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1	Short Original Communication
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3	Extraction of Cs bound with biotite by addition of oxalic acid without heating
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25	

27 Abstract

28 Purpose

29	There is a large quantity of contaminated soil with radionuclides due to the accident of Fukushima Daiichi
30	Nuclear Power Plant in Japan. Some previous studies reported that Cs ⁺ could be desorbed from soil with
31	high efficiency, however, these methods required huge input energy for operating. Therefore, we focused
32	on oxalic acid, which has shown relatively high desorption efficiency in previous thermal treatment, and
33	evaluated the potential of oxalic acid as a leaching reagent to reduce the volume of contaminated soil
34	without heating.
35	Methods
36	Stable isotope Cs ⁺ contaminated biotite was retained in plastic bag for 3 months for aging. Oxalic acid was
37	added to the contaminated biotite and the mixture was maintained around 20 °C. After each leaching time,
38	the mixture was separated using filter paper and the concentration of Cs ⁺ in filtrate was measured. Structural
39	changes in biotite before and after leaching were also analyzed. LiNO3 was also used for comparison studies.
40	Results
41	The desorption efficiency of Cs^+ from biotite by leaching with oxalic acid was 94.9% after 2 weeks and the
42	intensity of the specific reflection for biotite in the X-ray diffractograms decreased with leaching time.
43	These results indicate that oxalic acid can remove Cs ⁺ from biotite through the decomposition of the biotite

 $44 \qquad \text{structure. Since LiNO}_3 \text{ desorbed only } 32.2\% \text{ of } Cs^+ \text{ as ion-exchangeable form after } 2 \text{ weeks, oxalic acid}$

45	can	desorb	Cs+,	which	has a	strong	interaction	with	biotite	such	as the	e inner-	sphere	complex,	without
46	heati	ing.													

47Conclusion

- 48 Although the desorption of Cs⁺ from biotite by leaching with oxalic acid leaching takes a long time, it is a
- 49great advantage that this leaching method does not require external energy such as heating. Therefore,
- 50oxalic acid leaching has high potential to reduce the volume of contaminated soil with radionuclides.
- 51
- 52Keywords: Cesium desorption; Biotite; Oxalic acid leaching; Decomposition; Without heating; Volume
- 53reduction
- 54

56 **1** Introduction

57The accident at the Fukushima Daiichi Nuclear Power Plant happened in Japan on March 2011. The 58quantity of contaminated soil was estimated to be 13-14 million m³, of which the clayey soil discharged 59from agricultural fields and forests was estimated to be 6 million m³ (Japan Ministry of the Environment, 2019). The transportation of these soils from the temporary storage site to the interim storage facility started 60 in 2015 (Japan Ministry of the Environment, 2018). It is estimated that even after a ¹³⁷Cs half-life of 30 6162years, the volume of soil above 8,000 Bq/kg will be 105 thousand m³, of which 98 thousand m³ is clayey 63 soil (Japan Ministry of the Environment, 2018). The contaminated soils with less than 8,000 Bq/kg are 64 intended to be recycled as inorganic civil engineering material, while the contaminated soils with more than 658,000 Bq/kg are difficult to utilize as resources. In Japan where there are many natural disasters, the safe 66 storage of the large quantity of contaminated soil for a long time is limited. Therefore, the removal of 67 radioactive Cs from contaminated soil is urgently needed to reduce the volume of contaminated soil. 68However, Cs⁺ forms an inner-sphere complex, which directly coordinates without the intervention of water 69 molecules, with the interlayers of clay minerals, and this interaction is too strong to separate Cs⁺ from clay 70minerals (Mukai et al. 2018). Recently, low-pressure sublimation, thermal, and microwave treatments have 71been developed to desorb Cs⁺ from the contaminated soil (Shimoyama et al. 2014; Parajuli et al. 2015; Yin 72et al. 2017; Tamura et al. 2021; Akemoto et al. 2021a). These methods have high desorption efficiency of Cs⁺ from soil in relatively short time, however, heating at the boiling point of water or more is required. 73

74 Therefore, these method have limits to treat with large quantity of contaminated soil.

75	In this study, we focused on oxalic acid which has relatively high desorption efficiency in previous thermal
76	treatment without a special experimental environment (Kim et al. 2020). The potential of oxalic acid was
77	evaluated by the desorption experiment of Cs ⁺ from model contaminated biotite without heating.
78	
79	2 Materials and methods
80	2.1 Clay mineral and reagents
81	Biotite in Tamura-gun, Fukushima Prefecture, Japan (Geo-Science Materials Nichika Inc., Japan) was used
82	as the model clay mineral. Since the adsorption behavior of $\mathrm{Cs}^{\scriptscriptstyle +}$ on this biotite was established by the
83	Freundlich adsorption isotherm, the interlayer and surface of biotite acted as adsorption sites for Cs ⁺
84	(Akemoto et al. 2021b). For characteristics of biotite used in this study, cation exchange capacity is 380
85	mmol/kg and chemical component is shown in Table S1. From the K ₂ O content in the biotite, the biotite
86	used in our study is suggested to be more weathering and then it may be difficult to desorb Cs^+ from the
87	biotite by the ordinal condition (Kitayama et al. 2020). Cesium chloride (guaranteed reagent grade, >99%)
88	and oxalic acid (guaranteed reagent grade, 98%) were purchased from Fujifilm Wako Pure Chemical
89	Corporation, Japan. Lithium nitrate (guaranteed reagent grade, >98%) was purchased from Kanto Chemical
90	Co., Inc., Japan. All reagents were used without pretreatment.

92 **2.2 Desorption experiment**

93 The Cs⁺ contaminated biotite was prepared as described in a previous study (Akemoto et al. 2021b). The 94particle size of biotite was less than 53 µm and the adsorption capacity of biotite for Cs⁺ was 31.9 mg/g. 95 Biotite that adsorbed Cs⁺ was retained in plastic bag at room temperature for 3 months before experiments. 96 Oxalic acid (0.5 mol/L, 25 mL) was added to the contaminated biotite (0.05 g) and the mixture was 97maintained without any mechanical shaking under room temperature, which was measured continuously 98during the experiment (Thermo Recorder, TR-71U, T&D Corporation, Japan). After each leaching time (1 99 hour, 1 day, 3 days, 1 week, 2 weeks, 3 weeks, and 4 weeks), the mixture was separated using a filter paper 100(5C, 1 µm pore size, Toyo Roshi Kaisha Ltd., Japan), and the concentration of Cs⁺ in the filtrate was 101measured using a flame atomic absorption spectrophotometer (FAAS, ZA-3300, Hitachi Ltd., Japan) at 102 852.1 nm. Oxalic acid at low concentration (0.1 mol/L) and LiNO₃ (1.0 mol/L) were also used for 103 comparison studies for their effect on extraction of Cs⁺ with the same methodology as mentioned above. 104Since Li^+ has a high hydration energy, it is expected to desorb Cs^+ by expansion of the clay mineral 105interlayer due to the penetration of Li⁺ in it (Mukai et al. 2018). The desorption efficiency of Cs⁺ from 106 biotite was calculated by the following equation (Eq. 1). Desorption efficiency (%) = $\frac{C_t \times V}{Q \times W} \times 100$ 107(1)

108 where C_t is the concentration of Cs⁺ (mg/L) in the solution after each leaching time t, V is the volume of 109 the extract solution (L), Q is the adsorption capacity of biotite for Cs⁺ (31.9 mg/g), and W is the amount of 110 biotite (g). All experiments were performed in triplicates. The main elements of the clay minerals (Al, Fe, 111 Mg, and Si) in the filtrate were measured using inductively coupled plasma-atomic emission spectroscopy 112(ICP-AES, ICPS-8100, Shimadzu Corporation, Japan). The percentage of these elements (Al, Fe, Mg, and 113Si) extracted from the initial biotite was calculated based on the results of energy-dispersive X-ray 114fluorescence spectrometry (XRF, JSX-3100RII, JEOL Ltd., Japan) (Akemoto et al. 2021b). The solution 115pH was measured using a pH meter (D-51, Horiba Ltd., Japan). The biotite before and after leaching 116 treatment was measured using powder X-ray diffraction (XRD, SmartLab, Rigaku Corporation, Japan). 117XRD was performed using Cu K α radiation ($\lambda = 0.154$ nm) at 40 kV and 30 mA at a speed of 1 °/min and 118 an angular step of 0.01° .

119

120 **3** Results and discussion

The room temperature was observed to be at 20 ± 5 °C during the 4 weeks experiment. The pH of the solutions were 5.9, 1.3 and 0.8 in 1.0 mol/L LiNO₃, 0.1 mol/L oxalic acid and 0.5 mol/L oxalic acid, respectively, and these values were consistent throughout the experiment. The Cs⁺ adsorbed on clay minerals is considered to mainly take two forms (Fan et al. 2014). One is the outer-sphere complex on the interlayer or planar site, and this form is relatively easy to desorb by alkali metal ions. The other cannot be desorbed easily due to the formation of the inner-sphere complex on the interlayer or frayed edge site. The Cs⁺, which was not desorbed by Li⁺, was considered to exist as an inner-sphere complex in biotite.

128Figure 1 shows the desorption efficiency of Cs^+ from biotite at each leaching time. LiNO₃ could desorb 129approximately 31.6% of Cs⁺ after 1 h of leaching time, and the desorption efficiency was consistent 130throughout the experiment. According to a previous study, approximately 40% of Cs⁺ in biotite is existed 131as an ion-exchangeable form (Akemoto et al. 2021b). In this study, Li⁺ desorbed the ion-exchangeable form 132of Cs⁺ from biotite and reached equilibrium within 1 h of the leaching time. 133The desorption efficiency by oxalic acid increased with leaching time, and the value of 85.3% (0.1 mol/L) 134and 94.9% (0.5 mol/L) were obtained after 2 weeks, and 89.8% (0.1 mol/L) and 96.1% (0.5 mol/L) were 135obtained after 4 weeks of the experiment. These efficiencies were higher than that of Li⁺. The Cs⁺ has been 136considered to move into the interlayer of clay minerals and retain in an energy-stable site (Mukai et al. 1372018). Since this study used biotite that has been saturated with Cs^+ and aged for 3 months, Cs^+ could have 138occupied the energy-stable sites that were inaccessible. Therefore, the high efficiency of oxalic acid can be 139attributed to the ability of oxalic acid to desorb Cs⁺ in both outer-sphere complex and inner-sphere complex 140with biotite. 141Al, Mg, Fe, and Si are the main components of clay minerals, and the concentrations of each element in 142the extract solution was measured as a function of leaching time. When LiNO₃ was used as an extraction 143solution, Al, Mg, Fe, and Si was hardly observed in the solution (Fig. S1). Figure 3a indicates X-ray 144diffractograms of biotite after each leaching time by LiNO3. The basal spacing of biotite after Cs⁺ was 1.0

145 nm ($2\theta = 8.7^{\circ}$), indicating that Cs⁺ was still adsorbed in the interlayer (Kogure et al. 2012). The intensity

146	of this reflection (8.7°) was consistent even after 4 weeks, the biotite structure was unaltered. This result
147	also indicated that $LiNO_3$ desorbed only Cs^+ in an ion-exchangeable form. In contrast, oxalic acid desorbed
148	Al, Mg, Fe, and Si in the biotite. The desorption rates of Al, Mg, and Fe were 68%, 83%, and 94% by 0.5
149	mol/L oxalic acid after 2 weeks, and this desorption rate did not significantly change until 4 weeks (Fig. 2).
150	Since the desorption efficiency of Cs^+ by 0.5 mol/L oxalic acid also plateaued at a leaching time of 2 weeks
151	(Fig. 1), these results indicate that the decomposition of clay minerals by oxalic acid (0.5 mol/L) induced
152	the desorption of Cs^+ from biotite at room temperature. When oxalic acid was 0.1 mol/L, the concentrations
153	of Al, Mg, and Fe continued to increase even after 4 weeks (Fig. S1). At 4 weeks of leaching time, the
154	desorption rates of Al, Mg, and Fe were 60%, 78%, and 91%, respectively.
155	Buckwheat has been reported to detoxify Al with oxalic acid released from its root, which forms an oxalate
156	complex, with Al to oxalic acid ratio of 1:3 (Ma et al. 1997). The reaction ratio between Fe and oxalic acid
157	was 2:5 (Lee et al. 2006). The concentration of Al and Fe in biotite used in this study was 3.2 mmol/g and
158	2.0 mmol/g, respectively. Therefore, even 0.1 mol/L of oxalic acid was sufficient to react with all Al and
159	Fe to form their complexes with oxalic acid. The XRD results indicated that the intensity of the biotite
160	reflection decreased with leaching time (Figs. 3b and 3c), although this biotite reflection (1.0 nm) has not
161	completely disappeared with 0.1 mol/L of oxalic acid, even at a leaching time of 4 weeks (Fig. 3b). This
162	result implied that the decomposition of biotite by 0.1 mol/L oxalic acid takes long time over 4 weeks. On
163	the other hand, it was almost completely disappeared with 0.5 mol/L oxalic acid after 2 weeks (Fig. 3c).

164	The broad peak around 11.5 ° was estimated to be oriented from a hydrotalcite-like structure, which has
165	XRD reflection at 11.7 ° and is similar to octahedral sheet (Miyata 1983; Bernard et al. 2022). Assuming
166	that Al and Fe eluted by oxalic acid leaching are metals that have existed in the tetrahedral sheet by
167	isomorphous substitute with Si, it is considered that the tetrahedral sheet could not maintain its structure
168	due to this elution. Since this reflection appears with the decrease in biotite reflection, it is presumed to be
169	from the remaining octahedral sheet. These results indicated that oxalic acid (0.5 mol/L) could desorb
170	elements (Al, Mg, Fe, and Si) in biotite and consequently decompose its structure. The The Cs ⁺ desorption
171	from biotite due to decomposition of its structure by oxalic acid can be performed even without heating.
172	Oxalic acid is a natural organic acid present in various plants and is highly effective in the field of
173	environmental remediation (Huang et al. 2021; Costa et al. 2022). Since the results of this study confirmed
174	that Cs ⁺ strongly bound in clay minerals can be desorbed with oxalic acid without heating, there is the
175	possibility of introducing plants with oxalic acid content as a soil remediation technology option. Moreover,
176	this oxalic acid leaching has the potential to be a low energy soil remediation as it does not require heating.
177	

178Conclusions 4

179In this study, the Cs⁺ desorption from biotite using oxalic acid was investigated without heating. Oxalic 180acid (0.5 mol/L) could desorb Cs^+ from biotite which has been aged for 3 months, and its desorption 181 efficiency was 94.9% after 2 weeks. In the same procedure, LiNO3 could desorb only 32.2% of Cs⁺ after 2

182	weeks. These results indicate the ability of oxalic acid to desorb Cs^+ from biotite not only the outer-sphere
183	complex like ion-exchangeable form but also the inner-sphere complex. Although the desorption of Cs^+
184	from biotite by oxalic acid leaching takes a long time (at least 2 weeks), the fact that no external energy is
185	required for extraction is a great advantage. Therefore, oxalic acid leaching without heating can be used for
186	the volume reduction of contaminated soil with radionuclides.

187

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194	Author contribution
195	Yasuhiro Akemoto: Conceptualization, Methodology, Formal analysis and investigation, Writing - original
196	draft preparation, Visualization
197	Seira Takahashi: Methodology, Investigation
198	Toko Iwamura: Methodology, Investigation
199	Masahiko Kan: Writing - review and editing
200	Shunitz Tanaka: Methodology, Writing - review and editing, Supervision
201	
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206	The datasets used and/or analyzed during the current study are available from the corresponding author on
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208	
209	
210	Declarations
211	Competing Interests
212	The authors declare that they have no known competing financial interest or personal relationship that could
213	have appeared to influence the work reported in this paper.
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275	Figure captions
276	
277	Fig. 1 Desorption efficiency of Cs ⁺ from biotite at each leaching time at 20 \pm 5 °C. Biotite: 0.05 g, the
278	leaching solution: 25 mL. Error bars indicate standard deviation $(n = 3)$
279	
280	Fig. 2 Desorption rate of elements at each leaching time from biotite by 0.5 mol/L oxalic acid; Symbols,
281	purple triangle was Fe, blue circle was Mg, red square was Al, and green rhombus was Si, respectively
282	
283	Fig. 3 XRD patterns of biotite after each leaching treatment, a) 1.0 mol/L LiNO3, b) 0.1 mol/L oxalic acid,
284	c) 0.5 mol/L oxalic acid
285	
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287	



Fig. 1 Desorption efficiency of Cs^+ from biotite at each leaching time at 20 ± 5 °C. Biotite: 0.05 g, the leaching solution: 25 mL. Error bars indicate standard deviation (n = 3)



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299 Fig. 3 XRD patterns of biotite after each leaching treatment, a) 1.0 mol/L LiNO₃, b) 0.1 mol/L oxalic acid,

- 300 c) 0.5 mol/L oxalic acid