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A kinetic interpretation of the effect of extractant and acid on neodymium(III) extraction by a diglycolamide

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With an increasing demand for rare earth elements in modern technologies, it is crucial to develop a full understanding of the purification process for lanthanides. Recent studies have focused heavily on the separation of lanthanides using solvent extraction with diglycolamide extractant ligands. These completely incinerable extractants, especially *N,N,N',N'*-tetra(*n*-octyl)diglycolamide (TODGA), provide best-in-class thermodynamic separation for the early-middle lanthanides. Thermodynamic parameters are well characterized for lanthanide extraction from nitric acid media by TODGA, but there exists a lack of knowledge about the kinetics. We developed a greater understanding of the rates and mechanism of lanthanide extraction by TODGA using *operando* visible spectroscopy to monitor changes in a coordination-sensitive neodymium(III) absorbance band. Due to the dynamic interplay between equilibrium metal distribution and the rate of mass transfer, determining these parameters across a wide range of component concentrations revealed novel thermodynamic and kinetic findings. An activity-based treatment of the non-linear dependence of metal extraction on acidity allowed us to quantitatively identify the changes in speciation across the acidity range. Plateau and linear regions in the kinetic data are directly related to slopes in the equilibrium data, which, together, provide the reaction orders with respect to each component. Extraction and stripping rate laws follow the interfacial two-step consecutive reactions mechanism, where adsorption and desorption of neodymium(III) complexes at the interface govern the rate limiting steps, while TODGA, nitrate anions, and nitric acid adducts associate and dissociate in fast equilibria.

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Introduction

Lanthanide elements are critical to a range of high-performance magnetic and optical materials, but individual lanthanides are difficult to separate from one another due to their chemical similarity.^{1–3} Most separation chemistry for the purification of lanthanide elements uses organic extractants to form lanthanide complexes that are soluble in organic solvents, and the small differences in the lanthanide ionic radii can then be exploited to separate the lanthanides by liquid–liquid extraction.⁴ However, the selectivity for specific lanthanide elements is poor, and hundreds of individual extraction stages are often required to achieve high purity for one lanthanide.⁵ Understanding the physicochemical features underlying lanthanide separation processes and developing more efficient separating ligands are key areas of research for reducing the resources consumed in lanthanide production.

The diglycolamide (DGA) ligand *N,N,N',N'*-tetra(*n*-octyl)diglycolamide (TODGA)⁶ (Fig. 1) and its derivatives⁷ have emerged as promising extractants for lanthanide(III) cations with intra-lanthanide separation capabilities comparable to the best-in-class organophosphorus extractants. While some questions remain, thermodynamic aspects of the extraction, such as the degree of lanthanide(III) extraction and the equilibrium organic phase complexes formed by TODGA or other common tetra-alkyldiglycolamides, have received considerable attention.^{8–10} Since DGAs are solvating extractants, they tend to form supra-molecular complexes that also contain the lanthanide cation, charge-balancing anions, water, and mineral acids from the

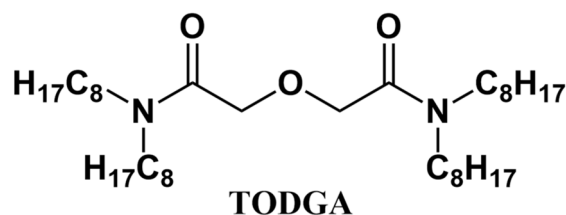


Fig. 1 The chemical structure of *N,N,N',N'*-tetra(*n*-octyl)diglycolamide (TODGA).

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aqueous phase.¹¹ Consequently, extractions involving tetraalkyl-DGAs have an unusually strong and complex dependence on the concentration of aqueous acid, which has been difficult to reproduce with an equilibrium thermodynamic model.^{12,13}

The kinetic aspects of DGA extraction systems, such as rate laws, rate constants, and mechanisms, have rarely been characterized, likely due to the difficulty in obtaining and interpreting quantitative kinetic data for liquid–liquid extraction systems.^{14,15} The Lewis cell¹⁶ and rotating membrane cell^{17,18} techniques have been used to probe the TODGA-HNO₃ system and found varying results. Using a Lewis cell, the extraction of Am(III) was found to be first order with respect to TODGA, HNO₃, and Am(III). A chemically-controlled reaction mechanism was proposed to rationalize the results;¹⁶ however, the authors conclude that the extraction is at least partially controlled by the rate of diffusion rather than chemical reactions in their Lewis cell.¹⁶ A comprehensive study of Eu(III) and Am(III) extraction by TODGA in 5 vol% 1-octanol/hydrogenated tetrapropylene using a rotating membrane cell indicated decreasing forward and reverse rate constants with increasing TODGA concentration, while increasing the acidity led to faster forward (extraction) rates and slower reverse (stripping) rates.¹⁸ In that study, the increasing metal extraction with increasing TODGA concentration was primarily attributable to the decrease in the rate of the stripping reaction.

To develop a better understanding of the reaction orders and mechanism under conditions where the chemical reactions are rate controlling, we used *operando* UV/visible absorption spectroscopy to study the kinetics of Nd(III) partitioning between solutions of TODGA in *n*-dodecane and aqueous nitric acid. Our findings with this highly-stirred technique shed new light on the mechanisms of lanthanide phase transfer in diglycolamide-based separations, and they provide insight into the extraction and equilibrium speciation in the bulk organic phase.

Experiment

TODGA (>99%) was synthesized and purified in our lab according to the procedure in the SI. Nitric acid (ACS grade) was purchased from Sigma-Aldrich and *n*-dodecane (99%) was purchased from ThermoFisher Scientific. DI water (18.2 MΩ) was obtained from a Millipore Direct-Q water purifier. A stock solution containing 0.478 M Nd(NO₃)₃ was diluted to prepare all the aqueous Nd-containing solutions.

Organic solutions were prepared by dissolving the appropriate mass of TODGA in *n*-dodecane. Aqueous phases were prepared by diluting appropriate masses of concentrated HNO₃ and the Nd(NO₃)₃ stock solution with DI water. For the acid dependency experiments, solutions were prepared over a range of HNO₃ concentrations (0.20–3.0 M) while the concentration of TODGA and Nd(III) were held at 0.10 M and 4.87 ± 0.04 mM, respectively. The concentration range for the TODGA dependency was 0.060–0.40 M TODGA, while the HNO₃ and Nd(III) concentrations were held at 0.50 M and 4.91 ± 0.04 mM, respectively. For each condition, 15 mL of organic phase was pre-equilibrated with an equal volume of the corresponding metal-free acid solution. Phases were vortexed at 2000 rpm for

10 min using a Benchmark Scientific Benchmixer Vortex Mixer and centrifuged for 5 min. The aqueous phase was removed, and the process repeated. After two pre-equilibrations, the organic phase was separated for continued use. Extraction experiments were performed by contacting equal volumes of pre-equilibrated organic phase and metal-loaded aqueous phase for 5 min and subsequent centrifugation. All extractions were performed in triplicate at 295 ± 1 K. The initial and equilibrium Nd concentrations were determined by Inductively Coupled Plasma – Optical Emission Spectroscopy using a PerkinElmer Avio 220 Max analyzing two emission wavelengths in both radial and axial viewing modes. The distribution ratio (*D*) at equilibrium was calculated according to eqn (1), where organic phase species are indicated by an overbar, while aqueous species have none, subscript init represents the initial concentration, and subscript eq represents the equilibrium concentration.

$$D_{\text{Nd}} = \frac{[\overline{\text{Nd}}]_{\text{eq}}}{[\text{Nd}]_{\text{eq}}} = \frac{[\text{Nd}]_{\text{init}} - [\text{Nd}]_{\text{eq}}}{[\text{Nd}]_{\text{eq}}} \quad (1)$$

Interfacial tension measurements of TODGA solutions in contact with varying concentrations of nitric acid were collected by the pendant drop method on a Biolin ThetaFlex at 293 ± 1 K. The densities of the organic and aqueous phase were determined using an Anton Parr DMA 1001 with a precision of ±0.1 mg mL⁻¹ to correct the recorded tension values. The pre-equilibrated organic phase (20 μL) was dispensed into fresh aqueous solution by a 14-gauge hooked needle, and 140 frames of images over 10 seconds were analyzed. All solutions were measured in duplicate.

Kinetic experiments were performed under pseudo first order reaction conditions with at least 12-fold excess of nitric acid and TODGA. The chemical form and phase distribution of the Nd(III) species in the mixture were followed as a function of time using the OLIS RSM-1000 spectrophotometer with CLARiTY attachment. This dual-monochromator instrument equipped with 1200 lines/mm gratings can acquire 1–100 scans per second over a wavelength range of approximately 70 nm with high resolution. The CLARiTY attachment uses an 8 mL round-bottom sample cell surrounded by a reflective coating that reflects the light throughout the cell multiple times before being collected and amplified by PMTs, leading to longer effective pathlengths and larger absorbance values for a given sample volume. A custom-made overhead stirrer was used in these experiments, which closed the system from stray light, and provided precise stirring speed control.

All kinetic experiments were performed following this sequence: the aqueous phase (3.95 mL), and then the pre-equilibrated organic phase (3.95 mL) were each carefully added to the cell to avoid pre-emptive mixing of the phases. The overhead stirrer was lowered into the cell, then data collection and stirring were started. The two phases were stirred vigorously at 57 rps in the round bottom cell at 295 ± 0.5 °C. Spectra were collected from 547 – 618 nm at 3.1 scans per second over 50 seconds to monitor the ⁴I_{9/2} → ⁴G_{5/2}, ²G_{7/2} hypersensitive



electronic transitions of Nd(III). Spectra were corrected with an instrumental background and further baseline corrected using the `msbackadj` function in the MATLAB Bioinformatics Toolbox. The resulting 3-dimensional data (spectral changes over time) were analyzed with the OLIS GlobalWorks software.

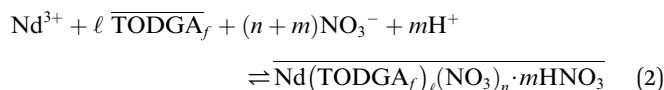
The specific interfacial area of the stirred samples was determined from the Sauter mean diameter of the organic droplets in the stirred mixture across the range of TODGA and nitric acid concentrations studied. Images of the stirred phases were acquired at 150 fps using a microscopic camera that was focused on an identical 8 mL round-bottom cell and stirrer without the external reflective coating or housing that replicated the dimensions of the CLARiTY cell. These images were randomly sampled and analyzed in ImageJ for the drop diameter after calibration of the images against those of $95 \pm 5 \mu\text{m}$ polystyrene beads. The mean diameter of the drops ranged from 169 to 199 μm across the solution conditions considered.

All uncertainties are given at the 95% confidence level.

Results and discussion

Equilibrium analysis

Equilibrium analysis of Nd extraction by TODGA is useful for understanding the stoichiometries of the Nd complexes present in the bulk aqueous and organic phases at equilibrium. It is also necessary for analysis of the kinetic experiments. A general form of the extraction equilibrium for the partitioning of Nd between aqueous nitric acid solutions and TODGA/*n*-dodecane can be written as



where species present in the organic phase are indicated with an overbar, and the equilibrium constant for the reaction is K_{ex} . Rearranging the equilibrium constant expression for this reaction and introducing the equilibrium Nd distribution ratio, D_{Nd} , the dissociation constant of nitric acid, and the stability constants of aqueous neodymium nitrate complexes (SI), expressions for the TODGA and nitrate dependencies of the measured equilibrium Nd distribution ratio can be derived such that

$$\log D_{\text{Nd}} = \ell \log [\overline{\text{TODGA}}_f] + (n+2m)\log\{\text{NO}_3^-\} + \log K_{\text{ex}} \quad (3)$$

where square brackets indicate molar concentrations and braces indicate molar scale activities such that $\{\text{NO}_3^-\} = \gamma_{\pm}[\text{NO}_3^-]$, $[\overline{\text{TODGA}}_f]$ represents the free concentration of TODGA in the organic phase that is neither bound to metal nor nitric acid, and the approximation $\{\text{H}^+\} \approx \{\text{NO}_3^-\}$ is used.

The effects of varying $[\overline{\text{TODGA}}_f]$ at constant acidity and varying the aqueous nitrate activity at constant TODGA concentration are shown in Fig. 2. Linear least squares fitting of eqn (3) to the data in Fig. 2A, yields a value of $\ell = 3.25 + 0.04$ for the average equilibrium Nd:TODGA stoichiometry in the organic phase. This indicates that there is a mixture of Nd

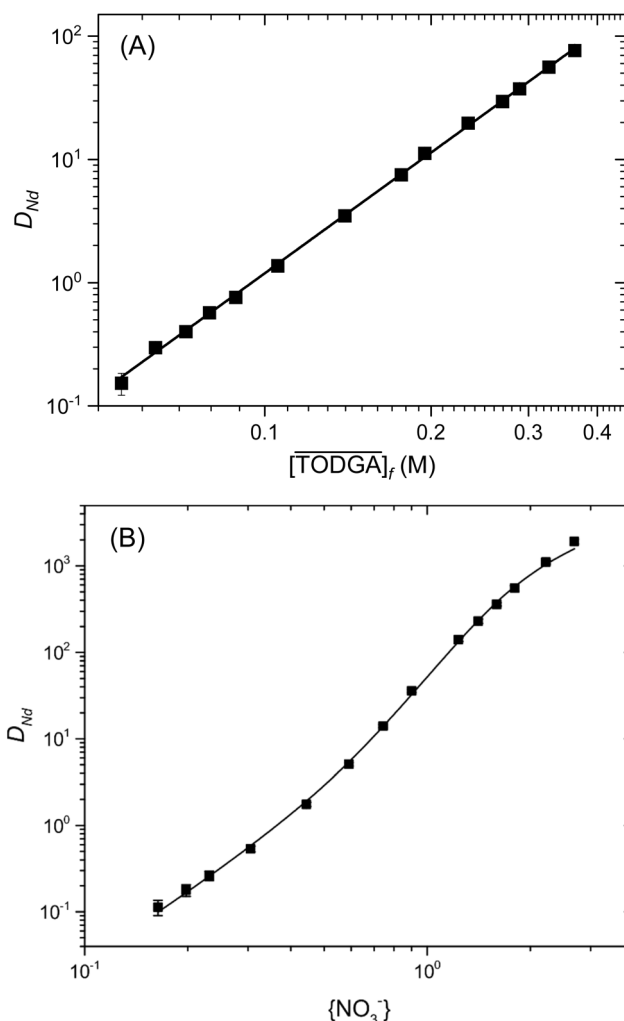


Fig. 2 Variation in equilibrium Nd distribution ratios between aqueous nitric acid and TODGA/*n*-dodecane at 295 K. (A) Extraction from 0.50 M HNO_3 ($\{\text{NO}_3^-\} = 0.37$) by varying concentrations of TODGA with an initial aqueous Nd concentration of 4.91×10^{-3} M. (B) Extraction from varying aqueous concentrations of nitric acid into 0.1 M TODGA/*n*-dodecane with an initial aqueous Nd concentration of 4.89×10^{-3} M. Lines represent the best fit of eqn (3) (panel A) or SI eqn S9 (panel B).

complexes containing 3 or 4 TODGA molecules in the organic phase under these conditions, which agrees well with the literature, where slopes between 3 and 4 are found for lanthanide(III) extraction by TODGA from nitric acid into aliphatic and aromatic diluents.^{19–21} Previous EXAFS studies of the extracted Nd-TODGA complexes in the bulk organic phase were consistent with coordination of three tridentate TODGA molecules with the counter anions residing in clefts between the coordinated TODGA molecules.⁹ With this core $\text{Nd}(\text{TODGA})_3(\text{NO}_3)_3$ structure, the fourth TODGA molecule and any coextracted HNO_3 would reside in the outer sphere of a lanthanide-centered supramolecular assembly.²²

The equilibrium and extraction rate experiments cover a wide range of aqueous phase nitric acid concentrations (0.20–3.0 M). The activity coefficients of the aqueous species are well-



known to vary significantly over this range, therefore the aqueous phase nitrate ion activity was used to model the nitric acid dependence of the extraction.²³ However, as expected from previous studies of lanthanide-DGA extraction, the equilibrium nitrate activity dependence of Nd extraction at constant TODGA concentration (Fig. 2B) is non-linear over the range of nitric acid concentrations studied^{6,19} and cannot be modeled well using eqn (3) with a single set of stoichiometric coefficients and K_{ex} values. At low nitrate activity, the Nd distribution ratios in Fig. 2B show the expected third-power dependence on the nitrate activity, but at higher nitrate activities, the measured distribution ratio exceeds a fourth-power dependence on the aqueous nitrate activity.

The possible reasons for this very steep acid dependence have been discussed elsewhere,^{12,13,19} but the presence of two distinct nitrate dependence slopes in this data implies that multiple species containing different numbers of nitrate anions (n) and nitric acid molecules (m) are present in the organic phase at equilibrium. Consequently, a more complete model that accounts for the presence of multiple organic phase complexes and also considers changes in the activity coefficients of other aqueous species, for example Nd, was used to model the data as summarized in the SI and Fig. S1–S3. The experimental distribution data is well fit by this equilibrium model with three Nd complexes in the organic phase (Fig. 2B and SI Fig. S2). The first complex, $\text{Nd}(\text{TODGA})_\ell(\text{NO}_3)_3$, predominates at low nitric acid concentrations ($[\text{HNO}_3] \leq 0.50 \text{ M}$, $\{\text{NO}_3^-\} \leq 0.37$) where the nitrate dependence slope of eqn (3) is $(n + 2m) = 3.03 \pm 0.08$, and from charge balance considerations $n = 3$ and $m = 0$. At higher aqueous nitrate activities, a mixture of $\text{Nd}(\text{TODGA})_\ell(\text{NO}_3)_3 \cdot \text{HNO}_3$ and $\text{Nd}(\text{TODGA})_\ell(\text{NO}_3)_3 \cdot 2\text{HNO}_3$ exists in the organic phase, and intermediate slopes are observed. For the highest nitrate activities studied, the nitrate dependence slope of eqn (3) corrected for the activity coefficients of the aqueous species is $(n + 2m) = 6.9 \pm 0.4$. This is consistent with organic phase complexes that contain 3 NO_3^- and $2.0 \pm 0.2 \text{ HNO}_3$ molecules at the highest acidities studied. These results agree with previous findings with TODGA¹⁹ and a similar branched tetraalkyldiglycolamide,¹² where an average of 1–2 HNO_3 molecules were determined to associate with organic phase lanthanide complexes when the acidity rises.

Therefore, from the perspective of equilibrium thermodynamics, the unusually large dependence of the measured distribution ratio on the aqueous nitrate activity can be understood as the result of the formation of organic phase Nd complexes that contain both the 3 nitrate anions required to form a neutral $\text{Nd}(\text{III})$ complex and an additional 1 or 2 nitric acid molecules.^{12,19}

Interfacial tension

The interfacial tensions between solutions of TODGA in *n*-dodecane and aqueous solutions of nitric acid were measured across a range of compositions (Fig. 3). The constant slope regions of the interfacial tension curves show that the aqueous-organic interface is saturated with TODGA molecules when the

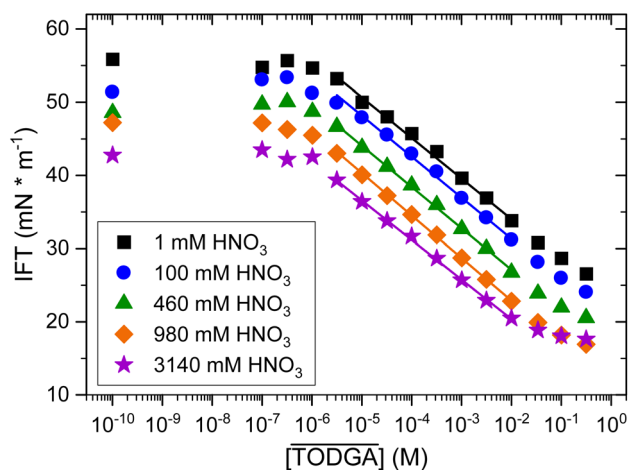


Fig. 3 Interfacial tensions of the nitric acid – TODGA/*n*-dodecane system.

TODGA concentration exceeds $1 \times 10^{-5} \text{ M}$ for all the aqueous acidities studied, with an interfacial excess corresponding to an average interfacial area of $166 \pm 4 \text{ \AA}^2/\text{TODGA}$. Furthermore, the interfacial behavior of TODGA is consistent and predictable to at least 0.1 M TODGA across the range of acidities studied. TODGA solutions in contact with 0.46 M HNO_3 , which represent the specific solution compositions used in the TODGA kinetic experiments below, also show no sign of reverse micelle formation in the organic phase at high TODGA concentrations.

Mass transfer kinetics

Measurements of the Nd reaction kinetics in this system provide a complementary perspective on the extraction process. The spectra for the extraction of Nd from 2.6 M HNO_3 into 0.10 M TODGA/*n*-dodecane are shown in Fig. 4, along with the changes in absorbance observed at three of the most prominent wavelengths. Similar data for Nd extraction from 0.4 M HNO_3 also are reported in SI Fig. S4. Increasing the aqueous acidity clearly increases the amount of metal extracted into the organic phase and the rate at which the reaction reaches equilibrium. At 0.40 M HNO_3 , $D_{\text{Nd}} = 0.53 \pm 0.04$ and the extraction reached equilibrium in approximately 25 s, leading to $k_{\text{obs}} = 0.18 \pm 0.01 \text{ s}^{-1}$. At 2.6 M HNO_3 , the equilibrium distribution ratio climbed to $D_{\text{Nd}} = 1106 \pm 54$ and Nd extraction reached equilibrium within 15 s with $k_{\text{obs}} = 0.30 \pm 0.05 \text{ s}^{-1}$. The spectra for Nd(III) isolated in either the aqueous phase or in the organic phase are provided in Fig. S5 for comparison.

The spectral changes over time were first analyzed by singular value decomposition of the data into absorbance and kinetic eigenvectors using the OLIS GlobalWorks software. Under all the conditions studied, this analysis always indicated the presence of two unique light-absorbing Nd species with two significant kinetic eigenvectors. Fitting all of the absorbance data from an experiment using a simple two-species irreversible model yielded an overall pseudo first order rate constant (k_{obs}) for the transfer of Nd between the aqueous and organic phases. This observed mass transfer coefficient was decoupled into



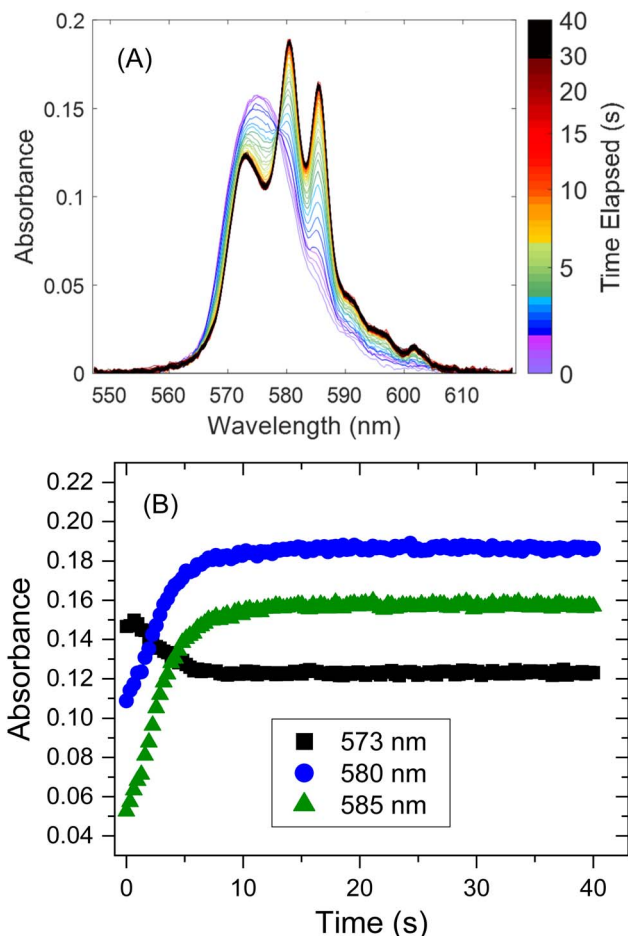


Fig. 4 Results of the *operando* UV/visible absorption spectroscopy of Nd extraction from 2.6 M HNO_3 by 0.10 M TODGA/*n*-dodecane, monitoring the hypersensitive $^4I_{9/2} \rightarrow ^4G_{5/2}, ^2G_{7/2}$ absorption bands of Nd(III) at 295 K. (A) Spectra collected from 547–619 nm over 40 s and (B) absorbance changes over time for three of the most prominent wavelengths.

a forward (extraction or aqueous \rightarrow organic) rate constant for mass transfer (k_{ao}) and a reverse (stripping or organic \rightarrow aqueous) rate constant for mass transfer (k_{oa}) using the distribution ratio, D_{Nd} , and eqn (4) and (5), where A/V is the specific interfacial area²⁴ for our systems.

$$D_{\text{Nd}} = \frac{[\text{Nd}]_{\text{eq}}}{[\text{Nd}]_{\text{aq}}} = \frac{k_{\text{ao}}}{k_{\text{oa}}} \quad (4)$$

$$k_{\text{obs}} = \frac{A}{V} (k_{\text{ao}} + k_{\text{oa}}) \quad (5)$$

The forward and reverse mass transfer rate constants derived from k_{obs} using eqn (5) were used as initial values for a two-species reversible kinetic fit of all the spectra from an experimental run in GlobalWorks, which fit the data well and yielded the final, refined k_{ao} and k_{oa} values for the Nd phase transfer reaction.

These interfacial area independent rate constants were used to determine the reaction orders in the rate law for the forward

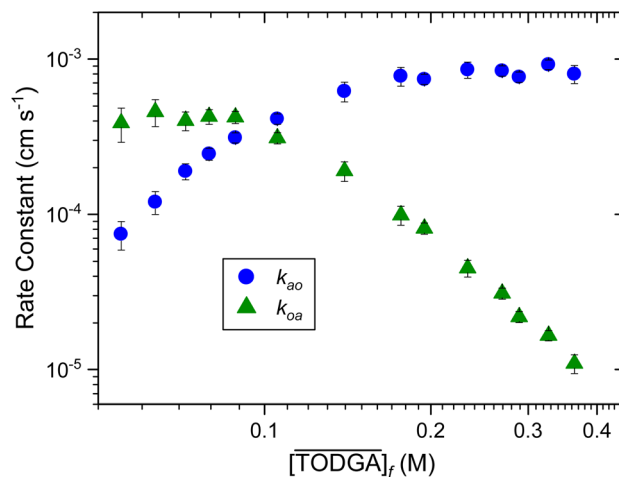


Fig. 5 Extractant dependence of the pseudo-first order rate constants of extraction (blue circles) and stripping (green triangles). The organic phase TODGA concentration varied from 0.06–0.40 M with constant 0.5 M HNO_3 and 4.9×10^{-5} M initial aqueous Nd at 295 K.

and reverse reactions.^{14,25,26} Logarithmic plots of the pseudo-first-order rate constants as functions of the concentration of free TODGA (Fig. 5) provide information about the reaction order. From the linear regions of the curves, limiting slopes of 3.0 ± 0.4 in the extraction direction and -3.1 ± 0.2 in the stripping direction are obtained. With respect to $[\text{TODGA}]_f$ and at constant $[\text{HNO}_3]$, the rate laws for the forward and reverse reactions take the general functional forms,

$$\text{rate aqueous} \rightarrow \text{organic} = k_{\text{ao}} = \frac{c_1 [\text{TODGA}]_f^3}{c_2 + c_3 [\text{TODGA}]_f^3} \quad (6)$$

and

$$\text{rate organic} \rightarrow \text{aqueous} = k_{\text{oa}} = \frac{c_4}{c_2 + c_3 [\text{TODGA}]_f^3} \quad (7)$$

The mathematical forms of the rate laws describing this solvent extraction system eqn (6)–(9) are consistent with the complex curvature and saturation at low and high concentrations observed in the $[\text{TODGA}]$ and $\{\text{NO}_3^-\}$ dependence data presented in Fig. 5 and 6. As will be discussed in more detail, this form of rate law is consistent with the interfacial two step consecutive reactions mechanism, which has been derived for metal solvent extraction systems previously.^{14,27} If the phase transfer reaction rate is controlled by chemical reactions, the c_n coefficients in eqn (6) and (7) will be composed of the appropriate rate constants for the individual elementary steps of the chemical reactions and the interfacial absorption coefficients of TODGA,¹⁴ since the interfacial tension data imply that the interface is saturated with TODGA molecules under these experimental conditions. If, on the other hand, the reaction is diffusion controlled, the values of c_n will be determined by the appropriate diffusion coefficients. The best fits of the data were obtained when parameters c_1 – c_4 and the TODGA reaction order



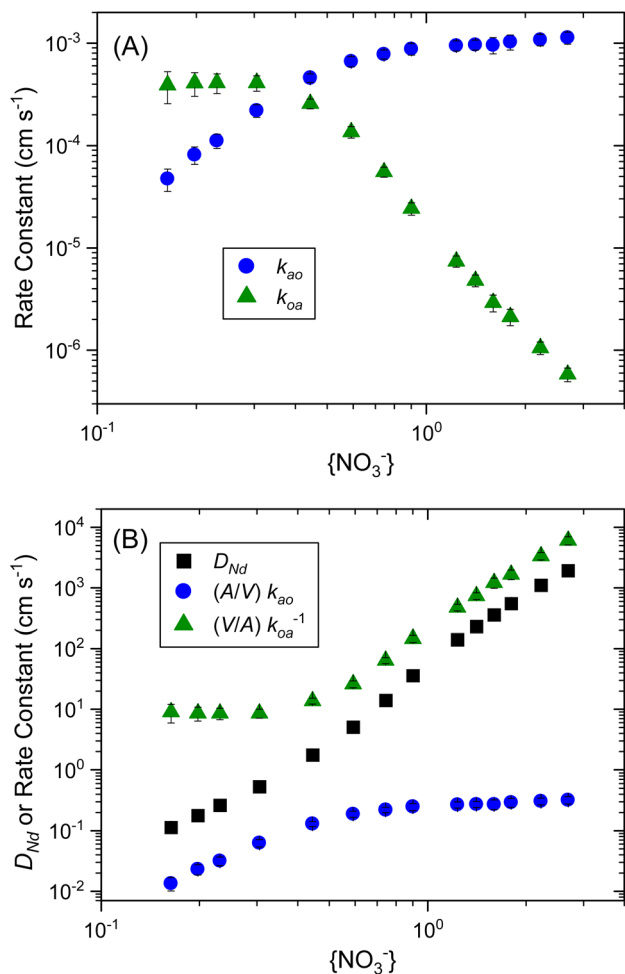


Fig. 6 Acid dependence varying the equilibrium aqueous nitrate anion activity from 0.16–2.69 (0.20–3.0 M HNO₃) with constant 0.10 M TODGA and 4.9 × 10⁻³ M initial aqueous Nd³⁺ at 295 K. (A) Pseudo-first order rate constants for extraction (blue circles) and stripping (green triangles). (B) A comparative plot of D_{Nd} , k_{ao} and k_{oa}^{-1} showing similar trends in the parameters.

were varied simultaneously. This decreased χ^2 for the fit by a factor of 2.5 and yielded a 3.25 ± 0.11 power dependence on TODGA for both the forward and reverse reaction rates. The data could also be modelled equally well assuming two parallel reaction paths involving 3 or 4 TODGA molecules. However, the presence of additional lower order TODGA-containing terms in the rate equation did not improve the fits.

The pseudo-first order kinetic results for the nitrate anion dependence are shown in Fig. 6A. They demonstrate that the extraction reactions are governed by different factors over different nitrate anion activity regimes. At low activities (*i.e.* {NO₃⁻} < 0.5 which is [HNO₃] < 0.67 M), the stripping rate constant, k_{oa} , is independent of nitrate activity, but is the primary contributor to the overall rate constant, k_{obs} , and the reaction rate over most of that range. Consequently, the extraction rate constant, k_{ao} , steadily rises over this range of nitrate activities with an effective reaction order that matches the equilibrium nitrate activity dependence, 3.03 ± 0.08 . The opposite behavior is observed in the higher nitrate activity

region (*i.e.* $0.5 < \{\text{NO}_3^-\} \leq 2.7$ or $0.67 \text{ M} < [\text{HNO}_3] \leq 3 \text{ M}$). In this range of nitrate activities k_{ao} plateaus but k_{oa} decreases steadily (Fig. 6A) with an observed reaction order of -4.73 ± 0.12 with respect to aqueous nitrate activity. The form of the rate laws with respect to the aqueous nitrate activity at constant TODGA concentration have the same general form as the TODGA-based rate expressions,

$$k_{ao} = \frac{c_5 \{\text{NO}_3^-\}^3}{c_6 + c_7 \{\text{NO}_3^-\}^3} \quad (8)$$

and

$$k_{oa} = \frac{c_8}{c_6 + c_7 \{\text{NO}_3^-\}^{4.73}} \quad (9)$$

Because k_{ao} is much larger than k_{oa} at high nitrate activities, and k_{ao} is constant, the overall mass transfer rate constant, k_{obs} , also plateaus (SI Fig. S6). Interestingly, although the kinetic constant, k_{obs} , plateaus at high nitrate activity, the thermodynamic quantity describing the overall extraction, D_{Nd} , continues to climb rapidly. Fig. 6B compares k_{oa}^{-1} and D_{Nd} to highlight the similarity between the rapid decrease in stripping rate and the large increase in D_{Nd} in the higher nitrate activity region. All of the increase in D_{Nd} with increasing nitrate activity is accounted for by the decrease in k_{oa} over the same range. Given the requirement of charge neutrality for the organic phase complexes, and the direct correspondence of the nitrate activity slopes of k_{oa}^{-1} and D_{Nd} depicted in Fig. 6B, the observed order of the stripping reaction with respect to nitrate implies the loss of all HNO₃ molecules bound to the extracted complex as well as the three NO₃⁻ in the overall stripping reaction.

Nature of the phase transfer reactions. These kinetic measurements shed light on the mechanism of the Nd(III) phase transfer reactions in the nitric acid/water/TODGA/*n*-dodecane system. In solvent extraction of metal ions, the rate controlling processes can be chemical reactions in either bulk phase, chemical reactions at or near the interface, or diffusion of the reactants or products. The key signature for extraction rates controlled by the rate of chemical reaction in the bulk phase, mass transfer rates that are independent of the interfacial area, is not easily observed with our system. However, few practical solvent extraction reactions are rate-controlled by bulk phase reactions, the solution complexation kinetics of lanthanides are often fast compared to diffusive processes,¹⁵ and TODGA has very low solubility in aqueous solutions (8×10^{-6} M in *n*-dodecane/water systems)²⁸ so bulk phase reactions between lanthanides and the extractant are not likely to be important. Instead, given the significant surface activity of TODGA indicated by the interfacial tension measurements, it appears that the rate controlling processes in this extraction system are occurring at or near the interface. This matches the conclusions of the previous kinetic studies with TODGA.^{16,18,29}

The mathematical forms of our observed rate laws are known to arise in two particular categories of solvent extraction reactions when interfacial processes dominate the rates.²⁴ For solvent extraction systems where the mass transfer kinetics is determined by the rates of chemical reactions at or near the



interface, rate laws with the forms of eqn (6)–(9) are expected for systems where the elementary steps for the adsorption and desorption of Nd^{3+} and a neodymium complex at the aqueous-organic interface are rate controlling.¹⁴ In solvent extraction, this type of reaction has been referred to as an interfacial two-step consecutive reaction (ITSCR).²⁷ However, this is not the only reaction mechanism that is consistent with eqn (6)–(9). Rate expressions with identical dependences on reactant concentrations also govern certain systems when the mass transfer rate is controlled by diffusion through the interface.^{25,27}

Solvent extraction reactions controlled by interfacial chemical reactions can be rigorously distinguished from diffusion-controlled reactions in systems where the rate can be studied as a function of the thickness of the diffusion layers, but that is not possible in our highly-stirred cell. Nevertheless, multiple lines of evidence are most consistent with partitioning rates that are controlled by the chemical reactions. First, we observe a plateau in the mass transfer rates at high stirring speeds (SI Fig. S7), which is a necessary, but not sufficient condition for chemical kinetic control of the reaction.³⁰ Second, using a rotating membrane cell with well-defined hydrodynamics, Vu *et al.*, observed that the mass transfer reactions of Am(III) by 0.2 M TODGA in 5% 1-octanol/hydrogenated tetrapropylene were substantially slower than the diffusion limit.¹⁸ Third, the average drop sizes, volume fractions, diffusion coefficients, and mass transfer rates, appear inconsistent with diffusive models that do not include slow chemical reactions. Correcting for differences in solvent viscosity, we used previously reported diffusion coefficients for Am¹⁸ to estimate the diffusion coefficient of $\text{Nd}(\text{TODGA})_3(\text{NO}_3)_3$ in *n*-dodecane to be 12.3% larger than the reported diffusion coefficient of $\text{Am}(\text{TODGA})_3(\text{NO}_3)_3$ in 5% 1-octanol/hydrogenated tetrapropylene, or $1.3 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$. The diffusion coefficient of Nd^{3+} in 0.5 M HNO_3 was taken to be $4.94 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$.¹⁸ With these values for the diffusion coefficients, the apparent thickness of the diffusion layer deduced from two-film theory³¹ appears too large in the continuous, well-stirred, aqueous phase of our system at 50–70 μm , when the average separation between the dispersed organic droplets is on the order of 150 μm . Moreover, the average surface residence time of the aqueous layer at the surface of the dispersed organic droplets predicted by Danckwartz surface renewal theory,³² 8.4 ± 0.7 seconds, appears much too long for our well-stirred reaction cell. Together these facts lead us to prefer the ITSCR model over diffusion-controlled mass transfer to explain the kinetic observations.

Mechanism of the phase transfer reactions. The presence of plateaus in our observed absolute extraction and stripping rates (Fig. 5 and 6) is consistent with a multistep mechanism composed of sequential, reversible, elementary steps,³³ which matches the nature of the ITSCR model well. The best fitting TODGA reaction order for the forward and reverse reactions, 3.25 ± 0.11 , matches the experimental average TODGA stoichiometry of the equilibrium Nd-TODGA complexes in the bulk organic phase (Fig. 2A). In this case, the non-integral reaction order implies parallel reaction paths involving 3 or 4 TODGA molecules. The slow step in the ITSCR model for this system on the organic side of the interface would be detachment

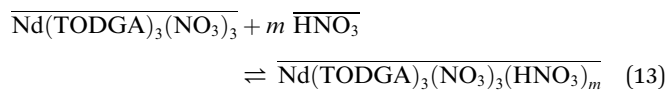
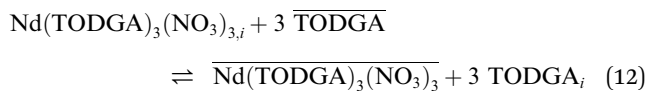
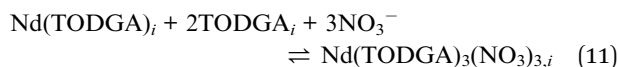
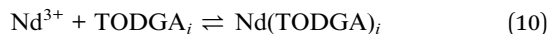
(extraction) and attachment (stripping) of fully coordinated 3 : 1 and 4 : 1 TODGA : Nd complexes at an average stoichiometry of 3.25 : 1. On the aqueous side of the interface, the other slow step would be coordination of Nd(III) to a ligand at the interface and release from the interfacial complex into the aqueous phase. Because lanthanide(III)-nitrate equilibria are very fast,³⁴ it is quite likely that the interfacial ligand that Nd binds to in this rate-limiting step is a TODGA molecule, which saturates the interface for all the solution compositions studied.

The reaction orders with respect to the nitrate present a more complicated picture of the reactions involved in the phase transfer. Three nitrates are required to form the requisite neutral Nd complex, but complexes containing 1 or 2 additional HNO_3 molecules also form in the equilibrium organic phase. Because the aqueous activity of H^+ varies directly with the activity of NO_3^- , each molecule of HNO_3 extracted should show a second power dependence of the equilibrium on $\{\text{NO}_3^-\}$ or $\{\text{H}^+\}$. This is clearly observed in the equilibrium D_{Nd} values when they are corrected for variations in the activity coefficients of Nd^{3+} , H_2O , and the formation of aqueous $\text{Nd}(\text{NO}_3)^{2+}$ and $\text{Nd}(\text{NO}_3)_2^+$, in addition to the changes in $\{\text{NO}_3^-\}$. These corrections are not easily applied to the kinetic reaction orders for the mass transfer. At low aqueous concentrations of HNO_3 , the reaction order in the extraction direction at low concentrations is 3 (eqn (8) and Fig. 6A). Combined with the TODGA dependence data discussed above, this is consistent with $\text{Nd}(\text{TODGA})_\ell(\text{NO}_3)_3$ being the complexes that detach from the interface in the extraction direction.

Different behavior is observed for the stripping reaction at higher nitrate activities. Under these conditions k_{ao} plateaus, and the stripping rate, k_{oa} , decreases with an observed order of 4.7 with respect to $\{\text{NO}_3^-\}$ without correction for changes in the activity coefficients of other aqueous species (eqn (9)). Because the reaction order for the stripping direction with respect to nitrate is greater than 3, the stripping reaction will involve dissociation of HNO_3 from the Nd complex prior to the rate determining steps. But, since the forward rate has plateaued at high nitrate activities (Fig. 6A), according to eqn (4) all the increase in D_{Nd} is associated with a strong decrease in k_{oa} (Fig. 6B), which makes the effective reaction order for k_{oa} match the nitrate dependence slope of D_{Nd} after the full activity correction, which is 6.9 at the highest acidities. Therefore, all three nitrates and all the coordinated nitric acid in the complex must be released in the stripping steps by the time the transition state forms in the final complex. This final step would then be release of Nd^{3+} from the aqueous side of the interface.

Combining these reaction orders, the rate laws imply that within an ITSCR mechanism (1) nitrate ions and TODGA molecules react with Nd adsorbed in the interface region in a series of fast elementary equilibria that occur between the rate-controlling interfacial adsorption and desorption elementary reactions, and (2) the addition of the solvating HNO_3 to the $\text{Nd}(\text{TODGA})_\ell(\text{NO}_3)_3$ complex takes place as a set of fast equilibria in the organic phase after desorption of the 1 : ℓ : 3Nd : TODGA : NO_3^- complex from the interface. For the complex containing 3 TODGA molecules the reactions would be





where the subscript i denotes interfacial species and reactions 10 and 12 are slow. Equilibria (11) and 13 represent cascades of multiple fast elementary reactions. In equilibrium 11, the sequence of NO_3^- and TODGA addition is unknown, but because nitrate dwells in the outer sphere of the $\text{Nd}(\text{TODGA})_3$ complex,⁹ three TODGA molecules likely add before the final nitrate counteranion can be accommodated in the outer sphere of the complex.

This multi-step mechanism is also consistent with the temperature dependence data for the extraction reactions. The overall extraction equilibrium (eqn (2)) is exothermic with $\Delta H^0 = -57 \pm 4 \text{ kJ mol}^{-1}$ for Nd extraction from 0.50 M HNO_3 into 0.1 M TODGA/*n*-dodecane (SI Fig. S8). However, the temperature dependencies of k_{ao} and k_{oa} show opposite trends from each other (SI Fig. S9). The forward rate slows down as the temperature increases with an apparent activation energy of $-40 \pm 2 \text{ kJ mol}^{-1}$, while the reverse rate increases with an apparent activation energy of $16 \pm 4 \text{ kJ mol}^{-1}$. The negative apparent activation energy observed for k_{ao} is consistent with mechanisms where a fast exothermic equilibrium, such as the one summarized by equilibrium (11), are followed by a slow step where the activation energy, E_{a} , for the slow step is less than $-\Delta H^0$ of the equilibrium.³⁵

Conclusions

The *operando* UV/visible spectroscopic measurement of solvent extraction kinetics under strong stirring allowed determination of the mass transfer parameters in the Nd(III)/ HNO_3 -TODGA/*n*-dodecane solvent extraction system for reactions that reached equilibrium in as little as 10 seconds. While these measurements were conducted under conditions resembling practical reaction conditions (*i.e.* TODGA and HNO_3 concentrations were varied over a practical range and efficient phase mixing was implemented), fundamental information about the extraction mechanism could be discerned. As expected for extraction of neutral nitrate salts by water-insoluble neutral ligands, the rate constants for extraction increased with a third order dependence on the concentrations of both TODGA and HNO_3 , eventually reaching plateaus at high concentrations. This is in contrast to the confounding findings of Vu *et al.*, who report Am(III) and Eu(III) extraction rates weakly decrease with increasing TODGA concentrations in a similar extraction system.¹⁸ The stripping rates showed the reverse behavior with plateaus in the TODGA and NO_3^- dependencies at low

concentrations and decreasing rates at higher concentrations. Similar decreasing stripping rates at high TODGA concentrations were also observed by Vu *et al.*¹⁸ Certain key differences exist between the current study and their investigations, such as our focus on neodymium(III), using *n*-dodecane as the organic diluent, and our employment of a highly-stirred tank technique. This differs from the study by Vu *et al.* which focused on europium(III) and americium(III), using the highly-branched hydrogenated tetrapropylene with 5 volume percent 1-octanol as the organic diluent, and a rotating membrane cell to study the kinetics. It is likely that the discrepancies between the two studies are due to a combination of these differences.

From a mechanistic perspective, the kinetic data can be interpreted according to the interfacial two-step consecutive reactions mechanism, where adsorption to and desorption from the interface of Nd species are rate-controlling. During extraction, 3 nitrate anions and 3 or 4 TODGA molecules bind to Nd at the interface in a series of equilibria before desorption of the neutral $\text{Nd}(\text{TODGA})_3(\text{NO}_3)_3$ complex from the interface into the organic phase. In the reverse process, strong inverse dependence of k_{oa} on the TODGA concentration and aqueous nitrate activity indicate that the $\text{Nd}(\text{TODGA})_3(\text{NO}_3)_3(\text{HNO}_3)_m$ complex fully dissociates, releasing Nd^{3+} to the aqueous phase in a slow step.

This kinetic picture of lanthanide extraction by TODGA from nitric acid solutions also provides an alternate interpretation of the varied and strong dependence of the equilibrium distribution ratio D_{Nd} on the nitric acid concentration. At the lowest nitric acid concentrations and nitrate activities, Nd complexes containing no nitric acid adducts are the most prevalent species in the organic phase (SI Fig. S3), k_{oa} is constant and all of the increase in D_{Nd} is attributable to the increase in the forward rate. At approximately $\{\text{NO}_3^-\} = 0.6$ (or $[\text{HNO}_3] = 0.8 \text{ M}$), the situation reverses. At that point, k_{ao} becomes constant, k_{oa} begins to decrease precipitously, and nitric acid adducts come to dominate the organic phase speciation. As illustrated in Fig. 6B, under these conditions it is the changes in k_{oa} that drive the changes in D_{Nd} , and the nitrate dependence of D_{Nd} achieves a much higher slope than it has when the forward rate determines the changes in D_{Nd} . Thus, from this perspective, the kinetic difficulty in losing all of the nitrate counter ions and nitric acid molecules from the organic phase Nd-TODGA complexes hinders the reverse reaction at high nitric acid concentrations which proportionately increases the equilibrium distribution ratio.

Author contributions

Wyatt S. Nobley – conceptualization, formal analysis, investigation, methodology, visualization, writing – original draft, writing – review and editing. Johnathan J. Klein – investigation, visualization, writing – original draft. Mark P. Jensen – conceptualization, formal analysis, supervision, funding acquisition, visualization, project administration, writing – review and editing

Conflicts of interest

There are no conflicts to declare.



Data availability

Data for this article, including the equilibrium distribution ratios is archived in the Separation Archive for Elements at <https://safe.lanl.gov/>, while interfacial tension measurements and rate constant data are given in the Supplementary Information (SI). Supplementary information is available. See DOI: <https://doi.org/10.1039/d5ra06672g>.

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