
Michael A. Bromley and Colin Boxall*

Lancaster University, Engineering, Gillow Avenue, Lancaster, LA1 4YW, U.K.

E-mail: m.bromley@lancaster.ac.uk, c.boxall@lancaster.ac.uk*

In the context of management of the waste streams arising from the reprocessing of spent nuclear fuel, studies have been conducted into the solvent extraction kinetics of lanthanide fission products by the oxygen-donor ligand N,N,N',N'-tetraoctyl diglycolamide (TODGA) and the effects of solvent phase acidity on the effectiveness of this organic extractant molecule. In this study, we observe a reduction in the rate of cerium extraction as solvent phase acidity increases, attributed to aggregation of the extractant. The effect is shown to be reversible through neutralization while a permanent prevention method is the subject of further work.

Keywords: TODGA; Rotating Diffusion Cell; Nuclear; Separation; Extraction; Reprocessing.

1. Introduction

Nuclear power is of great importance to the future of low carbon energy production and the ability to separate and recover the actinide elements from spent fuel is a key requirement for a sustainable nuclear fuel cycle. While the extraction of U and Pu for the fabrication of new fuel is well established with the PUREX process [1], recovery of the minor actinides, and particularly their separation from the chemically similar lanthanides, remains challenging.

A range of organic extractant molecules, such as N,N,N',N'-tetraoctyl diglycolamide (TODGA), have been developed for the recovery of trivalent actinides through solvent extraction processes with particular aims to achieve effective recycling of minor actinides from long-lived radioactive waste. Consequently, it is important to develop a comprehensive understanding of the extractant molecules and diluents as well as the associated chemical extraction mechanisms and kinetics involved in such processes. A more complete understanding will enable future process optimisation and assessment of the long-term suitability of proposed extractants and their potential for regeneration / recycling in order to minimize process waste and cost.

As such, studies of the interfacial and mass transport kinetics of the extraction of trivalent lanthanide fission products (as minor actinide surrogates) from aqueous HNO₃ solutions by the organic extractant molecule TODGA in organic solvent systems, such as dodecane, have been ongoing in our laboratories with the use of a rotating diffusion cell (RDC) [2].

The RDC is a compartmental diffusion apparatus consisting of two solution vessels separated by a Millipore GSWP04700 nitrocellulose membrane filter, Figure 1. This creates a defined solution phase interface across which the transport of species may be studied.

Rotation of the inner RDC vessel generates hydrodynamic flow conditions at each side of the interface membrane [2]; at the outer membrane surface, these conditions closely match those of widely studied rotating disc systems while in the inner vessel, a flow pattern towards and normal to the interface is maintained via a static baffle, Figure 1.
Establishment of a robust and controllable hydrodynamic environment at each side of the solution phase interface enables the study of interfacial kinetics and the interrogation of several kinetic regimes from mass transport to thermodynamic control as well as the transition between the two [3].

Through use of this technique, we have quantified the interfacial kinetics of the extraction of Ce(III) by TODGA [4] and have identified that the key Ce-TODGA complexation reaction takes place in the aqueous solution phase close to the aqueous-organic interface [5]. Further observations include the linear dependence of the rate of Ce extraction on the aqueous concentrations of both Ce(III) and HNO$_3$ (not shown), indicating the inclusion of a single nitrate ion in the first coordination sphere of the Ce(III)-TODGA complex; and, through variation of both hydrodynamic conditions and key solution concentrations, that the system operates under mixed mass transfer with chemical reaction (MTWCR) control [6]. These observations are in agreement with concurrent studies by Lelias et al and Simonin et al [4].

We also observe a reduction in the extraction kinetics of the Ce(III) / TODGA system where the organic solution phase is acidified both as a consequence of pre-contacting of the organic solution phase with an acidic aqueous phase, and from the increase in organic phase acidity due to simultaneous extraction of HNO$_3$ with Ce(III) by TODGA within the RDC.

Given the acidic environments associated with solvent extraction processes, further investigation into the effects of solvent phase acidity on the extraction properties of TODGA form the basis for this paper.

2. Experimental

All reagents used are AnaLaR grade or higher, and purchased from Sigma-Aldrich (Dorset, UK) or Alfa Aesar (Lancashire, UK) with the exception of TODGA (Technocomm Ltd., Falkland, UK). All water used is ultrapure doubly de-ionized water from a Direct-Q 3 UV Millipore water purification system (Millipore, Watford, UK) to a resistivity of 18.2 MΩ·cm.

RDC studies are performed with a freshly prepared aqueous solution phase of 10 mmol/dm$^3$ Ce(III) N$_2$O$_5$ in 1 mol/dm$^3$ HNO$_3$ in the outer vessel and an organic solution phase of 0.2 mol/dm$^3$ TODGA in n-dodecane with 5 % vol 1-octanol in the inner vessel. Where organic solution phases are to be used in a non-equilibrated state, no prior contact is made with the aqueous phase. Where the organic phase is to be pre-equilibrated, contacting is performed via 15 min shaking in an Eppendorf ThermoMixer C (Eppendorf UK Ltd, Stevenage, UK) and 5 min centrifugation in a MicroStar 12 centrifuge (VWR, West Sussex, UK).

In the RDC, the nitrocellulose membrane is fully permeated with the organic solution phase, ensuring that the phase boundary is positioned at the outer membrane surface while equal solution levels in each vessel avoid any hydrostatic pressure effects across the interface. The RDC is then operated at constant rotation speed of 2 Hz for up to 240 min with monitoring of each solution phase performed using a Gamry Spectro-115U UV-Vis spectrophotometer (Gamry Instruments, Cheshire, UK).

Validation of spectrophotometric measurement of [Ce(III)] in the organic solvent phase was performed by contacting fresh organic phase solutions with a range of known [Ce(III)] aqueous solutions. The organic phase absorbance at 346 nm was plotted against the quantified depletion of Ce(III) in the aqueous phase, as determined via both spectrophotometry and via ion chromatography with conductivity detection, Figure 2. These plots enable molar extinction coefficients for [Ce(III)]$_{org}$ of 0.0090 m$^2$/mol and 0.0094 m$^2$/mol to be determined respectively.

3. Discussion

Previous RDC studies in our laboratory of the kinetics of extraction of Ce(III) by the TODGA / dodecane system have demonstrated the system to be under MTWCR (mass transport with chemical reaction) control, with rate of extraction being controlled by (i) partitioning of the TODGA into the aqueous phase and (ii) subsequent Ce-TODGA complexation close to the aqueous / organic interface.

More recent RDC experiments with non-equilibrated TODGA, i.e. TODGA that has not been previously contacted / equilibrated with HNO$_3$ demonstrate the simultaneous extraction of both Ce(III) and HNO$_3$. Relative changes in species concentration in the organic solution phase are detected via UV-Vis spectrophotometry with HNO$_3$ represented by an increase in UV-Vis absorbance at a $\lambda_{max}$ of 300 nm and Ce(III) is represented by an increase in UV-Vis absorbance at a $\lambda_{max}$ of 346 nm, Figure 3.

While extraction of both species is seen to occur from the onset, a clear difference between the two processes is apparent. Organic HNO$_3$ concentration increases with a linear trend vs time for the duration of the RDC experiments, indicating a constant rate of HNO$_3$ extraction by TODGA with no evidence of saturation within the experimental period. In contrast, organic Ce(III) concentration increases at an initially steady rate at $t = < 60$ min, but begins to plateau as the rate of Ce extraction slows significantly at $t = > 60$ min.
This marked change in the rate of Ce(III) extraction is not attributable to complete extraction or to having reached the loading capacity of the extractant molecule as significant depletion of the concentration of Ce(III) in the aqueous phase is not observed and the quantity of TODGA in the RDC system should capably extract all available Ce(III) [7]. Rather, we postulate that the efficiency of the extraction process is inhibited by the increasing organic phase acidity due to extraction of HNO₃ from the aqueous solution phase resulting in micellization of the organic extractant molecule.

A number of recent papers discuss the aggregation of TODGA into tetrameric reverse micelles upon contact of the organic phase with HNO₃ reporting a critical micellization point at HNO₃ concentrations of ≥ 0.7 mol/dm³ [8-10]. Our studies demonstrate that a similar organic phase acidity is achieved in the RDC system at t = ~60 min through the linear extraction of the 1.0 mol/dm³ HNO₃ aqueous solution phase by TODGA. The resulting aggregation of the extractant molecule and the associated increase in solvent phase viscosity reported in our previous communication [11], negatively impact the mass transfer of the system - causing the plateau in the rate of Ce(III) extraction shown in Figure 3.

To further investigate the effect of solvent acidity on the rates of Ce(III) and HNO₃ extraction, RDC experiments were conducted using TODGA which was pre-equilibrated with 1.0 mol/dm³ HNO₃ and TODGA which was pre-equilibrated with HNO₃ and then repeatedly washed with 0.1 mol/dm³ NaOH in order to neutralize the acidity of the organic phase.

In the case of HNO₃ extraction, it was observed that very little acid is extracted by the 1 mol/dm³ HNO₃-equilibrated TODGA, as may be expected considering the solvent phase is already at equilibrium with the acidic aqueous phase. In contrast, the NaOH-washed TODGA extracts HNO₃ at a rate comparable to that of non-equilibrated TODGA, indicating that the solvent phase acidity established through acidic pre-equilibration has been neutralized, Figure 4.

In the case of Ce(III) extraction, the influence of acidifying the solvent phase is again demonstrated as the 1 mol/dm³ HNO₃-equilibrated TODGA shows a significant reduction in the initial rate of Ce(III) extraction when compared to fresh, non-equilibrated TODGA, Figure 5. At t = 60 min, 1.49 mmol/dm³ of Ce(III) is detected in the non-equilibrated solvent phase while in the HNO₃-equilibrated solvent phase, this is reduced by 77.85% to 0.33 mmol/dm³ of extracted Ce(III). At t = > 60 min, the rate of extraction by the non-equilibrated TODGA becomes comparable to that of the HNO₃-equilibrated TODGA which continues relatively linearly, extracting a further 0.32 mmol/dm³ of Ce(III) by t = 120 min. This supports the theory that the observed plateau in Ce(III) extraction can be attributed to a critical acidity level in the organic solvent phase.

Neutralization of the acidified solvent by NaOH-wash improves the initial extraction of Ce(III), extracting 1.88 mol/dm³ of Ce(III) by t = 60 min at a rate greater than that of the non-equilibrated TODGA. This clearly demonstrates a reversal of the effects of HNO₃ pre-equilibration and may also indicate the removal of a small amount of residual acidity in the TODGA remaining from synthesis. However, as seen in Figures 4 & 5, the simultaneous extraction of HNO₃ cannot be avoided and the in-process reduction in the rate of Ce(III) extraction still occurs as solvent phase acidity returns.

It is clear that the solvent phase acidity, either through pre-contacting with HNO₃ or through simultaneous HNO₃ extraction, influences the kinetics of Ce(III) extraction by TODGA in the RDC system. We conclude that, in the context of the MTWCR mechanism described above, HNO₃ induced aggregation of the
TODGA results in inhibition of both the diffusive mass transport of TODGA through the RDC interface membrane into the aqueous solution phase, and the key complexation reaction believed to take place there. While effects of HNO\textsubscript{3} pre-equilibration are reversible, TODGA remains susceptible to the in-process acidic environments associated with solvent extraction.

4. Conclusion

Comparison of RDC experiments conducted using non-equilibrated & HNO\textsubscript{3}-equilibrated TODGA reveals that the rate of Ce(III) extraction is significantly reduced by the increased solvent phase acidity in the pre-equilibrated experiments as both ligand mass transport to the aqueous phase and subsequent metal-ligand complexation within the system are inhibited by aggregation of the extractant molecule.

While this effect is shown to be reversible through neutralization of the solvent phase prior to use in the RDC, the simultaneous extraction of HNO\textsubscript{3} by TODGA means that solvent phase acidity increases in-process until a critical acidity level is reached and extractant aggregation occurs in non-equilibrated and in NaOH-neutralized TODGA.

In both the pre-equilibrated case and in non-equilibrated and neutralized cases at long extraction times, the net result is a reduction in the Ce(III) extraction kinetics and continued extraction at a sub-optimal rate. As there is no reason to expect that the thermodynamics of the Ce(III) / TODGA system would be changed then, given adequate time, the distribution values in each case would remain constant though further work may be required in this regard.

This study offers important information regarding the optimal use of TODGA in nuclear waste management processes where the extractant is expected to perform under acidic conditions. Clearly, in an acidic solvent phase, the extraction kinetics of TODGA are non-optimal and may reduce overall process efficiency.

Furthermore, the viscosity increase and change in surface tension associated with solvent phase acidity and extractant aggregation may influence the efficiency of other plants, such as centrifugal contactors, where a change in droplet size and viscous drag could again result in a reduction in efficiency of the system. In any reprocessing system, it is desirable to operate in the most effective and efficient conditions possible in order that the time, energy consumption and cost may be kept to a minimum and hence further investigation into the effects of solvent acidity on solvent extraction kinetics are warranted.

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