundwater and uptake by plants and organism. The most common agents for lead immobilization are phosphate-based compounds [5]-[7]. Although these studies have successfully demonstrated that phosphates effectively immobilize lead in various contaminated soils, there are several concerns about the use of phosphate as agents for lead immobilization. First, from an environmental protection point of view, excessive supplied phosphate can lead to eutrophication in the natural environment. Second, from a resource conservation point of view, phosphate is becoming a precious element, especially in countries such as Japan that import a major portion of their phosphate requirements. For these reasons, research on other types of lead immobilization agents is needed. Up to now, soil amendments, for example, lime [8], calcium carbonate [9], red-mud [10], fly ash [11] and so on, may decrease leachable concentrations of contaminants and thus reduce the detrimental effects of heavy metals on environmental receptors, such as microorganisms, plants, animals, water and humans [12].

In previous studies, sulfur-impregnated adsorbents with high removal abilities for heavy metals were prepared from paper sludge, cedar bark and rice husk using K₂S solution [13]-[15]. According to the Pearson theory, the sulfur, as a soft base, should interact with heavy metals such as Zn²⁺, Pb²⁺, Cd²⁺ and Ni²⁺ (soft acids) rather than with oxygen (a hard base) in the activated carbon. From these results, it would be possible to produce a low cost agent to immobilize...
the metal ions in contaminated soils from agricultural waste. From these background, in this study, we attempted to prepare the lead immobilization agent from rice straw, which is produced as a by-product of rice production, using sulfur-impregnation, and lead immobilization ability of the sulfur-impregnated product was estimated using artificial lead contaminated soils.

II. MATERIALS AND METHODS

A. Raw Sample

Raw rice straw, which was collected from a cultivated area in Akita prefecture, Japan, first cut into 1 cm length, was then washed with distilled water, and dried and stored for use. Properties of rice straw sample are shown in Table I, and mineralogical composition of rice straw is shown in Fig. 1, which indicates that rice straw mainly composed of cellulose. All reagents were purchased from Wako Chemical Co., Japan at analytical grade.

![Graph showing XRD patterns of rice straw.](image)

Fig. 1. XRD patterns of rice straw.

TABLE I: PROPERTIES OF RICE STRAW SAMPLE

| Moisture | Ash | Volatile matter + Fixed carbon |
|----------|-----|-------------------------------|
| Content (%) | 7.3 | 11.6 |
|          | 39.2 | 6.3 |
|          | 0.6  | 0.8 |

B. Sulfur Impregnation

20 g of sample was immersed in 200 mL of 0.1 - 1 M K$_2$S solution for 0 - 24 h, then filtered, and dried in a drying oven overnight to obtain sulfur-immersed samples. These samples were pyrolyzed using a horizontal reactor. Sulfur-immersed samples were put in a ceramic board, and installed in a transparent quartz tube of 0.45 mm inside diameter and 1 m in length. Before pyrolysis, N$_2$ gas was injected into the tube for 30 min at a rate of 1.0 L/min to replace the air in the tube. The board was heated in an electric furnace at 400 °C for 1 h, with a continuous flow of N$_2$ gas at a rate of 1.0 L/min. After heating, the solid was cooled to room temperature with a steady N$_2$ gas flow (1.0 L/min) in the tube, then washed with distilled water and dried in drying oven overnight to obtain the sulfur-impregnated material.

The abilities of the material for immobilization of lead ion from aqueous solution were examined as follows. 0.1 g of the sample was added to 10 mL of Pb(NO$_3$)$_2$ solution with 10 mM in 50 mL centrifuged tube, and was shaken in a reciprocal shaker for 24 h. After shaking, the slurry was centrifuged, and the pH of the supernatant and the concentration of Pb$^{2+}$ in the supernatant were measured by pH meter (D-53, Horiba) and atomic absorption spectrophotometer (AAS) (AAAnalyst200, PerkinElmer), respectively. The immobilization ratios of Pb$^{2+}$ were calculated using the following equation:

\[ R = \frac{C_0 - C_e}{C_0} \times 100 \]  

Here, \( R \) is Immobilization ratio of Pb$^{2+}$ (%), \( C_0 \) is Initial concentration of Pb$^{2+}$ in the solution (mg/L), and \( C_e \) is Measure concentration of Pb$^{2+}$ in the solution (mg/L).

The samples were analyzed by powder X-ray diffraction (XRD) with monochromate CuKα radiation (Ultima IV, Rigaku) and Fourier transform infrared spectrometry (FT-IR) (Nicolet iS5, Thermo Fisher Scientific). The morphologies of the samples were observed using a scanning electron microscope (SEM) (JSM-6510A, JEOL) equipped with an energy-dispersive X-ray spectrometer (EDS).

C. Lead Immobilization

Five soil samples, kaolinite, Akadama soil, Kanuma soil, black soil and river sand, were used to prepare artificially lead-contaminated soils. Lead-contaminated soil was artificially prepared by mixing 30 g of soil sample and 300 mL of 1000 mg-Pb/L aqueous solution (provided by Pb(NO$_3$)$_2$) using a rotary stirrer for 6 h. After mixing, the mixture stands at room temperature for 7 days, and then filtrate. The concentration of Pb$^{2+}$ in the filtrate was measured by AAS to calculate the content of lead in artificially contaminated soils. The solid was dried in air for 7 days to obtain artificially contaminated soil.

1 g of artificially contaminated soil without or with 0.01 – 0.1 g of sulfur-impregnated product was added into 50 mL of a conical flask, and 10 mL of distilled water was then poured. The flask was shaken with shaking incubator at 50 °C for 6 h, and then filtrate. The pH of the filtrate was measured by pH meter, and the eluted concentration of Pb$^{2+}$ in the filtrate is measured by AAS.

III. RESULTS AND DISCUSSION

A. Lead Immobilization of Sulfur-Impregnated Product

![Graph showing Lead immobilization (%) vs. K$_2$S concentration (mol/L).](image)

Fig. 2. Lead immobilization of the product from rice straw via pyrolysis of the straw immersed in K$_2$S solution with various concentrations.
Fig. 2 shows the lead immobilization of the product from rice straw via pyrolysis of the straw immersed in K$_2$S solution with various concentrations. With increasing K$_2$S concentration to 0.5 mol/L, lead immobilization of the product increases, and be almost constant (approximately 80 %) above 0.5 mol/L K$_2$S solution.

Fig. 3 shows the XRD patterns of (a) immersed straws, (b) pyrolysed immersed straws, (c) the products and (d) the products after lead immobilization test. The straws after K$_2$S immersion have cellulose peaks, and the height of the peaks for cellulose decreases with increasing K$_2$S concentration (Fig. 3 (a)). After pyrolysis, the pyrolyzed straws have amorphous structure, and the peaks of carbon disulfide [CS$_2$] appears in the pyrolyzed straw immersed in above 0.5 mmol/L K$_2$S solution (Fig. 3 (b)). The product after washing and drying the pyrolyzed immersed straws also indicates amorphous peak, and the peaks of carbon disulfide remain in the product from the straw immersed in above 0.5 mmol/L K$_2$S solution (Fig. 3 (c)). After lead immobilization test, leadhilite [Pb$_4$SO$_4$(CO$_3$)$_2$(OH)$_2$] and anglesite [PbSO$_4$] appear on the product from the straw immersed in 0.1 - 0.75 mol/L K$_2$S solution and 0.75 - 1.0 mol/L K$_2$S solution, respectively. It would be considered that lead was immobilized by the formation of leadhilite and anglesite due to the reaction of lead ion with sulfur functional on the surface of the product to leadhilite and carbon disulfide to anglesite. In addition, carbon disulfide would precipitate by high concentration of K$_2$S in the solution. From these results, the appropriate product for lead immobilization can be prepared from the straw immersed in 0.5mol/L K$_2$S solution.

Fig. 4 shows the lead immobilization of the product from rice straw via pyrolysis of the straw immersed in 0.5 mol/L K$_2$S solution for various times. With increasing immersion time, lead immobilization of the product rapidly increases, and then be almost constant (about 80 %) after 15 min.

Fig. 5 shows the XRD patterns of (a) immersed straws, (b) pyrolysed immersed straws, (c) the products and (d) the products after lead immobilization test. The straws after K$_2$S immersion have cellulose peaks (Fig. 5 (a)), the pyrolyzed straws have amorphous structure with the peaks of carbon disulfide (Fig. 5 (b)) and the product indicates amorphous peak without the peaks of carbon disulfide (Fig. 5 (c)). After lead immobilization test, leadhilite [Pb$_4$SO$_4$(CO$_3$)$_2$(OH)$_2$]
appears on the product from the straw immersed in 0.5 mol/L K$_2$S solution after 15 min immersion, which means that the appropriate product for lead immobilization can be prepared from the straw immersed in 0.5 mol/L K$_2$S solution within 15 min immersion.

Fig. 5 shows the lead immobilization of rice straw, sulfur-immersed straw and product. Lead immobilization of rice straw and sulfur-immersed straw are 38 % and 42 %, respectively, while that of the product is 80 %, which means that pyrolysis with sulfur content is important for lead immobilization.

Fig. 7 shows the lead immobilization of rice straw, sulfur-immersed straw and product. Lead immobilization of rice straw and sulfur-immersed straw are 38 % and 42 %, respectively, while that of the product is 80 %, which means that pyrolysis with sulfur content is important for lead immobilization.

Fig. 8 shows that FT-IR spectra of rice straw, sulfur-immersed straws, and the products. These exhibits absorption bands with 3350 – 3550 cm$^{-1}$ due to hydroxyl groups. Rice straw and immersed straw indicate the absorption bands of methyl and methylene groups in the range 2960 - 2850 cm$^{-1}$, while these bands disappear in pyrolyzed immersed straw and product due to the pyrolysis of organic contents. Aromatic C=C group absorbs in the range around 1640 cm$^{-1}$ and 1360 cm$^{-1}$, carboxyl C=O groups in the range around 1730 cm$^{-1}$, and these bands become stronger after pyrolysis due to the carbonization. After sulfur-immersion, the absorption band of 710 cm$^{-1}$ for C=S group and that of 1275 - 1303 cm$^{-1}$ for C=S group appear. Therefore, carboxyl and sulfur functional group was formed on the surface of product.

Fig. 9 shows the SEM photos of (a) rice straw, (b) sulfur-immersed straws, and (c) the products. It can be observed that rice straw has spherical particles on the surface (Fig. 9 (a)) and these particles are covered with many short needle-like crystals after K$_2$S immersion (Fig. 9 (b)). The product has some pores without spherical particles with needle-like crystals (Fig. 9 (c)). It is noted that the high Si
content in raw rice straw, high contents of K and S in sulfur-immersed straw, and low Si content were detected by EDS analysis. It would be considered that needle-like K$_2$S crystals cover the silica particles on the surface of rice straw after K$_2$S immersion, and silica particle is reacted with K to form potassium silicate during the pyrolysis to remove by dissolving on washing process.

![FT-IR spectra](image)

**Fig. 8.** FT-IR spectra of rice straw, sulfur-immersed straws, and the products.

Therefore, the lead immobilization of the product is caused by porous structure with sulfur functional groups.

![SEM photos](image)

**Fig. 9.** SEM photos of (a) rice straw, (b) sulfur-immersed straws, and (c) the products.

### B. Lead Immobilization

The lead immobilization of the obtained product was examined using artificial contaminated soils. Table II shows the lead content of artificial contaminated soils used in this experiment. The order of lead content is black soil (9799 mg/kg) > akadama soil (8347 mg/kg) > kanuma soil (6969 mg/kg) > river sand (4185 mg/kg) > kaolinite (2693 mg/kg).

| Soil          | Lead content (mg/kg) |
|---------------|-----------------------|
| Kaolinite     | 2693                  |
| Akadama soil  | 8347                  |
| Kanuma soil   | 6969                  |
| Black soil    | 9799                  |
| River sand    | 4185                  |

Table II: Lead Contents of Artificial Contaminated Soil in This Experiment

The effect of sulfur-impregnated product addition on the lead immobilization were determined. Table III shows the pH of the solution after elution test with addition of sulfur-impregnated product. Without addition of the product, pH of the solution is weak acidic (pH 4-5), and with increasing addition of the product, pH of the solution increased. With addition of 0.1 g product, pHs of the solution using akadama soil, kanuma soil and black soil are neutral, while those using kaolinite and river sand are weak alkaline. It would be considered that the product is alkaline material and three soils have pH buffering ability.

| Lead concentration (mg/L) |
|---------------------------|
| Kaolinite                 |
| 26.7                      |
| 63.5                      |
| 85.5                      |
| 1.0                       |
| 1.1                       |
| 0.5                       |
| 11.7                      |
| 14.4                      |
| 0.9                       |
| 1.0                       |
| 0.3                       |
| 0.6                       |
| 0.7                       |
| 0.2                       |
| 1.0                       |
| N.D.                      |
| N.D.                      |
| N.D.                      |
| N.D.                      |

Table IV shows the eluted lead concentrations after elution test using artificial contaminated soils. It is noted that Japanese standard of soil elution is less than 0.01 mg/L. Without addition of the product, high concentrations of lead were eluted from kanuma soil (85.5 mg/L), akadama soil (63.5 mg/L) and kaolinite (26.7 mg/L), while elution from black soil (1.0 mg/L) and river sand (1.1 mg/L) are low. With increasing addition of the product, the lead elution decreases, and with addition of 0.1 g product, the lead elution from all samples are not detected using AAS. It would be considered that the product can immobilized lead ion in various soils.

| Lead concentration (mg/L) |
|---------------------------|
| Kaolinite                 |
| 26.7                      |
| 63.5                      |
| 85.5                      |
| 1.0                       |
| 1.1                       |
| 0.5                       |
| 11.7                      |
| 14.4                      |
| 0.9                       |
| 1.0                       |
| 0.3                       |
| 0.6                       |
| 0.7                       |
| 0.2                       |
| 1.0                       |
| N.D.                      |
| N.D.                      |
| N.D.                      |
| N.D.                      |

From these results, it would be possible that, by mixing lead contaminated soil with sulfur-impregnated product, the eluted solution becomes neutral and the concentration of lead drops below the Japanese eluted standard for soil.

### IV. Conclusions

A novel carbonaceous immobilizing agent for heavy metal contaminated soil can be prepared from rice straw using sulfur immersion and pyrolysis, and can immobilize lead ion in various artificial lead contaminated soils. The product prepared from material immersed in more than 0.5 M K$_2$S solution for less than 15 min has a maximum immobilization ability for lead ion, and the lead immobilization using the sulfur-impregnated product is sustainable due to the formation of leadhillite and anglesite. With addition of the product, the lead elution from all artificial contaminated soil is lower than Japanese standard of soil elution. In future, long term stability of lead immobilization will be examined to use for field verification test.
CONFLICT OF INTEREST
The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS
Please state each author’s contribution to this work, it can be up to several sentences long and should briefly describe the tasks of individual authors. e.g., T. Wajima conducted the research, analyzed the data, and wrote the paper. Author had approved the final version.

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