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Coordination characteristics of trivalent lanthanides and actinides in molten hydrate salts of Ca(NO₃)₂ and CaCl₂

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Abstract

Distribution behavior of Ce(III), Am(III), and Cm(III) between tri-*n*-butyl phosphate solution and molten calcium nitrate hydrate Ca(NO₃)₂·RH₂O or molten calcium chloride hydrate CaCl₂·RH₂O was studied radiochemically. In Ca(NO₃)₂·RH₂O systems, maximum separation factors of Ce and Cm to Am were observed to be 12 (Ce/Am) and 1.7 (Cm/Am). The distribution ratios of these elements increased with the decrease of water activity in the hydrates, and the extractabilities at the water deficient region was less sensitive compared to those at the water abundant region. This trend was similar to the coordination circumstance change observed in electronic absorption spectra of Nd(III) in the hydrates.

Keywords

Americium · Curium · Hydrate salt · Extraction · Activity

Introduction

Extraction of lanthanides and actinides in molten hydrate salts (generally called "hydrate melts") by an organic solution of tri-*n*-butyl phosphate (TBP) has been investigated for Ca(NO₃)₂·4H₂O [1-7], Mg(NO₃)₂·6H₂O [2], and Mn(NO₃)₂·6H₂O [8] media. A typical PUREX system (HNO₃-TBP system) shows low extractability for trivalent lanthanides and actinides, while using hydrate melts as the inorganic phase gives higher extractability of these elements. This is attributed to the enhanced chemical activities of nitrates due to the water deficiency of the system. The extractability is equivalent to a PUREX system with highly concentrated HNO₃ solutions [9,10]. The results have attracted technological interests for the purpose of recovering and separating lanthanides as fission products and trivalent actinides such as Am³⁺ and Cm³⁺ from high level radioactive wastes (HLLW) generated in the nuclear fuel cycle.

Beyond the HLLW technological interest, chemical behavior of trivalent lanthanides and actinides in hydrate melts is additional specific interest, especially, the coordination chemistry of f-elements in concentrated electrolytes. Cations, anions, and water molecules in hydrate melts contact each other continuously by forming a quasi-lattice structure [11]. A solute cation, *e.g.*, trivalent lanthanide or actinide, in a hydrate melt is surrounded by anions and water molecules which are contacted by cations of the hydrate melt matrix. The mean activity coefficient of salts is much larger than unity [6], while the activity coefficient of water is much smaller than unity [6,12], and they are drastically changed by changing water content. The specific coordination circumstance of solutes in the hydrate melts would make the chemical behavior of solute cation different from that in the usual aqueous solutions.

In the present study, we investigated the extraction distribution of trivalent Eu,

Am, and Cm between $Ca(NO_3)_2$ ·4H₂O and PUREX solvents (mixtures of TBP and *n*-dodecane). The extraction distribution of Am and Eu between $CaCl_2$ ·6H₂O and PUREX solvents was studied for comparison. Additional information on the coordination circumstance was obtained via the *f*-*f* hypersensitive transition of Nd(III) in the melts.

Experimental

Solvent extraction

All chemicals including Ca(NO₃)₂·4H₂O and CaCl₂·6H₂O were reagent grade from Wako Pure Chemical Industries, Ltd., and were used without further purification. The nitric acid solution containing ²⁴³Am, ²⁴⁴Cm (~100 kBq, respectively) was prepared and this was once evaporated to dryness. A few dm⁻³ (mL) of pure water was added and evaporated again. This process was repeated several times to remove nitric acid completely. The final residue, ²⁴³Am(NO₃)₃ and ²⁴⁴Cm(NO₃)₃ were dissolved into the molten Ca(NO₃)₂·4H₂O. An organic solution of 0.004 mol dm⁻³ (M) TBP diluted with *n*-dodecane (PUREX solvent) was prepared. This organic solution was pre-equilibrated with fresh Ca(NO₃)₂·4H₂O at room temperature.

Fifteen mL of the hydrate melt and 15 mL of the PUREX solvent were contacted by magnetic stirring in a glass centrifuge tube. The extraction was performed in a water bath at 343 K. Extraction equilibrium was attained in 2 hours. Under the equilibrium, two phases were separated by centrifugation and weighed portions of both phases were taken for γ -spectrometry to determine the distribution ratios (*D*) of ²⁴³Am and ²⁴⁴Cm. Intensities of γ rays (75 keV and 43 keV for ²⁴³Am and ²⁴⁴Cm, respectively) were analyzed. The self-shielding effect, which is the loss of transmittance of the γ rays

inside samples, was corrected as shown in our previous study [7]. All quantifications of hydrate melts and PUREX solvents were performed gravimetrically to avoid the large errors associated with volumetric treatment. The gravimetric radioactivities were converted to volumetric radioactivities using the density of solutions, in which literature values [13] or densities measured by employing a picnometer were used. Then, weighed TBP was added to the glass centrifuge tube to increase its molarity in the organic phase, and the same procedure was repeated. The extraction experiments with [TBP] ranging from 0.004 to 0.47 M were performed. Since Eu is a 4*f* element having similar ionic radius compared with those of Am and Cm [14], the same extraction experiment was performed with a radioactive tracer ¹⁵²Eu. *D* was determined from the γ -ray counts at 121.8 keV.

A similar extraction experiment of ²⁴³Am was performed for hydrated CaCl₂ medium. A portion of concentrated HCl was added to a dried ²⁴³Am(NO₃)₃ to convert its chemical form to be ²⁴³AmCl₃, then the solution was evaporated to dryness. This was dissolved in molten CaCl₂·6H₂O. An organic solution of 0.1 M TBP diluted with *n*-dodecane was prepared. This organic solution was pre-equilibrated with fresh CaCl₂·6H₂O at room temperature. The extraction procedure was the same with the Ca(NO₃)₂·4H₂O case. By adding weighed water into the system to increase water content, the extraction experiments with CaCl₂·*R*H₂O ranging from *R*=6 to 10 were performed at 343 K. In order to know the extraction stoichiometry, 0.007 M EuCl₃ dissolved in CaCl₂·6H₂O was extracted at 343 K by PUREX solvents including 0.1-0.3 M TBP. The extracted Eu and Ca in the organic phase were stripped into a diluted HCl and the concentrations of Eu and Ca were analyzed by ICP-AES (ICPS-1000TR, Shimadzu). [Eu] and [Ca] in the equilibrated CaCl₂·*R*H₂O phase were also analyzed.

Visible absorption spectrophotometry

The *f*-*f* hypersensitive transition, ${}^{4}G_{5/2}$, $\leftarrow {}^{4}I_{9/2}$, of Nd was analyzed by visible absorption spectrophotometry. Ca(NO₃)₂·*R*H₂O (*R* = 3.4-15) and CaCl₂·*R*H₂O (R = 5.7-15) containing Nd with [Ca]/[Nd] ~ 1000 were prepared. 0.1 M HClO₄, 0.01 M HNO₃, and 0.01 M HCl solutions containing ~0.01 M Nd were also prepared. [Nd] was determined by ICP-MS (HP-4500, Hewlett Packard). Each sample was transferred into a quartz cell and the absorption spectrum from 560 to 600 nm was measured at 0.1 nm interval by using an absorption spectrophotometer (UV-3100PC, Shimadzu). All measurements were performed at room temperature. In order to evaluate the oscillator strength, the refractive index of samples at room temperature was determined by using a refractometer (Fisher Scientific).

Results and discussion

Extraction behavior of Eu, Am, and Cm in $Ca(NO_3)_2$ hydrate melt by TBP

The extraction reaction of trivalent lanthanides and actinides (M^{3+}) in the PUREX system can be written [15],

$$M^{3+} + 3NO_3^{-} + 3TBP \rightleftharpoons M(NO_3)_3 \cdot 3TBP$$
⁽¹⁾

It is natural that the distribution of M (D_M) increases with the decrease of water content R of Ca(NO₃)₂·RH₂O by depressing the hydration to M. In the present study, the water activity, $a_{H_{2}O}$, is used as the representative of water deficiency. $a_{H_{2}O}$ is shown in Fig. 1 as a function of R of Ca(NO₃)₂·RH₂O. Since $a_{H_{2}O}$ of concentrated Ca(NO₃)₂ is not

available [12], the $a_{H_{2}O}$ values in Ca(NO₃)₂ hydrate melts were evaluated in the present study by using an electric hygrometer (Novasina IC-500, AW-LAB). Small $a_{\rm H_{2}O}$ can be seen at the water deficient region corresponding to hydrate melts. D_{Eu} , D_{Am} , and D_{Cm} obtained in our previous study [6,7] are shown in Fig 2a as functions of $a_{\rm H,O}$ newly estimated in the present study. The ionic radii of Eu3+, Am3+, and Cm3+ of 6-coordination in crystals [14] are similar but are in the order $Am^{3+} > Cm^{3+} > Eu^{3+}$. The D values are in the same order. The D values increase with the decrease of $a_{\rm H_{2}O}$ and log D > 1 can be achieved at the smaller log $a_{\rm H_2O}$ region. In this region, the extractability of Eu turned out to be smaller than that of Am and Cm. This is due to the inflection of log D vs. log $a_{\rm H,0}$ at ca. -0.33 for lanthanides [6] and ca. -0.4 for actinides [7]. This extraction trend change has been reported to be a coordination circumstance change of lanthanides and actinides in the Ca(NO₃)₂ hydrate melt [6]. The degree of the inflection is clearer for Eu than for Am and Cm. This may be attributable to different physicochemical properties between 4*f*-orbitals and 5*f*-orbitals.

The *D* values are shown in Fig. 2b as functions of [TBP]. The slopes of log *D vs*. log [TBP] were found to be 3 at small [TBP] region. This proves the validity of stoichiometry shown in reaction 1 for the hydrated $Ca(NO_3)_2$ system. The slope is moderated to be 2 with the increase of [TBP]. This is because the consumption of free TBP occurs by the extraction of Ca [4]. Though the coextraction of Ca occurs, the extractability of trivalent *f*-elements is improved to be log *D* > 2 at high [TBP] region.

The separation factors compared to Am, $D_{\rm Eu}/D_{\rm Am}$ and $D_{\rm Cm}/D_{\rm Am}$, are shown in Fig. 3a and 3b as functions of $a_{\rm H,O}$ and [TBP]. As a maximum separation factor shown

in Fig 3a, $D_{\rm Eu}/D_{\rm Am} = 6.6$ was achieved at log $a_{\rm H_2O} \sim -0.25$ ($R \sim 7$), while $D_{\rm Eu}/D_{\rm Am} =$

1.4 was achieved at log $a_{\rm H_2O} \sim -0.40$ ($R \sim 5$). The $D_{\rm Eu}/D_{\rm Am}$ value decreased to 1.1 at the water abundant region. The dependence of separation factor on [TBP] was not found for $D_{\rm Cm}/D_{\rm Am}$ which was ranged in 1.6-1.7 (Fig 3b). $D_{\rm Eu}/D_{\rm Am}$ increased with the decrease of [TBP] which reached to 12, but their extractability was small as log D < -1 (see Fig. 2b). The separation factor $D_{\rm Eu}/D_{\rm Am}$ was reported to be 8.7-84 for an extraction system with an extractant TPEN (N,N,N',N'-tetrakis(2-methylpyridyl)-ethylenediamine) [16] and 960-7900 for a system with Cyanex 301 (bis(2,4,4-trimethylpentyl)dithiophosphinic acid) [17]. Our results would be too small on the viewpoint of Eu/Am separation. The separation factor $D_{\rm Eu}/D_{\rm Am}$ was reported to be 1.25-1.78 for several complexing reagents [15,18, and references therein]. Our result $D_{\rm Eu}/D_{\rm Am} \sim 1.7$ showed a high value in the reported range, though the typical extractant TBP was used. Water deficient media may enhance the difference of chemical activities between Am and Cm.

Extraction behavior of Am in CaCl₂ hydrate melt by TBP

The water activity a_{H_2O} in CaCl₂·6H₂O is almost the same with that in Ca(NO₃)₂·4H₂O (Fig. 1). The extraction behavior of trivalent *f*-elements in CaCl₂ hydrate melts by TBP therefore is of interest. D_{Am} obtained is shown in Fig. 4a as functions of a_{H_2O} . Low extractability of Am in the CaCl₂ hydrate melt system is natural by considering the small *D* values of trivalent *f*-elements reported for HCl-TBP systems [19,20]. Though [TBP] was 5 times larger than that of the Ca(NO₃)₂ case, D_{Am} was found to be ~100 times smaller. Inflection of log *D* vs. log a_{H_2O} at *ca*. -0.40 was also seen and its degree

looks larger than that of the $Ca(NO_3)_2$ case.

In order to know the extraction stoichiometry, dependence of D_{Eu} on [TBP] was studied. The results are shown in Fig. 5. In the [TBP] region less than 0.1 M, the extraction of Ca was negligibly small. The consumption of TBP by the extraction of Ca was found to increase with [TBP], and at [TBP] = 0.1 M, 3% of TBP was consumed. The slope of log D_{Eu} vs. log [TBP] showed 3 at small [TBP] region, and hence, the extracted species would be EuCl₃·3TBP.

Dependence of $D_{\rm M}$ *on water activity*

The extraction reaction 1 can be rewritten,

$$M(H_2O)_m^{3+} + 3X^- + 3TBP \neq MX_3 \cdot 3TBP + mH_2O$$
⁽²⁾

where X⁻ represents NO₃⁻ or Cl⁻. The stability constant K_{ex} of this reaction is,

$$K_{\rm ex} = \frac{\gamma_{\rm MX_3 \cdot 3TBP} [\rm MX_3 \cdot 3TBP] a_{\rm H_2O}{}^{\rm m}}{\gamma_{\rm M(H_2O)_m{}^{3+}} [\rm M(H_2O)_m{}^{3+}] \gamma_{\rm X^{-}}{}^{\rm 3} [\rm X^{-}]^{\rm 3} a_{\rm TBP}{}^{\rm 3}}$$
(3)

Because of $[MX_3 \cdot 3TBP]/[M(H_2O)_m^{3+}] = D_M$ and $\gamma_{M(H_2O)_m^{3+}} \gamma_{X^-}^3 = \gamma_{MX_3}^4$, Eq. 3 is

rewritten,

$$K_{\rm ex} = \frac{\gamma_{\rm MX_3:3TBP} \ D_{\rm M} \ a_{\rm H_2O}}{\gamma_{\rm MX_3}^{4} \ [\rm X^-]^3 \ a_{\rm TBP}^{3}}$$
(4)

Logarithm of $D_{\rm M}$ is hence,

$$\log D_{\rm M} = \log K_{\rm ex} + 4 \log \gamma_{\rm MX_3} + 3 \log [{\rm X}^-] + 3 \log a_{\rm TBP}$$
(5)
$$- \log \gamma_{\rm MX_3,3TBP} - m \log a_{\rm H_2O}$$

For the experiments with constant [TBP] = 0.02 M for hydrated Ca(NO₃)₂ systems and with [TBP] = 0.1 M for hydrated CaCl₂ systems, γ and a of species in the organic phases ($\gamma_{MX_3,3TBP}$ and a_{TBP}) can possibly be treated as constants. In Eq. 5, (log K_{ex} + 3 log a_{TBP} – log $\gamma_{MX_3,3TBP}$) is treated as a constant value.

$$\frac{1}{4} (\log D_{\rm M} - 3 \log [{\rm X}^-]) = -\frac{m}{4} \log a_{\rm H_2O} + \log \gamma_{\rm MX_3} + \text{Const.}$$
(6)

 $\frac{1}{4}$ (log $D_{Am} - 3 \log [X^-]$) is shown in Fig. 4b as functions of log a_{H_2O} . The relationship $\frac{1}{4}$ (log $D_{Am} - 3 \log [X^-]$) vs. log a_{H_2O} shows a similar inflection property with Fig. 4a. This means that the correction of D_M by $[X^-]$ is not the origin of this inflection. According to Eq. 6, the breakdown of the linear relation of $\frac{1}{4}$ (log $D_M - 3 \log [X^-]$) vs. log a_{H_2O} would be due to a specific change in γ_{MX_3} .

Variation of γ_{MX_3} results from the coordination circumstance change of M^{3+} in hydrate melts with various water contents. In order to know it, electronic absorption spectra of Nd³⁺ in Ca(NO₃)₂·*R*H₂O and CaCl₂·*R*H₂O were measured. The electric dipole transitions between *f*-orbitals that show abnormal variations in intensity are classified as hypersensitive transitions [21]. These variations are attributable to the action of an inhomogeneous electromagnetic field from the surrounding medium. In the case of neodymium, transitions from the ground level ${}^4I_{9/2}$ to excited levels of ${}^2G_{7/2}$ and ${}^4G_{5/2}$, which are abbreviated as ${}^4G_{5/2}, {}^2G_{7/2} \leftarrow {}^4I_{9/2}$, are sensitive to the coordination circumstances, and they are commonly used to examine the coordination of Nd³⁺ [21]. The transition ${}^{4}G_{5/2} \leftarrow {}^{4}I_{9/2}$ is the hypersensitive transition, which is accompanied by and overlapped with a less intensive non-hypersensitive transition to the level ${}^{2}G_{7/2}$. The information on coordination environment of Nd³⁺ would be helpful to understand that of trivalent lanthanides and actinides having similar ionic radii.

The measured spectra of the hypersensitive transition are shown in Fig. 6a and 6b. The peak position of ${}^{4}G_{5/2} \leftarrow {}^{4}I_{9/2}$ transition is shown in Fig. 6b by referring literature [22,23]. The spectra of Nd³⁺ in low acidity solutions (0.01 M HNO₃ and HCl) were almost the same and were similar to that in HClO₄ solutions [24,25]. The spectra were clearly changed with the decrease of water content. The oscillator strength (the index for the transition probability) of the hypersensitive *f*-*f* transition of Nd is known to be larger for stronger interaction between Nd³⁺ and ligands [21 and references therein]. The oscillator strength can be defined as,

$$f = 4.319 \times 10^{-9} \frac{9n}{(n^2 + 2)^2} \int \varepsilon(v) dv$$
⁽⁷⁾

where $\varepsilon(v)$ is the molar absorptivity at energy $v(\text{cm}^{-1})$ and *n* is the refractive index of the solvent medium. The *f* value of the results shown in Fig. 6 was evaluated (Fig. 7a). The *f* values increased with the decrease of water content for both systems. At a fixed temperature, the factors influencing *f* are: (i) symmetry of Nd(III) complex and (ii) electron donating ability of ligands around Nd³⁺. The increase of *f* shown in Fig. 7a suggests that the tricapped trigonal prism [22], which is a typical 9 hydration for lighter lanthanides, is distorted. The substitution of hydrated waters of Nd³⁺ by NO₃⁻ or Cl⁻ would have occurred. As shown in Fig. 6a, the intense increase of ${}^4G_{5/2} \leftarrow {}^4I_{9/2}$ transition probability of Nd(III) in Ca(NO₃)₂·*R*H₂O is suggestive of the substitution of hydrated H₂O around Nd³⁺ by NO₃⁻. *f* may have been enhanced by a direct interaction between Nd³⁺ and NO₃⁻.

The structural studies on Ca(NO₃)₂·*R*H₂O by x-ray diffraction [26,27] and neutron diffraction [28] analyses have pointed out the contact ion pair of Ca²⁺ and NO₃⁻. Their results suggest that the contact ion pair of Nd³⁺ and NO₃⁻ is also possible in Ca(NO₃)₂·*R*H₂O. The hydration number of M³⁺ in Ca(NO₃)₂·*R*H₂O has been found to decrease with water content [29]. Hence, the changes in spectra (Fig. 6a) and *f* (Fig. 7a) of Ca(NO₃)₂·*R*H₂O system would be consistent with the occurrence of contact ion pair of Nd³⁺ and NO₃⁻.

The spectra shown in Fig. 6b do not show the drastic increase of the ${}^{4}G_{5/2} \leftarrow {}^{4}I_{9/2}$ transition probability of Nd(III) in CaCl₂·*R*H₂O. The reason is reported to be a weaker interaction of Cl⁻ with H₂O molecules in the inner hydration sphere of Nd³⁺ [30]. The specific feature of Fig. 6b may be the energy shift of the ${}^{4}G_{5/2} \leftarrow {}^{4}I_{9/2}$ transition peak. This may be correlated with the nephelauxetic effect [31]. It is known that the red shift (depression of transition energy) is caused by the strong interaction between *f*-elements (lanthanides and actinides) and ligands [31,32]. The stronger interaction reduces the values of the free-ion parameters and causes a contraction of energy level structures of the *f*-element ions in matrix. In the present study, the spectra shown in Fig. 6b were decomposed into 6 absorption peaks by Gaussian/Lorentzian sum function, and the peak position of the ${}^{4}G_{5/2} \leftarrow {}^{4}I_{9/2}$ transition was estimated. It should be noted that, since there are many transitions from sub-levels of ${}^{4}I_{9/2}$ transition by referring literatures [22,23] might be interfered by the transitions between sub-levels. The parameter $\beta = v_{\text{sample}}/v_{\text{reference}}$ [23,31], where v is the wavenumber (cm⁻¹) at the ${}^{4}G_{5/2} \leftarrow {}^{4}I_{9/2}$ transition, was determined. The subscript "reference" means 0.01 M HNO₃ for nitrate systems or 0.01 M HCl for chloride systems. Figure 7b shows $1 - \beta$ as a function of $a_{\text{H}_{2}\text{O}}$. The $1 - \beta$ value increased with the decrease of $a_{\text{H}_{2}\text{O}}$. This means that the red shift occurs with the decrease of water content. This trend is consistent with a report on Nd³⁺ in concentrated LiCl [23].

Structural studies on CaCl₂·6H₂O by x-ray diffraction [33,34] showed that ~1 Cl⁻ ion exists in the first hydration sphere of Ca²⁺. Their results suggest that the contact ion pair of Nd³⁺ and Cl⁻ is possible in CaCl₂·*R*H₂O. In a study on Nd³⁺ in concentrated LiCl solutions, it was suggested that, with the decrease of water content, the hydrated H₂O around Nd³⁺ is substituted by Cl⁻ and the coordination number decreases from 9 to 8 [23]. Since the electron donating ability of O in H₂O is different from that of Cl⁻, varying the water content changes the ligand field around Nd³⁺ and this may have caused the shift in peak position shown in Fig. 6b and also the increase of *f* (Fig. 7a).

The correlations $f vs. \log a_{H_2O}$ and $1 - \beta vs. \log a_{H_2O}$ (Fig. 7) possess similar trends of $\frac{1}{4} (\log D_{Am} - 3 \log [X^-]) vs. \log a_{H_2O}$ shown in Fig. 4b. However the inflection property shown in Fig. 7a and 7b is not clear compared with that shown in Fig. 4b. This suggests that the inflection depends not only on the short distance as the first coordination sphere of M³⁺ but also the long distance like second and far coordination spheres.

Conclusions

Coordination characteristics of trivalent lanthanides and actinides in hydrate melts were investigated by solvent extraction method and spectrophotometry. In PUREX experiments, distribution behavior of trivalent lanthanides and actinides in water deficient system was different from that in water abundant system. Contact ion pairing of trivalent *f*-elements with anions in inorganic phase occurs with the decrease of water content, and the coordination circumstance change would result in the change of distribution behavior. The distribution behavior may also be affected by changes in far coordination spheres.

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Figure captions

Fig. 1 Water activities in Ca(NO₃)₂·*R*H₂O and CaCl₂·*R*H₂O at 343 K. Open marks were literature values [12]. $a_{H_{2}O}$ of solid marks was estimated in the present study by using an electric hygrometer (Novasina IC-500, AW-LAB).

Fig. 2 Distribution ratios of Am, Cm, and Eu between Ca(NO₃)₂·*R*H₂O and PUREX solvent. a) Dependence of log *D* on log a_{H_2O} . D_{Am} , D_{Cm} , and D_{Eu} measured in our previous study [6,7] were reproduced as functions of a_{H_2O} obtained in the present study. Experimental conditions are [TBP] = 0.02 M and *T* = 343 K. b) Dependence of log *D* on log [TBP]. 1 σ errors due to γ -analysis are shown together. The inorganic phase is Ca(NO₃)₂·4H₂O and experimental temperature is *T* = 343 K.

Fig. 3 Separation factors of Eu/Am and Cm/Am. a) Dependent of separation factor on log $a_{H,O}$. D_{Eu}/D_{Am} and D_{Cm}/D_{Am} were calculated from fitting curves shown in Fig.

2a. b) Dependence of separation factor on log [TBP]. D_{Eu}/D_{Am} was calculated from data shown in Fig. 2b. Error bars of D_{Cm}/D_{Am} are attributable to errors of D_{Am} and D_{Cm} .

Fig. 4 Distribution ratios of Am between CaCl₂·*R*H₂O and PUREX solvent. a) D_{Am} . D_{Am} between Ca(NO₃)₂·*R*H₂O and PUREX solvent (Fig. 2a) is shown together. b) $\frac{1}{4}$ (log D_{Am} – 3 log [X⁻]). X⁻ means NO₃⁻ and Cl⁻. Details of derivation can be seen in Eqs. 3-6.

Fig. 5 Distribution ratios of Eu between CaCl₂·6H₂O and PUREX solvent with various [TBP].

Fig. 6 Electronic absorption spectra of hypersensitive transition of Nd(III) in hydrate melts. The absorption spectra are measured at room temperature. a)

 $Ca(NO_3)_2 \cdot RH_2O. b) CaCl_2 \cdot RH_2O.$

Fig. 7 Oscillator strength of hypersensitive transition and shift in peak position for absorption spectra of Nd(III) in hydrate melts. a) Oscillator strength of the ${}^{4}G_{5/2}, {}^{2}G_{7/2} \leftarrow {}^{4}I_{9/2}$ transition. The *f* values (Eq. 7) are shown as functions of $a_{H_{2}O}$. b) Energy shift of the ${}^{4}G_{5/2} \leftarrow {}^{4}I_{9/2}$ transition peak for Nd³⁺ in CaCl₂·*R*H₂O. The absorption spectra shown in Fig. 6b were decomposed into 6 absorption peaks by Gaussian/Lorentzian sum function, and the peak position of the ${}^{4}G_{5/2} \leftarrow {}^{4}I_{9/2}$ transition was estimated referring literatures [22,23]. The 1 – β value (see text) is shown as a function of $a_{H_{2}O}$.



Figure 1 (Fujii et al.)



Figure 2 (Fujii et al.)



Figure 3 (Fujii et al.)



Figure 4 (Fujii et al.)



Figure 5 (Fujii et al.)



Figure 6 (Fujii et al.)



Figure 7 (Fujii et al.)