


 Cite this: *RSC Adv.*, 2023, **13**, 17001

Extraction of Se(IV) and Se(VI) from aqueous HCl solution by using a diamide-containing tertiary amine†

 Hirokazu Narita, ^a Motoki Maeda, ^b Chiharu Tokoro, ^b Tomoya Suzuki, ^a Mikiya Tanaka, ^c Hideaki Shiwaku ^d and Tsuyoshi Yaita ^d

Here, we investigated the mechanism underlying the extraction of Se(IV) and Se(VI) from aqueous HCl solutions by *N*-2-ethylhexyl-bis(*N*-di-2-ethylhexyl-ethylamide)amine (EHBAA). In addition to examining extraction behavior, we also elucidated structural properties of the dominant Se species in solution. Two types of aqueous HCl solutions were prepared by dissolving a Se^{IV} oxide or a Se^{VI} salt. X-ray absorption near edge structure analyses revealed that Se(VI) was reduced to Se(IV) in 8 M HCl. Using 0.5 M EHBAA, ~50% of Se(VI) was extracted from 0.5 M HCl. In contrast, Se(IV) was hardly extracted from 0.5 to 5 M HCl; however, at molar concentrations above 5 M, the extraction efficiency of Se(IV) increased drastically, reaching ~85%. Slope analyses for the distribution ratios of Se(IV) in 8 M HCl and Se(VI) in 0.5 M HCl showed that apparent stoichiometries of Se(IV) or Se(VI) to EHBAA were 1:1 and 1:2, respectively. Extended X-ray absorption fine structure measurements revealed that the inner-sphere of the Se(IV) and Se(VI) complexes extracted with EHBAA was [SeOCl₂] and [SeO₄]²⁻, respectively. Together, these results indicate that Se(IV) is extracted from 8 M HCl with EHBAA via a solvation-type reaction, whereas Se(VI) is extracted from 0.5 M HCl via an anion-exchange-type reaction.

 Received 28th February 2023
 Accepted 1st May 2023

 DOI: 10.1039/d3ra01341c
rsc.li/rsc-advances

1. Introduction

Selenium (Se) has been widely used in various industrial fields.^{1,2} Industrial production of Se is mainly achieved by recovery as a byproduct from the anode slimes produced during copper smelting.^{3,4} Solvent extraction is one of the important methods for the recovery of Se; however, our understanding of the extraction of Se ions from acidic solutions and the precise speciation of Se ions in relatively concentrated HCl solutions remains largely incomplete. Se is also an essential trace element for humans and animals; however, its consumption in large amounts is detrimental to health. Therefore, there have been many studies conducted to determine the concentration and oxidation states of Se in the natural environment and to examine how best to facilitate its removal.⁵⁻⁷ Thus, more precise

speciation data will facilitate the development of more efficient solvent extraction methods for Se.

Many microextraction approaches have been reported for the solvent extraction of trace amounts of Se from media such as food and environmental water.⁷ However, approaches for the extraction of Se from highly acidic solutions are limited. Tri-*n*-butyl phosphate (TBP) has been successfully used for solvent extraction of Se(IV) from HCl solution.⁸⁻¹⁰ Interestingly, Chowdhury and Sanyal⁹ reported that the dominant complex extracted changes depending on the concentration of the HCl solution, with SeOCl₂(TBP)₂ being extracted from 3 to 6 M HCl and H₂SeOCl₄(TBP)₂ being extracted from 6 to 10 M HCl. A TBP analog, tri-*iso*-butyl phosphate, has also been shown to extract Se(IV) from chloride aqueous solution at around pH 8.5.¹¹ A protonated *N*-*n*-octylaniline has been shown to form an ion-pair complex with [HSeO₃]⁻.¹² In contrast, diammmoniumcalix[4]arene has been shown to extract Se(VI) via ion-pair formation with [HSeO₄]⁻ or [SeO₄]²⁻.^{13,14} The reactions of these extractants with Se were estimated mainly by slope analyses of the distribution ratio of Se between the organic and aqueous phases. Moving forward, direct structural information of the dominant Se species in the organic and aqueous phases is needed to understand the extraction behavior in more detail.

In acidic chloride solutions, Se(IV) and Se(VI) form selenous acid (H₂SeO₃) and selenic acid (H₂SeO₄), respectively, and dissociate to anions such as HSeO₃⁻, SeO₃²⁻, HSeO₄⁻, and

^aGlobal Zero Emission Research Center, National Institute of Advanced Industrial Science and Technology (AIST), 16-1 Onogawa, Tsukuba, Ibaraki 305-8569, Japan.
 E-mail: hirokazu-narita@aist.go.jp

^bSchool of Creative Science and Engineering, Waseda University, 3-4-1 Okubo, Shinjuku, Tokyo 169-8555, Japan

^cNational Institute of Advanced Industrial Science and Technology (AIST), 1-1-1 Umezono, Tsukuba, Ibaraki 305-8560, Japan

^dMaterials Science Research Center, Japan Atomic Energy Agency (JAEA), 1-1-1 Koto, Sayo, Hyogo 679-5148, Japan

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d3ra01341c>



SeO_4^{2-} with a decrease in the acid concentration.¹⁵ For Se(iv), it has been suggested that an SeOCl_2 complex forms in highly concentrated HCl solutions (4–12 M), and at even higher concentrations (~ 15 M) SeCl_5^- is produced.¹⁶ Compared with the number of reports providing structural information for some precious metals and base metals, which are present in the anode slimes, there are fewer reports providing structural information about Se ions in acidic chloride solutions. Knowledge of the structure of Se ions in the aqueous phase is needed for the development of new extractants because extraction type is highly dependent on the dominant Se complex in solution.

Our group has been examining the possibility of using amide-containing tertiary amines (ACTAs) as extractants of metal ions.^{17–22} ACTAs efficiently extract chlorometalates *via* an anion-exchange reaction because of their stable protonation ability.²¹ They also extract rare earth elements from HNO_3 *via* a solvation reaction.²² In particular, ACTAs have excellent extraction efficiency for platinum-group metals from HCl solutions. For example, diamide- and triamide-type ACTAs are able to extract $[\text{RhCl}_5(\text{H}_2\text{O})]^{2-}$, an aqua-chloro complex, despite this complex generally being considered unextractable.^{18,21} ACTAs are also much more efficient extractants of metal complex anions than the typical tertiary amine extractant tri-*n*-octylamine (TOA). We therefore expected that ACTAs could be used to efficiently extract Se from HCl.

Here, we examined the extraction of Se(iv) and Se(vi) from aqueous HCl solutions by using a diamide-containing tertiary amine, *N*-2-ethylhexyl-bis(*N*-di-2-ethylhexyl-ethylamide)amine (EHBA) (Fig. 1). We discuss the underlying extraction mechanism based on a slope analysis of the distribution ratio of Se ions between the organic and aqueous phases and structural data for the dominant Se complexes in the solutions obtained by X-ray absorption fine structure (XAFS) analysis. To allow comparison, the efficacies of the extractants TOA and TBP were also examined.

2. Experimental

2.1. Reagents

SeO_2 (FUJIFILM Wako Pure Chemical Co.) and $\text{Na}_2\text{SeO}_4 \cdot 10\text{H}_2\text{O}$ (Alfa Aesar) were used as an Se(iv) oxide and Se(vi) salt, respectively. EHBA (purity: 96%) was provided by Chemicrea Inc. TBP and TOA were purchased from the Tokyo Chemical Industry Co.

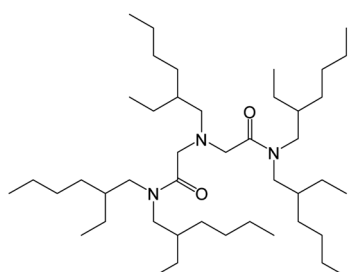


Fig. 1 Structure of *N*-2-ethylhexyl-bis(*N*-di-2-ethylhexyl-ethylamide)amine.

The extractants were used without further purification. All the other chemicals used were of reagent grade.

2.2. Solutions of Se in aqueous HCl

Aqueous solutions of Se ions were prepared by dissolving SeO_2 or $\text{Na}_2\text{SeO}_4 \cdot 10\text{H}_2\text{O}$ in aqueous HCl. Hereafter, the aqueous solutions are abbreviated as SolAxH and SolBxH, where 'A' = SeO_2 , 'B' = $\text{Na}_2\text{SeO}_4 \cdot 10\text{H}_2\text{O}$, and 'x' = HCl concentration (M). The Se concentration was 0.001 M for the solvent extraction experiments and 0.1 M for the XAFS measurements. For the solvent extraction experiments, the aqueous solutions were used at 12 h after oxide or salt dissolution. For the XAFS measurements, mainly the aqueous solutions were used at 3 w after oxide or salt dissolution.

2.3. Solvent extraction

An 80/20 (v/v) *n*-dodecane/2-ethylhexanol mixture was used as the diluent for the extractants. Equal volumes of one of the diluted extractants (EHBA, TOA, or TBP) and of Se-free HCl solution were added to a 10 mL glass tube, and the tube was shaken vertically for 30 min at an amplitude of 100 mm and a frequency of 200 stroke per min and then centrifuged. Next 1 mL of the HCl-equilibrated organic phase and the same volume of HCl solution containing 0.001 M Se (SolAxH or SolBxH) was similarly shaken for 60 min and then centrifuged. The shaking time was sufficient to ensure that the extraction equilibrium was reached. The concentrations of Se ions in the aqueous phase were measured by ICP-AES (Horiba ULTIMA2). The concentrations in the organic phase were calculated from the mass balance of the metal ions before and after extraction. The data accuracy was within 5%. The extractions were performed at room temperature (23 ± 2 °C). The extraction percentage ($E\%$) and distribution ratio (D) were calculated by using eqn (1) and (2), respectively, where $[\text{Se}]_{\text{init},\text{aq}}$, $[\text{Se}]_{\text{eq},\text{org}}$, and $[\text{Se}]_{\text{eq},\text{aq}}$ denote the Se concentrations in the initial aqueous phase, the equilibrated organic phase after extraction, and the equilibrated aqueous phase after extraction, respectively.

$$E\% = \frac{[\text{Se}]_{\text{eq},\text{org}}/[\text{Se}]_{\text{init},\text{aq}} \times 100}{\{([\text{Se}]_{\text{init},\text{aq}} - [\text{Se}]_{\text{eq},\text{aq}})/[\text{Se}]_{\text{init},\text{aq}}\}} \times 100 \quad (1)$$

$$D = [\text{Se}]_{\text{eq},\text{org}}/[\text{Se}]_{\text{eq},\text{aq}} = \frac{([\text{Se}]_{\text{init},\text{aq}} - [\text{Se}]_{\text{eq},\text{aq}})/[\text{Se}]_{\text{eq},\text{aq}}}{[\text{Se}]_{\text{eq},\text{aq}}} \quad (2)$$

2.4. XAFS measurements

Four aqueous solutions, SolA0.5H, SolA8H, SolB0.5H, and SolB8H ([Se] = 0.1 M), were prepared. For SolB8H, the dependence on standing time was investigated because a change in the oxidation state was suggested during the solvent extraction experiment. Organic solutions containing Se complexes extracted with EHBA were prepared by the solvent extraction method detailed in Section 2.3. SolA8H and SolB0.5H ([Se] = 0.1 M) and 0.5 M EHBA were used for the aqueous and organic phases, respectively. The concentration of the Se complexes extracted with EHBA from SolA8H and SolB0.5H (hereafter



abbreviated to ExSolA and ExSolB, respectively) was 0.08 M for ExSolA and 0.05 M for ExSolB. The solutions were transferred to 10 mm diameter polyethylene cells, which were then tightly sealed. Solid reference samples were prepared by grinding the Se oxide or salt with boron nitrate and then pelleting the resulting powder.

XAFS measurements were performed at the BL11XU beamline of SPring-8 (8 GeV at ~99 mA). Synchrotron X-ray radiation was monochromatized using Si(111) crystal monochromators. Se K-edge XAFS spectra were collected in transmission mode using two ion chambers (both filled with N₂) at room temperature (~25 °C). The beam size on the sample was about 1 × 1 mm². The SeO₂ was used for energy calibration prior to X-ray absorption near edge structure (XANES) measurements; the peak top of the Se K-edge (white line) was aligned to 12 664 eV.²³ XAFS data were collected three times and then averaged for each sample.

Data processing was performed with the EXAFS data analysis software package WinXAS version 4.0.²⁴ Theoretical phases and amplitudes were calculated by using the software FEFF 8.²⁵ For the FEFF calculations, the crystal structures of (Se^{IV}O₃)²⁻ in [Na₂SeO₃],²⁶ (Se^{VI}O₄)²⁻ in [(NH₄)₄H₂(SeO₄)₃],²⁷ and (Se^{IV}OCl₂) in [SbCl₅·SeOCl₂]²⁸ were used as model compounds. Multiple scattering paths were ignored because their contribution was not significant for the studied systems. Thus, we considered only single scattering paths: Se–O and/or Se–Cl. The *k*-range for the Fourier transforms was 4–12 Å⁻¹ for SolA0.5H and 4–13 Å⁻¹ for the others, and the curve-fitting *R*-range was 0.5–2.5 Å for SolA8H and ExSolA, and 0.8–2.0 Å for the others. The coordination geometry of Se in 0.5 M HCl solution (SolA0.5H) is tetragonal SeO₃ because SeO₃²⁻, HSeO₄⁻, and/or H₂SeO₄ are present.¹⁵ Therefore, the coordination number was kept constant at 3 and the amplitude reduction factor, *S*₀², was refined; the obtained *S*₀² value (0.86(3)) was used for all fits. This value was considered reasonable because the obtained number of correlations (coordination number), *N*, for Se–O for the solid reference samples, Se^{IV}O₂ and Na₂Se^{VI}O₄·10H₂O, was 3.1(3) and 4.0(4), respectively (Fig. S3 and Table S1†). The structural parameters (*N*; bond distance, *r* (Å); Debye–Waller factor squared, σ^2 (Å²); and the shift in threshold energy, ΔE_0 (eV)) were allowed to vary in the fit; the same ΔE_0 value was used for each shell. The difference in bond lengths between Se–O and Se–Cl was distinguishable because their difference is more than the smallest separation,²⁹ given by $\Delta r = \pi/2\Delta k$ ($\Delta k \approx 9$ Å⁻¹, $\Delta r \approx 0.17$ Å). The fitting quality was checked by using the *R*-factor: $\{\sum |k^3 \chi(k)_{\text{obs}} - k^3 \chi(k)_{\text{calc}}| / \sum |k^3 \chi(k)_{\text{obs}}|\} \times 100$.

3. Results and discussion

3.1. Effect of HCl concentration on Se ion extraction

The extraction percentage of Se as a function of HCl concentration with 0.5 M extractants was determined for SolAxH and SolBxH. Three extractants were examined: EHBAA, TOA, and TBP. Third phase formation was observed at 0.5 M and 8 M HCl in the TOA system. In the SolAxH system, very little Se was extracted at HCl concentrations below 5 M; however, the extraction percentage rapidly increased at HCl concentrations

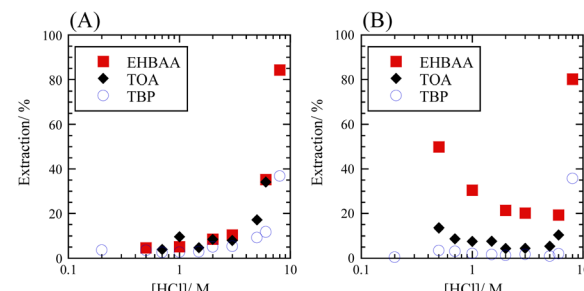


Fig. 2 Extraction percentage of Se as a function of HCl concentration. (A) SolAxH: Se^{IV}O₂ dissolved and (B) SolBxH: Na₂Se^{VI}O₄·10H₂O dissolved. Organic phase, 0.5 M EHBAA, TOA, or TBP pre-equilibrated with HCl; aqueous phase, [Se] = 0.001 M.

above 5 M for each of the extractants (Fig. 2A). In the SolBxH system, similar rapid increases were observed at HCl concentrations above 5 M for each of the extractants (Fig. 2B); however, extraction percentages up to 50% were obtained at HCl concentrations below 5 M, depending on the extractant. The extraction efficiency in this region decreased in the order of EHBAA ≫ TOA > TBP, indicating that anion-exchange extraction predominantly occurred at concentrations of 0.5–5 M because TBP is a weak anion exchanger,³⁰ and EHBAA is a more efficient extractant of metal complex anions than TOA.^{17–19,21} According to a previous study examining the reduction of Se(vi), Se(vi) is rapidly reduced to Se(iv) by chloride in a strongly acidic medium;³¹ a similar reduction of Se(vi) to Se(iv) was indicated to have occurred at HCl concentrations above 5 M in the SolBxH system.

3.2. Dominant Se species in aqueous HCl solution

Fig. 3 shows the Se K-edge XANES spectra obtained for SolA0.5H, SolA8H, SolB0.5H, and SolB8H, together with the

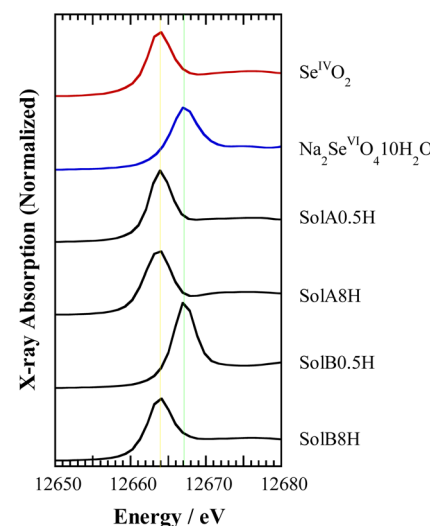


Fig. 3 Se K-edge XANES spectra for the aqueous solutions SolA0.5H, SolA8H, SolB0.5H, and SolB8H and the reference solids Se^{IV}O₂ and Na₂Se^{VI}O₄·10H₂O.



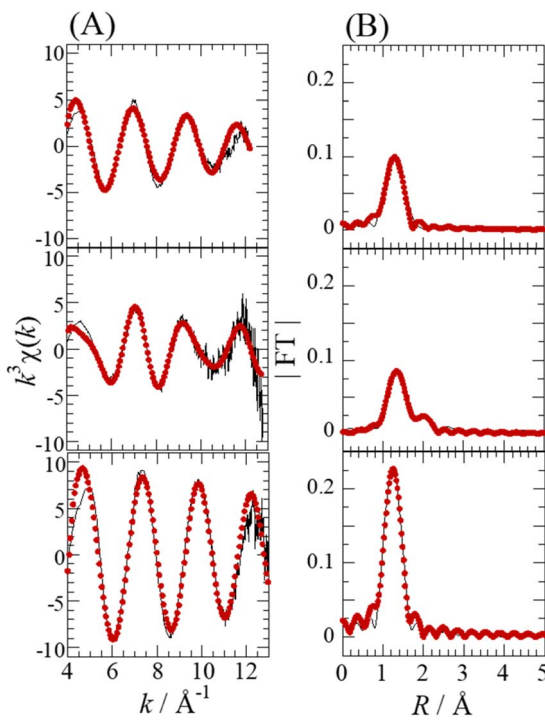


Fig. 4 Se K-edge k^3 -weighted EXAFS spectra (A) and the corresponding Fourier transforms (FT) (B) for SolA0.5H (top), SolA8H (middle), and SolB0.5H (bottom). The phase shifts are not corrected. Experimental data (solid line) and theoretical fit (dotted line).

spectra for solid samples of $\text{Se}^{\text{IV}}\text{O}_2$ and $\text{Na}_2\text{Se}^{\text{VI}}\text{O}_4 \cdot 10\text{H}_2\text{O}$ for reference. The peak positions in the spectra for SolA0.5H and SolA8H were in good agreement with that of the solid $\text{Se}^{\text{IV}}\text{O}_2$ sample. In contrast, although the peak position in the spectrum for SolB0.5H was in good agreement with that of the solid $\text{Na}_2\text{Se}^{\text{VI}}\text{O}_4 \cdot 10\text{H}_2\text{O}$ sample, the peak position for SolB8H was in agreement with that of $\text{Se}^{\text{IV}}\text{O}_2$, indicating that Se(vi) in 8 M HCl was reduced to Se(iv) after dissolution. This is consistent with

the findings from the extraction experiment discussed in Section 3.1.

The time for this reduction to occur was investigated by XANES measurement using SolB8H solutions that were left to stand before analysis (Fig. S1†). The spectra show that Se(vi) was reduced to Se(iv) within 12 h after dissolution.

Fig. 4 shows the Se K-edge k^3 -weighted EXAFS spectra and corresponding Fourier transforms for SolA0.5H, SolA8H, and SolB0.5H. An intense peak seen at ~ 1.3 Å in each spectrum (Fig. 4B) was assigned to the Se–O correlation. For SolA8H, in addition to the intense peak, there was a small peak at ~ 2 Å, which was assigned to the Se–Cl correlation. The curve-fitting results are shown in Table 1. According to the $\text{p}K_{\text{a}_1}$ value of selenous acid (~ 2.5),¹⁵ H_2SeO_3 is predominant in 0.5 M HCl solution. In contrast, in >7 M HCl solution both $[\text{SeOCl}_2]$ and $[\text{H}_2\text{SeO}_3]$ are present, and in 15 M HCl solution SeCl_5^- is dominant.¹⁶ For SolA0.5H, both the N value (3) and Se–O bond length ($1.71(1)$ Å) were almost the same as those for the crystal structure of $[\text{SeO}_3]^-$.²⁶ The small peak at ~ 2 Å for SolA8H was confirmed to represent the Se–Cl correlation by the obtained structural parameters. The obtained N values for Se–O and Se–Cl were 1.8(1) and 1.1(1), which although not consistent with the structure of the $[\text{SeOCl}_2]$ complex, are consistent with a mixture of $[\text{SeOCl}_2]$ and $[\text{H}_2\text{SeO}_3]$. Although Raman studies have inferred the existence of a $[\text{SeO}_2\text{Cl}]^-$ complex anion,³² that study conducted measurements not in HCl aqueous solution but in solid and acetonitrile. For SolB0.5H, the curve-fitting results (Se–O: $N = 4.4(4)$, $r = 1.633(4)$ Å) indicated that the inner-sphere structure is tetrahedral SeO_4 , which is consistent with the XANES spectrum. According to the $\text{p}K_{\text{a}_2}$ of selenic acid (1.6–2.0),¹⁵ both $[\text{SeO}_4]^{2-}$ and $[\text{HSeO}_4]^-$ would exist in low-pH solutions.

3.3. Structure of the Se complexes extracted with EHBAA

The Se complexes extracted with EHBAA from SolA8H and SolB0.5H (abbreviated as ExSolA and ExSolB, respectively) were

Table 1 Curve-fitting results for selenium EXAFS data

		N^a	r^b (Å)	$\sigma^2 c$ (Å ²)	ΔE^d (eV)	R -factor ^e
SolA0.5H	Se–O	3 ^f	1.71(1)	0.0048(9)	9(1)	18
SolA8H	Se–O	1.8(1)	1.70(1)	0.0016(5)	5.8(5)	7.9
	Se–Cl	1.1(1)	2.29(1)	0.0065(6)		
SolB0.5H	Se–O	4.4(4)	1.633(4)	0.0014(6)	8(1)	13
ExSolA	Se–O	1.0(2)	1.64(2)	0.002(1)	5(1)	14
	Se–Cl	2.0(3)	2.28(2)	0.006 (1)		
ExSolB	Se–O	3.8(3)	1.632(4)	0.0018(5)	5(1)	8.5
Crystal structures						
$[\text{Na}_2\text{SeO}_3]^g$	Se–O	3	1.71 ^h			
$[\text{SbCl}_5 \cdot \text{SeOCl}_2]^i$	Se–O	1	1.69			
	Se–Cl	2	2.12 ^h			
$[\text{TiCl}_3(\text{SeOCl}_2)_2]_2(\mu\text{-O})^j$	Se–O	1	1.644			
	Se–Cl	2	2.145 ^h			
$[(\text{NH}_4)_4\text{H}_2(\text{SeO}_4)_3]^k$	Se–O	4	1.64 ^h			

^a Coordination number. ^b Bond distance (Å). ^c Debye–Waller factor squared (Å²). ^d The shift in threshold energy (eV). ^e Residual: R -factor = $\sum |k^3 \chi(k)_{\text{obs}} - k^3 \chi(k)_{\text{calc}}| / \sum |k^3 \chi(k)_{\text{obs}}| \times 100$. ^f Fixed parameter. ^g Ref. 26. ^h Averaged bond distances. ⁱ Ref. 28. ^j Ref. 33. ^k Ref. 27. Estimated errors are shown in parentheses.



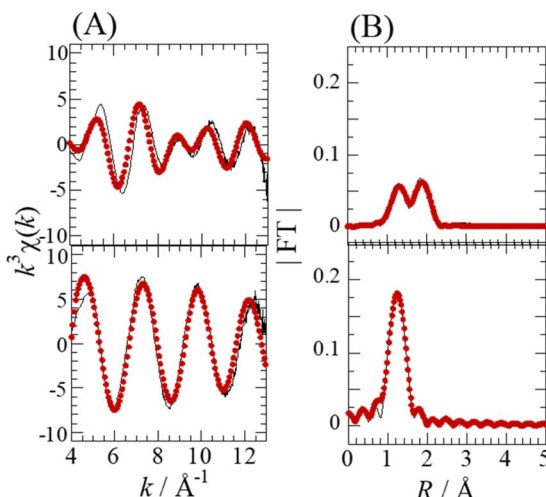
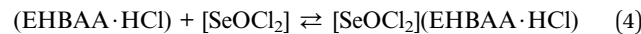


Fig. 5 Se K-edge k^3 -weighted EXAFS spectra (A) and the corresponding Fourier transforms (FT) (B) for ExSolA (top), and ExSolB (bottom). $[\text{Se}] = 0.08 \text{ M}$ in ExSolA and 0.05 M in ExSolB. The phase shifts are not corrected. Experimental data (solid line) and theoretical fit (dotted line).

examined by XAFS. The XANES spectra obtained for these extracts indicated that the oxidation state of Se is retained after being extracted into the organic phase (Fig. S2†). The Se K-edge k^3 -weighted EXAFS spectra and the corresponding Fourier transforms for ExSolA and ExSolB are shown in Fig. 5. In the Fourier transform for ExSolA, two intense peaks were seen at ~ 1.3 and $\sim 2 \text{ \AA}$. The $\sim 2 \text{ \AA}$ peak was more intense than that observed in the spectrum for the aqueous phase (*i.e.*, in the spectrum for SolA8H). The ratio of $N(\text{Se}-\text{O})$ to $N(\text{Se}-\text{Cl})$ for ExSolA was about 1:2, which is different from that for SolA8H (2:1). In the reported crystal structures of $[\text{SeOCl}_2]$, the bond lengths of Se-O (1.644–1.69 \AA) and Se-Cl (2.12–2.145 \AA) vary slightly.^{28,32} The parameter of Se-Cl obtained in the present study is longer than these values, suggesting that the slight structural change of the SeOCl_2 moiety in solution. In contrast, the XANES and EXAFS spectra for ExSolB are very similar to those for SolB0.5H, which is consistent with the obtained structural parameters indicating the presence of SeO_4 , as was observed for SolB0.5H.

3.4. Extraction mechanism of Se(IV) and Se(VI)

Fig. 6 shows the dependence of the distribution ratio of Se on the EHBAA concentration for SolA8H and SolB0.5H. The slope values are different between SolA8H and SolB0.5H, indicating that 1:1 Se(IV):EHBAA and 1:2 Se(VI):EHBAA complexes are formed, respectively. Since EHBAA is readily protonated as contacting with HCl solution,^{19,21} ion-pair formation with a complex anion preferentially occurs. Also, neutral complexes may be extracted by EHBAA *via* a solvation reaction.²² For the EHBAA-SolA8H system, the EXAFS results suggested that a SeOCl_2 moiety is dominant in the organic phase. Therefore, the extraction reaction can be presented as



The SolA8H solution probably contains neutral $[\text{H}_2\text{SeO}_4]$ and $[\text{SeOCl}_2]$, as mentioned in Section 3.2. The very low extraction percentage of Se in the HCl concentration range of 0.5–5 M, indicates that $[\text{H}_2\text{SeO}_4]$, which was the dominant species at those concentrations, was unextractable. Although $[\text{SeOCl}_2]$ is not anionic, a higher reactivity can be expected than that of $[\text{H}_2\text{SeO}_4]$.¹ A previous study using TBP also showed that TBP did not extract $[\text{H}_2\text{SeO}_4]$, but instead it extracted $[\text{SeOCl}_2]$ and $[\text{H}_2\text{SeOCl}_4]$.⁹

Regarding the protonated structure of EHBAA,²¹ an H^+ is chelated by two amide oxygen atoms and an amine nitrogen atom, and a Cl^- weakly bonds to an array of polarized C-H^{δ+} units on the opposite side of EHBAA from the protonation site. Considering the stable protonation ability of EHBAA, solvation with $[\text{SeOCl}_2]$ possibly occurs at the site opposite the diamide oxygen atoms. The crystal structure of $[\text{TiCl}_3(\text{SeOCl}_2)_2]_2(\mu\text{O})\cdot\text{CH}_2\text{Cl}_2$ has three long-distance Se···Cl interactions (2.930–3.465 \AA) and two direct Se-Cl bonds (avg. 2145 \AA).³³ The Cl^- in the EHBAA·HCl complex might have a weak interaction with $[\text{SeOCl}_2]$ at the outer-sphere of the Se, although the interaction was too weak to produce a peak in the EXAFS spectrum.

For the EHBAA-SolB0.5H system, the EXAFS results show that SeO_4 was dominant in both the organic and aqueous phases. In the aqueous phase, both $[\text{HSeO}_4]^-$ and $[\text{SeO}_4]^{2-}$ existed, as mentioned in Section 3.2. Two protonated EHBAA may preferentially extract a $[\text{SeO}_4]^{2-}$ complex *via* an anion-exchange reaction, as supported by the formation of a 1:2 complex in the organic phase. In addition, the decreasing extraction percentage

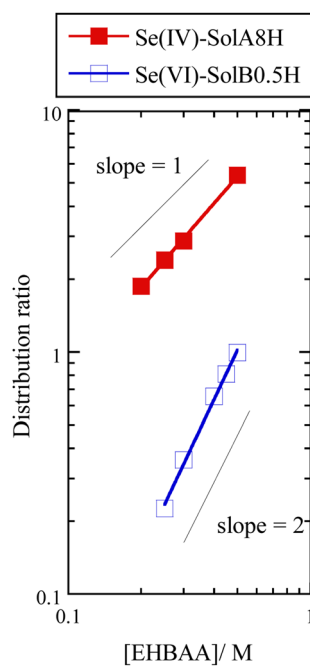
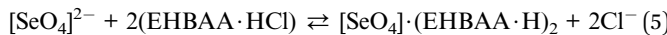


Fig. 6 Dependence of distribution ratio of Se(IV) and Se(VI) on EHBAA concentration. Organic phase, EHBAA pre-equilibrated with 0.5 or 8 M HCl; aqueous phase, SolA8H or SolB0.5H, $[\text{Se}] = 0.001 \text{ M}$.

of Se(vi) in the low HCl concentration range of 0.5–5 M (Fig. 2B) probably stems from the change in the dominant Se species in the order of $[\text{SeO}_4]^{2-} \rightarrow [\text{HSeO}_4]^- \rightarrow [\text{H}_2\text{SeO}_4]$.¹⁵ Similarly, the low extraction percentage reported for TBP in this low concentration region can be explained by its lower basicity (*i.e.*, its low protonation ability).³⁰ Therefore, the extraction reaction of Se(vi) from 0.5 M HCl can be represented as



4. Conclusions

Here, we used XAFS to investigate the extraction of Se by EHBA from two types of aqueous HCl solution (SolAxH made by dissolving SeO_2 and SolBxH made by dissolving $\text{Na}_2\text{SeO}_4 \cdot 10\text{H}_2\text{O}$). SolB8H mostly contained Se(iv) species due to a reduction of Se(vi) to Se(iv). In the EHBA-SolAxH system, the extraction percentage of Se rapidly increased at HCl concentrations above 5 M, but was poor below that limit. In contrast, in the SolBxH systems, ~50% of Se was extracted with EHBA at 0.5 M HCl; the extraction percentage of Se gradually decreased with an increase in HCl concentration up to 5 M; and then it rapidly increased at HCl concentrations above 5 M, although this rapid increase was due to the reduction to Se(iv). The XAFS measurements indicated that the inner-sphere structures of the dominant Se complexes were SeO_3 for SolA0.5H, a mixture of SeO_3 and SeOCl_2 for SolA8H, and SeO_4 for SolB0.5H. The EXAFS curve-fitting results showed that the Se complexes extracted with EHBA have inner-sphere structures of SeOCl_2 for SolA8H and of SeO_4 for SolB0.5H. The relationship between the distribution ratio of Se and EHBA concentration revealed that 1 : 1 and 1 : 2 Se : EHBA complexes were dominant in the organic phase of the SolA8H and SolB0.5H systems. The obtained results suggested that the dominant Se complexes with EHBA are $[\text{SeOCl}_2](\text{EHBA}\cdot\text{HCl})$ for SolA8H and $[\text{SeO}_4](\text{EHBA}\cdot\text{H})_2$ for SolB0.5H; the former was a solvation-type extraction, and the latter was an anion-exchange-type.

Author contributions

Hirokazu Narita: conceptualization, methodology, validation, formal analysis, investigation, writing – original draft, visualization, writing – review & editing; Motoki Maeda: validation, investigation, Chiharu Tokoro: methodology, visualization; Tomoya Suzuki: data curation, formal analysis; Mikiya Tanaka: writing – review & editing; Hideaki Shiwaku: methodology, formal analysis; Tsuyoshi Yaita: methodology, formal analysis.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors wish to thank Ms. Tomoko Saito for technical assistance. A part of this work was performed under the Shared

Use Program of JAEA Facilities (Proposal No. 2013B-E08) supported by JAEA Advanced Characterization Nanotechnology Platform as a program of “Nanotechnology Platform” of the Ministry of Education, Culture, Sports, Science and Technology (MEXT), Japan. The synchrotron radiation experiments were performed at JAEA beamline BL11XU in SPring-8 (Proposal No. 2013B3518).

Notes and references

- W. H. Midura and M. Midura, in *Encyclopedia of Inorganic and Bioinorganic Chemistry*, John Wiley & Sons, Ltd., 2014.
- S. Chaudhary, A. Umar and S. K. Mehta, *Prog. Mater. Sci.*, 2016, **83**, 270–329.
- G. Liu, Y. Wu, A. Tang, D. Pan and B. Li, *Hydrometallurgy*, 2020, **197**, 105460.
- Z. Dong, T. Jiang, B. Xu, J. Yang, Y. Chen, Q. Li and Y. Yang, *Chem. Eng. J.*, 2020, **393**, 124762.
- A. Polatajako, N. Jakubowski and J. Szpunar, *J. Anal. At. Spectrom.*, 2006, **21**, 639–654.
- Y. Liu, M. He, B. Chen and B. Hu, *Talanta*, 2015, **142**, 213–220.
- E. A. Azooz, M. Tuzen, W. I. Mortaba and N. Ullah, *Crit. Rev. Anal. Chem.*, 2022, DOI: [10.1080/10408347.2022.2153579](https://doi.org/10.1080/10408347.2022.2153579).
- Y.-C. Hou, C.-C. Chang, W.-L. Cheng and I.-S. Shaw, *Hydrometallurgy*, 1983, **9**, 381–392.
- M. R. Chowdhury and S. K. Sanyal, *Hydrometallurgy*, 1993, **32**, 189–200.
- M. R. Chowdhury and S. K. Sanyal, *Hydrometallurgy*, 1994, **34**, 319–330.
- A. Sattari, M. Kavousi and E. K. Alamdar, *Trans. Indian Inst. Met.*, 2017, **70**, 1103–1109.
- B. M. Sargan, S. V. Mahamuni and M. A. Anuse, *J. Saudi Chem. Soc.*, 2011, **15**, 177–185.
- W. Aeungmaitrepirom, A. Hagege, Z. Asfari, L. Bennouna, J. Vicens and M. Leroy, *Tetrahedron Lett.*, 1999, **40**, 6389–6392.
- W. Aeungmaitrepirom, A. Hagege, Z. Asfari, J. Vicens and M. Leroy, *J. Inclusion Phenom. Macrocyclic Chem.*, 2001, **40**, 225–229.
- F. Séby, M. P. Gautier, E. Giffaut, G. Borge and O. F. X. Donard, *Chem. Geol.*, 2001, **171**, 173–194.
- J. Milne and P. LaHaie, *Inorg. Chem.*, 1979, **18**, 3180–3183.
- H. Narita, K. Morisaku and M. Tanaka, *Chem. Commun.*, 2008, **45**, 5921–5923.
- H. Narita, K. Morisaku and M. Tanaka, *Solvent Extr. Ion Exch.*, 2015, **33**, 407–441.
- M. Maeda, H. Narita, C. Tokoro, M. Tanaka, R. Motokawa, H. Shiwaku and T. Yaita, *Sep. Purif. Technol.*, 2017, **177**, 176–181.
- T. Suzuki, T. Ogata, M. Tanaka, T. Kobayashi, H. Shiwaku, T. Yaita and H. Narita, *Metals*, 2018, **8**, 558.
- H. Narita, R. M. Nicolson, R. Motokawa, F. Ito, K. Morisaku, M. Goto, M. Tanaka, W. T. Heller, H. Shiwaku, T. Yaita, R. J. Gordon, J. B. Love, P. A. Tasker, E. R. Schofield, M. R. Antonio and C. A. Morrison, *Inorg. Chem.*, 2019, **58**, 8720–8734.



22 S. Uchino, H. Narita, K. Kita, H. Suzuki, T. Matsumura, H. Naganawa, K. Sakaguchi and K. Ohto, *Solvent Extr. Res. Dev., Jpn.*, 2023, **30**, 39–46.

23 F. E. Huggins, C. L. Senior, P. Chu, K. Ladwig and G. P. Huffman, *Environ. Sci. Technol.*, 2007, **41**, 3284–3289.

24 T. Ressler, *J. Synchrotron Radiat.*, 1998, **5**, 118–122.

25 A. L. Ankudinov, B. Ravel, J. J. Rehr and S. D. Conradson, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1998, **58**, 7565–7576.

26 R. B. Helmholdt, E. J. Sonneveld and H. Schenk, *Z. Kristall.*, 1999, **214**, 151–153.

27 A. Pawłowski and A. Haznar, *J. Solid State Chem.*, 2001, **160**, 189–194.

28 Y. Hermodsson, *Acta Chem. Scand.*, 1967, **21**, 1313–1327.

29 J. E. Penner-Hahn, *Coord. Chem. Rev.*, 1999, **190–192**, 1101–1123.

30 F. L. Bernardis, R. A. Grant and D. C. Sherrington, *React. Funct. Polym.*, 2005, **65**, 205–217.

31 J. Pettersson and Å. Olin, *Talanta*, 1991, **38**, 413–417.

32 P. Lahaie and J. Milne, *Inorg. Chem.*, 1979, **18**, 632–637.

33 F. Calderazzo, M. D'Attoma, F. Marchetti and G. Pampaloni, *J. Chem. Soc., Dalton Trans.*, 2000, 2497–2498.

