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The synergic effect of neutral organophosphorus ligands combined with acidic β -diketones for the extraction and separation of trivalent actinides

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Separating trivalent f-block elements remains a central challenge due to their similar ionic radii and chemical behaviors. Historically, these separations have been achieved using single extractants, either alone or in combination with ion exchange chromatography. However, recent studies, including this work, have explored the potential of using synergic combinations of multiple extractants to enhance extraction and separation efficiencies for trivalent actinide separations. This study investigated synergic solvent extraction (SX) systems for extracting and separating americium and curium using three neutral organophosphorus ligands: octyl (phenyl)-N,N-diisobutylcarbamoylmethylphosphine (CMPO), dibutyl N,N-diethylcarbamylmethylenephosphonate (DBDECMP), and dihexyl N,N-diethylcarbamylmethylenephosphonate (DHDECMP), combined with either 2-thenoyltrifluoroacetone (HTTA, $pK_a = 6.25$) or 4-benzoyl-3-methyl-1-phenyl-2-pyrazolin-5-one (HP, $pK_a = 3.95$). Distribution ratios (D) were determined for ²⁴¹Am³⁺ and ²⁴²Cm³⁺ as functions of nitric acid pH using 1,2-dichloroethane as the solvent. The combination of these ligands resulted in varying degrees of synergy as demonstrated by their synergic extraction enhancement coefficients (SEC). A maximum separation factor (SF $_{Am/Cm}$) of 2.65 \pm 0.21 was achieved with 0.05 M HTTA and 0.05 M DBDECMP at pH 2.50. This extractant combination was impregnated into an inert macroporous support at various ligand ratios using rotary evaporator methods to produce novel extraction chromatographic (EXC) resins. Various parameters affecting the adsorption of ²⁴¹Am³⁺ and ²⁴²Cm³⁺ onto EXC resins, such as solution pH, ionic strength, contact time, γ-irradiation dose, and temperature, were studied. Metal extraction and synergism were retained upon conversion to EXC resins, with increasing extraction observed at higher pH levels. Thermodynamic studies showed increased adsorption and decreased Gibbs free energy (ΔG) with rising temperature. Kinetic investigations indicated rapid and consistent uptake after 10 minutes. The EXC resins exhibited excellent metal retention in preliminary column experiments, demonstrating a promising potential to separate americium and curium with a maximum decontamination factor of 88. Overall, this work successfully demonstrated the identification and conversion of synergic SX systems into novel synergic EXC resins for adjacent trivalent actinide separations

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1. Introduction

Trivalent f-block separations have applications across many disciplines, including nuclear fuel reprocessing, nuclear forensics, stockpile stewardship, and nuclear medicine.^{1–4} Nuclear power remains an integral option in meeting the increasing demand for clean energy. The United States currently operates

on a once-through fuel cycle where, after irradiation, the Used Nuclear Fuel (UNF) is destined to decay in a deep geologic repository. Currently, there are no approved repositories in the United States. An alternative to the long-term repository option is recycling UNF; however, one of the main challenges is the High-level Radioactive Liquid Waste (HLW) generated during the reprocessing of UNF. UNF contains a challenging matrix of uranium, plutonium, minor actinides (MA) (Np, Am, and Cm), lanthanides, and fission products. One option for managing the high activity and long-lived nuclear waste is a closed fuel cycle, where partitioning and transmutation are used to recycle valuable uranium and plutonium resources, while addressing the minor actinides' long-term radiotoxicity and storage concerns. 5 While comprising the smallest proportion

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Dalton Transactions Paper

of UNF by mass, the MA significantly drive long-term radiotoxicity and heat load.6 A variety of well-established separation processes have been developed to recover and isolate uranium (Uranium Extraction, UREX), strontium/cesium (Fission Product Extraction, FPEX), neptunium/plutonium (Neptunium-Plutonium Extraction, NPEX), remaining fission products (Transuranic Extraction, TRUEX), and trivalent actinides/lanthanides (Trivalent Actinide Lanthanide Separation with Phosphorus-Reagent Extraction from Aqueous Komplexes, TALSPEAK, and Actinide Lanthanide Separation Process, ALSEP).7-10 In order to effectively transmute and recycle Am into shorter lived radionuclides, a further separation of the remaining Am and Cm stream is required; however, an appropriate separation method still needs to be developed as the current processes put Am and Cm into the same stream.

The trivalent actinides (An³⁺) are analogous to the trivalent lanthanides (Ln³⁺) with similar size and charge. Due to similar chemical properties within each series, inter- and intragroup separations of trivalent f-block elements have been a longstanding challenge.11 The challenge in separating americium and curium lies in the trivalent oxidation state that dominates the chemical species in conjunction with similar bonding characteristics and ionic radii. 12 The slight decrease in ionic radius with increasing atomic number is often exploited to achieve a separation of Am(III) and Cm(III). Trivalent actinide separations have been explored using a variety of radiochemical techniques such as ion-exchange chromatography, extraction chromatography (EXC), selective oxidation, and solvent extractions (SX). 11,13-15 SX has been studied extensively in the separation of actinides. 16 In the simplest SX separations, the dissolved actinide is transferred from an aqueous phase to an immiscible organic phase through ligand coordination, with the charge balance maintained to ensure that a neutral actinide complex is extracted.

Previously, neutral bifunctional organophosphorus extractants such as octyl (phenyl)-N,N-diisobutyl carbamoylmethylphosphine (CMPO), dibutyl N,N-diethylcarbamylmethylenephosphonate (DBDECMP), and dihexyl N,N-diethylcarbamylmethylenephosphonate (DHDECMP), have been used alone as efficient extractants of trivalent actinides from acidic solutions.17-21 Other types of extractants, such as acidic β-diketones like 2-thenoyltrifluoroacetone (HTTA, $pK_a = 6.25$) and 4-benzoyl-3-methyl-1-phenyl-2-pyrazolin-5-one (HP, pK_a 3.95), have also been extensively applied to trivalent f-block separations.22-25

Recently, our group reported synergic SX of Gd(III) and Tb (III) with neutral bifunctional organophosphorus (CMPO, DBDECMP, or DHDECMP) and acidic β-diketones (HTTA or HP) extractants (Fig. 1), as a promising technique to enhance extraction and/or separation selectivity of adjacent lanthanides. Several works have explored and characterized the synergism observed when combining bifunctional neutral and acidic extractants, such as CMPO, DBDECMP, DHDECMP, HTTA, and HP for f-element extraction and inter-f-elements separations, but seldom report on the synergic separation of adjacent trivalent actinides.26-31 Numerous studies have proposed the potential utility of synergic extractions using a variety of extractant types for separating f-elements. 27,32-37 However, further research is necessary to explore the potential of synergic separations for adjacent actinides using the proposed extractant combinations. This work looks to extend synergic SX techniques to enhance extraction and separation selectivity of Am(III) and Cm(III).

In terms of enhanced extraction, synergism is observed when the combination of extractants results in a distribution ratio (D) that is greater than the sum of the ratios for the individual components (eqn (1)). Conversely, antagonism occurs when the combined effect of the extractants results in a D that is less than the sum of the individual components (eqn (2)). This indicates that the combination is less effective than the individual extractants working independently. When the combined D equals the sum of the individual D (eqn (3)), the effect is neither synergic nor antagonistic, suggesting that the extractants function independently without enhancing each other's performance. Regarding selectivity, synergism is defined when the combined extractants produce a larger separation factor $(SF_{(1,2)})$ than either of the individual components $(SF_{(1)})$ or SF(2) or when they significantly improve separation efficiency in scenarios where previous extraction was ineffective.

$$D_{(1,2)} > D_{(1)} + D_{(2)} \tag{1}$$

$$D_{(1,2)} < D_{(1)} + D_{(2)} \tag{2}$$

$$D_{(1,2)} = D_{(1)} + D_{(2)} \tag{3}$$

EXC is a popular separation technique that combines the high selectivity of SX with the potential for straightforward automation, while also eliminating the generation of mixed liquid organic waste through its column-based chromatography procedures.38 EXC is a form of reverse-phase partition chromatography. In this work, the stationary phase is nonpolar and comprised of an inert support in which a single or mixture of extractants, modifiers, and/or solvents are physically adsorbed inside the pores of the support. Modifying the mobile phase's acidity and/or including complexing reagents alters the retention affinity of each element differently, facilitating the separation between adjacent elements. SX investigations are often the first step in finding potential extractants for EXC materials.

Effective SX systems with promising separation characteristics can be readily transformed into EXC resins by impregnating an inert support with the extractant system. The preparation and utilization of EXC resins have been thoroughly showcased in the literature by numerous commercial applications. 39-43 While synergism in SX has been well investigated, the extension of synergism to EXC resins has rarely been studied. 44-46 In 1970, Aly et al. demonstrated one of the first examples of synergism in extraction chromatography using HTTA and DBDECP (dibutyl diethyl carbamoyl phosphonate). In their study, when HTTA and DBDECP were sorbed on Celite to produce an EXC resin, a separation factor of 2.06 was

$$H_{3}C$$
 CH_{3}
 $H_{3}C$
 CH_{3}
 $H_{3}C$
 CH_{3}
 $H_{4}C$
 CH_{3}
 $H_{5}C$
 CH_{5}
 $H_{5}C$
 CH_{7}
 CH_{7}
 CH_{8}
 CH_{9}
 C

Fig. 1 Chemical structures of extractants used in this work. (1) octyl (phenyl)-*N*,*N*-diisobutyl carbamoylmethylphosphine, (CMPO), (2) dibutyl *N*,*N*-diethylcarbamylmethylenephosphonate (DBDECMP), (3) dihexyl *N*,*N*-diethylcarbamylmethylenephosphonate (DHDECMP), (4) 2-thenoyltrifluoroacetone (HTTA), and (5) 4-benzoyl-3-methyl-1-phenyl-5-pyrazolone (HP).

obtained for the adjacent lanthanides, samarium, and europium, compared to a <1 separation factor with either HTTA or DBDECP. 47,48 More research is needed on the systematic study of characteristics and translation of synergism between SX and EXC resins. The work presented here aimed to identify and characterize synergism in the SX of Am(III) and Cm(III) by combining neutral organophosphorus compounds (CMPO, DBDECMP, or DHDECMP) with β -diketones (HTTA or HP). The extractant combination yielding the best separation characteristics was later converted to novel synergic EXC resins to investigate selectivity in chromatography columns. By studying the conversion, characterization, and optimization of synergic systems, we can design more efficient and selective methods for adjacent Am(III) and Cm(III) separations.

Materials and methods

2.1. Reagents and materials

Concentrated nitric acidic (TraceMetalTM grade, Fisher ChemicalTM) was used to prepare HNO₃ solutions from pH 0.00–3.00 using volumetric flasks. The pH of each solution was measured with an OrionTM StarTM A211 (Thermo Scientific) pH meter and OrionTM PerpHecTTM ROSSTM Combination pH micro electrode (Thermo Scientific) calibrated with pH 1.68, 4.01, and 7.00 buffers. The pH of each solution was adjusted using standardized 1.0 M NaOH (Fisher ChemicalTM) to ±0.02 units of the desired pH. 1,2 − Dichloroethane (99.8%, Sigma-Aldrich) was used as an organic diluent during solvent extraction. Dodecane (≥99%, TCI America) and methanol (99%, Sigma-Aldrich) were used during resin preparation. 4-Benzoyl-3-methyl-1-phenyl-2-pyrazolin-5-one (HP) (99%, AA Blocks) and

2-thenoyltrifluoroacetone (HTTA) (99%, Sigma-Aldrich) were used as the β-diketone extractants, while N,N-diisobutyl-2-(octylphenylphosphoryl)acetamide (CMPO) (95%, AmBeed), dibutyl N,N-diethylcarbamoylmethylphosphonate (DBDECMP) (97.1%, TCI America), and dihexyl N,N-diethylcarbamylmethylenephosphonate (DHDECMP) (92.7%, TCI America) were used as the neutral organophosphorus extractants. Pre-filter resin (20-50 µm) was purchased from TrisKem International. Sodium nitrate (99%, Sigma-Aldrich) was used to control ionic strength. All reagents were used as received without any additional purification. The radionuclides ²⁴¹Am and ²⁴²Cm were supplied from Lawrence Livermore National Laboratory stocks and dissolved in HNO₃ to yield stocks containing ≈555 dpm 241 Am $^{3+}$ per μ L (1.0 M HNO₃) and \approx 82 dpm 242 Cm $^{3+}$ per μL (0.1 M HNO₃), respectively. The presence of ²³⁸Pu in the ²⁴²Cm³⁺ stock was verified by alpha spectroscopy, and it was found to constitute less than 0.6% of the total activity. All solutions and samples were prepared using Milli-Q® water (Millipore, USA, resistivity >18.2 M Ω cm⁻¹).

2.2. Actinide solvent extraction with synergic systems

The distribution ratios (*D*) of Am(III) and Cm(III) were measured using a standard solvent extraction technique, involving the equilibration of equal volumes of an organic phase and an aqueous phase, followed by phase separation and analysis of metal concentrations in each phase. Diluted $^{241}\text{Am}^{3+}$ and $^{242}\text{Cm}^{3+}$ stocks were prepared before each experiment such that the nominal activity was $\approx\!15$ dpm $^{241}\text{Am}^{3+}$ per μL and $\approx\!7.5$ dpm $^{242}\text{Cm}^{3+}$ per μL , respectively. Each dilute stock was adjusted to ± 0.02 units of the desired pH, as described above.

Weighed amounts of each neutral and β -diketone extractant were dissolved in 1,2-dichloroethane to form 0.2 M stocks by

Paper **Dalton Transactions**

mixing at 1500 RPM for 15 minutes using a digital vortex. Organic extraction solutions were prepared using the stocks such that each extractant had a final concentration of 0.05 M. Extraction solutions were made with each extractant (CMPO, DBDECMP, DHDECMP, HTTA, and HP) alone and combined (CMPO + HTTA, CMPO + HP, DBDECMP + HTTA, DBDECMP + HP, DHDECMP + HTTA, and DHDECMP + HP). All organic extraction solutions were pre-equilibrated with equal volumes of nitric acid at each respective pH value by mixing at 1500 RPM for 30 minutes using a digital vortex, followed by centrifugation at 4500 RPM for 5 minutes. Equal volumes of 500 µL of pre-equilibrated organic phase and adjusted pH diluted metal stocks were mixed at 2000 RPM for 30 minutes at 22 ± 1 °C using a digital vortex, followed by centrifugation at 6000 RPM for 30 seconds.

The extracted aqueous phase was separated from the organic phase and retained, and an aliquot (50-250 µL) was taken for analysis. For extraction with 241Am3+, individual extractants were analyzed in duplicate, while combined extractant solutions were analyzed in triplicate. Conversely, all ²⁴²Cm³⁺ samples were analyzed in duplicate. The ²⁴¹Am³⁺ and ²⁴²Cm³⁺ activity was determined using liquid scintillation counting (LSC) with Ultima GoldTM AB scintillation cocktail and Tri-Carb 3110TR LSC analyzer (PerkinElmer). The aqueous samples were each counted for 120 minutes or until a 1% counting uncertainty was attained.

The SX distribution ratios (D) (eqn (4)) were calculated as the ratio of activity between the organic ([An]_{org}) and aqueous ([An]aq) phases.

$$D = \frac{[An]_{org}}{[An]_{aq}} = \frac{[An]_{i, aq} - [An]_{f, aq}}{[An]_{f, aq}}$$
(4)

The activity in the organic phase was determined as the difference between the initial and final activities. The initial activity was determined from aliquots (50 µL) of standards from the dilute metal stocks. All errors on the distribution ratios are reported as one standard deviation between replicates or the propagated uncertainty in counts per minute (CPM), whichever is greater.

The synergic enhancement coefficient (SEC) that describes the change in distribution ratios between the single extractants and combined extractant system is found by the following equation:

$$SEC = Log_{10} \left[\frac{D_{(1,2)}}{(D_{(1)} + D_{(2)})} \right]$$
 (5)

where $D_{(1)}$, $D_{(2)}$, and $D_{(1,2)}$ are the distribution ratios of individual extractants and combined system, respectively. Values of SEC > 0 indicate a synergic interaction, whereas SEC < 0 indicates an antagonistic interaction and SEC = 0 represents no effect on extraction when combined. For individual extractants at pH values with negligible Am extraction, a limit was defined as two times the minimum detectable activity at a 95% confidence interval (MDA₉₅).

The separation factors (SF_{Am/Cm}) (eqn (6)) for Am(III) and Cm(III) were calculated as the ratio of individual SX distribution ratios obtained from liquid scintillation counting analysis.

$$SF_{Am/Cm} = \frac{D_{(Am)}}{D_{(Cm)}} \tag{6}$$

where $D_{\text{(Am)}}$ and $D_{\text{(Cm)}}$ represent the D of ²⁴¹Am³⁺ and ²⁴²Cm³⁺, respectively.

2.3. Optimization of separation behavior for Am/Cm using HTTA-DBDECMP solvent extraction systems

The synergic SX system, utilizing HTTA and DBDECMP, exhibited the most promising separation factors at pH 2.50 and was thus chosen for further optimization. In these studies, the concentration of DBDECMP remained constant at 0.05 M, while the concentration of HTTA varied from 0.0125, 0.025, 0.075, to 0.150 M. Additionally, a separate optimization was conducted by maintaining [HTTA] at 0.05 M while varying the concentration of DBDECMP from 0.0125, 0.025, 0.075, to 0.150 M.

2.4. Preparation of HTTA and/or DBDECMP extraction chromatographic (EXC) resins

Based on the findings of the solvent extraction investigation and optimization, the combination of HTTA and DBDECMP at 0.05 M each yielded the highest SF_{Am/Cm} at pH 2.50. This concentration ratio was converted to a mole ratio (g of diketone (R_n) per g of neutral (R_d) by multiplication of the respective molecular mass.

An extractant loading (EL) of 30% (0.3 g of total extractant per g of inert support) was used for all resin preparations. The mass (g) of each extractant for a desired mass of inert support (g) was calculated from eqn (7).

$$(T) \cdot (EL) = R_{d}x + R_{n}x \tag{7}$$

where T is the desired mass of inert support (g), EL extractant loading fraction, R_d is the diketone extractant mass ratio, R_n is the neutral extractant mass ratio, and x is the total mass (g) of extractant for T and EL parameters.

EXC resins were prepared with 10 wt% dodecane to enhance the retention of the ligands in the inert support. A maximum organic loading of 40 wt% was used based on resin manufacturers' recommendations. For resin preparation, appropriate masses of each extractant were dissolved separately in 5 mL of 1,2-dichloroethane or methanol and mixed (15 minutes at 2500 RPM, digital vortex). Both organic extractant solutions were transferred to a conical tube containing dodecane and mixed (30 minutes at 2500 RPM, digital vortex). Each extractant tube was rinsed with one, 1 mL fractions of 1,2-dichloroethane or methanol.

The corresponding mass of inert support was slurried with a 30% (v/v) methanol and water solution at liquid-to-solid ratio of 5 mL per g of support. The combined extractant/dodecane solution was transferred to the slurried resin and rinsed with two, 2 mL fractions of 1,2-dichloroethane or methanol. The

Dalton Transactions Paper

resin slurry and extractant/dodecane solution were transferred to a 500 mL round-bottom flask and rinsed with two, 5 mL fractions of 30% (v/v) methanol/water. The bulk solvent was removed under vacuum (50-75 Torr) at 45 °C. As the resin approached visual dryness (≈1 hour), the vacuum was decreased (100 Torr) to prevent bumping and continue residual solvent removal. After bumping was no longer observed (≈1 hour), the maximum vacuum (20 Torr) was drawn and held overnight (12 hours). This procedure was repeated for each mass ratio and solvent type studied. Individual extractant EXC resins were prepared in a similar manner. These materials are referred to hereafter as HTTA-DBDECMP resin 1-5. Table 1 summarizes the naming convention, mass ratios, and relevant characteristics for the EXC resins prepared.

2.5. Determination of the weight distribution ratio (D_w) for **HTTA-DBDECMP** resins

The weight distribution ratio, D_w (mL g⁻¹), of Am(III) and Cm (III) were measured on HTTA-DBDECMP resins 1-5 from pH 1.50-2.50 solutions of nitric acid via batch extraction experiments. The $D_{\rm w}$ values were calculated from the following equation:

$$D_{\mathbf{w}} = \frac{[\mathbf{Metal}]_{\text{resin}}}{[\mathbf{Metal}]_{\text{aq}}} = \left(\frac{C_{i} - C_{\text{eq}}}{C_{\text{eq}}}\right) \times \left(\frac{V}{m}\right)$$
(8)

where $[Metal]_{resin}$ and $[Metal]_{aq}$ are the equilibrium analyte concentrations on the resin, and in the aqueous phase, C_i is the initial analyte concentration of the aqueous phase, C_{eq} is the equilibrium analyte concentration of the aqueous phase, V is the volume of the aqueous phase (mL), and m is the mass of the resin (g). For batch experiments with ²⁴¹Am³⁺ and ²⁴²Cm³⁺, measured count rates replace analyte concentrations as shown in eqn (9).

$$D_{W} = \frac{[\text{Metal}]_{\text{resin}}}{[\text{Metal}]_{\text{aq}}} = \left(\frac{A_{i} - A_{\text{eq}}}{A_{\text{eq}}}\right) \times \left(\frac{V}{m}\right)$$
(9)

where A_i is the initial and A_{eq} is the equilibrium count rates (CPM) of the aqueous phase, respectively.

Microcentrifuge tubes (2.0 mL) were loaded with 50 ± 2 mg of resin and 1.0 mL of HNO₃ (pH 1.50-2.50, prepared as described above) with nominal concentrations of ≈14 dpm $^{241}\text{Am}^{3+}$ μL^{-1} and ≈ 5.6 dpm $^{242}\text{Cm}^{3+}$ per μL . These experiments analyzed each element separately and in duplicate. The tubes were incubated on a digital rotator (30 RPM for 1 hour, 26 \pm 1 °C). After equilibration, the resin was filtered using 0.2 μm PTFE syringe filters. For all resin experiments, a 250 μL aliquot of the aqueous filtrate for 241Am3+ and 242Cm3+ for each replicate was counted via LSC as described above. The aqueous samples were each counted for 120 minutes or until a 1% counting uncertainty was attained. The $D_{\rm w}$ errors for all resin experiments are reported as one standard deviation between replicates or the propagated uncertainty in CPM, whichever is greater.

2.6. Thermodynamics, kinetic, stability, and ionic strength experiments

2.6.1. Thermodynamic investigation for HTTA-DBDECMP resin 2. The $D_{\rm w}$ values for $^{241}{\rm Am}^{3+}$ on HTTA-DBDECMP resin 2 were measured at 303 \pm 1, 318 \pm 1, and 333 \pm 1 K from a nitric acid solution of pH 2.00 in triplicate. All resin samples and metal solutions were pre-incubated for 1 hour to obtain thermal equilibrium before contact.

The pre-heated solutions of ²⁴¹Am were added to each tube and incubated for 1 hour on a digital shaker (30 RPM). To calculate thermodynamic parameters of adsorption, such as change in Gibb's free energy (ΔG) , change of enthalpy (ΔH) , and change in the entropy (ΔS), the Van't Hoff equation (eqn (10)) was employed.

The Van't Hoff equation⁴⁹ can be derived from the fundamental thermodynamic relationship between Gibbs Energy and equilibrium. Through mathematical manipulation, the Van't Hoff equation relates changes in the equilibrium constant to temperature by the following equation:

$$\ln[K_{\rm eq}] = -\frac{\Delta G}{RT} = -\frac{\Delta H}{RT} + \frac{\Delta S}{R}$$
 (10)

where R is the universal gas constant (8.314 J mol⁻¹ K⁻¹), T is the temperature (K) and K_{eq} is the equilibrium constant.

The K_{eq} is a dimensionless quantity that provides a measure of the ratio of concentrations of products to reactants at equilibrium for a reversible chemical reaction. The $D_{\rm w}$ can serve as an analog to K_{eq} , reflecting the equilibrium distribution of the adsorbate between the adsorbent and the surrounding phase. From a plot of $\ln[D_{\rm w} \times \rho_{\rm wet}]$ versus the reciprocal of temperature (K), the ΔS and ΔH can be calculated. There has been some disagreement over applying the Van't Hoff equation to adsorption data because of the need to have the equilibrium constant with dimensionless units.49 To produce a dimensionless equilibrium constant, the $D_{\rm w}$ values were multiplied by the wet resin density ($\rho_{\text{wet}} = 0.704 \text{ g mL}^{-1}$). 38,42,50

2.6.2. Kinetic investigation for HTTA-DBDECMP resin 2. For kinetic investigations, the $D_{\rm w}$ values for $^{241}{
m Am}^{3+}$ on

Table 1 HTTA and/or DBDECMP extraction chromatographic (EXC) resins summary

Resin name	Mass ratio $(R_d:R_n)$	Solvent type	HTTA (g)	DBDECMP (g)	Support (g)	Extractant loading (%)	Organic loading (%)
1	1.00:0.88	1,2-DCE	0.4780	0.4210	3.0044	29.9	39.9
2	1.00:1.38	1,2-DCE	0.7564	1.0454	6.0054	30.0	40.0
3	1.00:1.38	Methanol	0.7322	1.0092	6.0010	29.0	39.0
4	0.00:1.00	Methanol	_	0.7510	2.5016	30.0	40.0
5	1.00:0.00	Methanol	0.7496	_	2.4996	30.0	40.0

Paper

HTTA-DBDECMP resin 2 were measured at 299 ± 1 K in HNO₃

HTTA-DBDECMP resin 2 were measured at 299 \pm 1 K in HNO₃ (pH 2.00) at various contact times.

2.6.3. γ -Irradiation and acid stability investigation for HTTA-DBDECMP resin 2. The $D_{\rm W}$ values for $^{241}{\rm Am}^{3+}$ on γ irradiated HTTA-DBDECMP resin 2 were measured from a nitric acid solution at pH 2.00. A polypropylene conical tube loaded with 500 mg of HTTA-DBDECMP resin 2 and 5.0 mL of HNO₃ (pH 2.00) was irradiated in the University of Utah's J.L. Shepherd Mark I, Model 30 $^{137}{\rm Cs}$ irradiator for two weeks. The estimated γ exposure (X_{γ}) was 3.818×10^7 R based on recent exposure mapping of the irradiated position. The absorbed dose $(Q_{\rm Resin})$ of 401 ± 60 kGy was calculated from X_{γ} using eqn (11).

$$Q_{\text{Resin}} = 0.88X_{\gamma} \frac{\left(\frac{\mu_{\text{en}}}{\rho}\right)_{\text{Resin}}}{\left(\frac{\mu_{\text{en}}}{\rho}\right)_{\text{Air}}}$$
(11)

where Q_{Resin} is the dose (rads) due to photons, X_{γ} is exposure is roentgen (R), $\left(\frac{\mu_{\text{en}}}{\rho}\right)_{\text{Resin}}$ is mass energy absorption co-

efficient for the resin (0.031 cm 2 g $^{-1}$), and $\left(\frac{\mu_{\rm en}}{\rho}\right)_{\rm Air}$ is mass energy absorption coefficient for the air at 661.7 keV (0.026 cm 2 g $^{-1}$), respectively.

Air and individual elemental mass energy absorption coefficient $\left(\frac{\mu_{\rm en}}{\rho}\right)$ values were interpolated from National Institute of Standards and Technology (NIST) Standard Reference Database 126. These were used to calculate $\frac{\mu_{\rm en}}{\rho_{\rm Resin}}$ based on the estimated elemental composition of the prepared resin. Due to the proprietary nature of the macroporous support, the exact elemental composition was difficult to determine. Combined with potential uncertainties in the calculation method, a 15% uncertainty is reported for $Q_{\rm Resin}$.

Alongside the γ irradiation stability test, an acid stability test was conducted. The resins were exposed to nitric acid for the same duration in both tests. After each test, the acid was decanted, and the resins were dried in a laboratory oven at 50 °C.

2.6.4. Ionic strength investigation for HTTA-DBDECMP resin 2. To investigate the influence of ionic strength on adsorption the $D_{\rm w}$ values for $^{241}{\rm Am}^{3+}$ on HTTA-DBDECMP resin 2 were measured at 299 \pm 1 K in HNO $_3$ (pH 2.00) from solutions with a total ionic strength between 0.05–1.0 M. The total ionic strength was controlled using (Na,H)NO $_3$.

2.7. Column separations with HTTA-DBDECMP resin 2

The elution profiles of Am(III) and Cm(III) on HTTA-DBDECMP resin 2 columns were investigated separately via different column types. Each column type was wet-packed with resin slurried in a nitric acid solution at pH 3.00 (60 minutes at 1500 RPM, digital vortex). For all column experiments, a 250 μ L aliquot of each fraction was counted via LSC as described above.

2.7.1. Column type 1 (inner diameter: 4.5 mm, length: 45 mm). Column type 1 had a wet-packed volume of $\approx 800 \ \mu L$ and an elution fraction size of $\approx 500 \ \mu L$. Elution experiments for both actinides were conducted concurrently, with each actinide being evaluated separately. After loading, each column was prepared by washing with two fraction volumes of pH 3.00 HNO₃, followed by loading one fraction volume of $^{241}Am^{3+}$ or $^{242}Cm^{3+}$ at pH 3.00 HNO₃.

Following metal loading, each column was washed with approximately 0.5 column volumes of pH 3.00 HNO₃, followed by an isocratic elution at pH 1.75 HNO₃. After approximately 11 column volumes, the column was stripped with approximately 0.5 column volumes of 1.0 M HNO₃.

A relative column decontamination factor for fractions between i and j, $\mathrm{DF}_{i,j}$ was determined by eqn (12) for each column type.

$$DF_{i,j} = \frac{A_{Am}(\text{initial})/A_{Cm}(\text{initial})}{\sum_{k=i}^{j} A_{Am}(\text{final})/\sum_{k=i}^{j} A_{Cm}(\text{final})}$$
(12)

where $A_{\rm Am}$ (initial) and $A_{\rm Cm}$ (initial) are the activities of $^{241}{\rm Am}^{3+}$ and $^{242}{\rm Cm}^{3+}$ initially loaded onto the column, respectively.

2.7.2. Column type 2 (inner diameter: 2.5 mm, length: 60 mm). Column type 2 had a wet-packed volume of \approx 400 μ L and an elution fraction size of \approx 300 μ L. Type 2 columns were prepared and eluted similarly to type 1, but the loading of 241 Am $^{3+}$ or 242 Cm $^{3+}$ occurred at pH 3.25 HNO₃. After 15 column volumes of isocratic elution at pH 1.75, the column was stripped with about 0.5 column volumes of 1.0 M HNO₃.

Results and discussion

3.1. Synergism in solvent extraction

The investigated pH ranges were tailored for each synergic system based on previously studied extraction efficiencies of lanthanides with these systems from pH 0.00–3.00.⁵² HTTA and DBDECMP did not display significant extraction (>5%); however, HP, DHDECMP and CMPO demonstrated varying extraction levels over their studied pH regions. HP and DHDECMP displayed increasing extraction from pH 1.50 ($D = 0.09 \pm 0.01$) to 2.00 ($D = 0.14 \pm 0.02$) and pH 2.00 ($D = 0.11 \pm 0.01$) to 3.00 ($D = 0.33 \pm 0.05$), respectively. CMPO displayed decreasing extraction from pH 0.50 ($D = 0.15 \pm 0.02$) to 0.75 ($D = 0.05 \pm 0.03$).

Synergic effects were observed for all pairings of neutral extractants with $\beta\text{-}diketones.$ Fig. 2 shows the SEC as a function of acid concentration and demonstrates that as the pH of extraction increased, the SEC value also increased. At high pH values, as much as a thousandfold increase in extraction was observed. As indicated in Fig. 2, at a constant pH of 1.50 CMPO had the largest SEC values among all three neutral ligands, followed by DHDECMP, and DBDECMP. The synergism between the neutral extractants and HP resulted in higher SEC values when compared to HTTA.

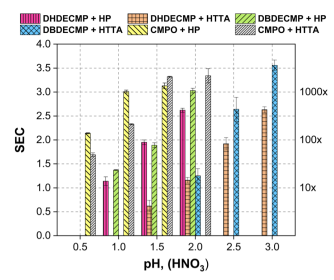


Fig. 2 Synergic enhancement coefficients (SEC) for combinations of neutral organophosphorus and β -diketone extractants as a function of pH value (HNO₃).

3.2. Solvent extraction separation behavior analysis and optimization

Tables 2–4 summarize the separation factors for each neutral and β-diketone combination. For CMPO and HTTA, separation factors were constant (\approx 1.90) across the studied pH range, whereas decreased from 1.87 to 1.19 with HP as the pH increased. For DBDECMP and HTTA, the SF_{Am/Cm} increased from 1.05 to 2.65 as pH increased, whereas with HP, decreased from 1.44 to 1.17. The separation potential with DHDECMP

was not realized, as low SF_{Am/Cm} were observed with both diketones across the studied pH regions. The highest unoptimized SF_{Am/Cm} was 2.65 \pm 0.21 at pH 2.50 with 0.05 M DBDECMP and 0.05 M HTTA with a corresponding SEC value of 2.50 \pm 0.26.

Table 5 shows the results of the SX optimization performed with DBDECMP and HTTA at pH 2.50. For the HTTA screen, the concentrations were grouped into two categories: (1) with a higher separation factor at 0.0125-0.0250 M, and (2) with a lower separation factor at 0.0750-0.1500 M. Lower concentrations of HTTA had higher SFAm/Cm than higher HTTA concentrations. Similar conclusions could not be drawn for the DBDECMP screen, where SF_{Am/Cm} was generally within experimental error. Generally, increasing the concentration of HTTA caused a higher increase in the D relative to a similar DBDECMP increase. Previous slope analysis experiments of these extractant combinations using Tb suggest that one neutral extractant is found in the extracted complex, whereas 2 diketone ligands are used. 52 This suggests that the increasing the amount of HTTA relative to DBDECMP should produce additional active ligand sites able to extractant metal. The optimization experiments suggested that the highest separation factor was obtained at 0.05 M HTTA and 0.05 M DBDECMP. These concentrations were the basis for converting the solvent extractant system to the EXC resins.

The SF_{Am/Cm} for the solvent extraction systems in this work are low but typical (generally between 2 and 3) for extractant/complexant combinations used in redox-free Am(III)/Cm(III) separations.^{53–55} While this study did not explore hydrophilic complexing agents with reverse affinity, identifying potential options could provide further improvements in separation selectivity. Although this study did not demonstrate significant

Table 2 Separation factors (SF_{Am/Cm}) for Am(III) and Cm(III) as a function of pH in HNO₃ with CMPO and HTTA or HP

	0.05 M CMPO							
	0.05 M HTTA			0.05 M HP				
pH value (HNO ₃)	$D_{\mathrm{Cm}} \pm \sigma_{\mathrm{Cm}}$	$D_{ m Am} \pm \sigma_{ m Am}$	$\mathrm{SF_{Am/Cm}} \pm \sigma_{\mathrm{SF_{Am/Cm}}}$	$D_{\mathrm{Cm}} \pm \sigma_{\mathrm{Cm}}$	$D_{ m Am} \pm \sigma_{ m Am}$	$\mathrm{SF_{Am/Cm}} \pm \sigma_{\mathrm{SF_{Am/Cm}}}$		
0.25	_	_	_	0.83 ± 0.02	1.55 ± 0.03	1.87 ± 0.06		
0.50	0.22 ± 0.03	0.43 ± 0.04	1.94 ± 0.29	1.58 ± 0.11	2.37 ± 0.08	1.50 ± 0.12		
1.00	0.94 ± 0.02	$\boldsymbol{1.83 \pm 0.05}$	1.94 ± 0.06	13.98 ± 0.34	16.70 ± 1.07	$\boldsymbol{1.19 \pm 0.08}$		
1.50	9.54 ± 0.34	17.97 ± 0.38	1.88 ± 0.08	_	_	_		

Table 3 Separation factors (SF_{Am/Cm}) for Am(III) and Cm(III) as a function of pH in HNO₃ with DBDECMP and HTTA or HP

	0.05 M DBDECMP							
	0.05 M HTTA			0.05 M HP				
pH value (HNO ₃)	$D_{\mathrm{Cm}} \pm \sigma_{\mathrm{Cm}}$	$D_{ m Am} \pm \sigma_{ m Am}$	$\mathrm{SF_{Am/Cm}} \pm \sigma_{\mathrm{SF_{Am/Cm}}}$	$D_{\mathrm{Cm}} \pm \sigma_{\mathrm{Cm}}$	$D_{ m Am} \pm \sigma_{ m Am}$	$SF_{Am/Cm} \pm \sigma_{SF_{Am/Cm}}$		
1.00	_	_	_	0.27 ± 0.02	0.39 ± 0.01	1.44 ± 0.10		
1.25	_	_	_	1.46 ± 0.02	1.80 ± 0.06	1.24 ± 0.04		
1.50	_	_	_	6.85 ± 0.14	8.04 ± 0.41	1.17 ± 0.06		
2.00	0.42 ± 0.01	0.44 ± 0.04	1.05 ± 0.10	_	_			
2.25	1.61 ± 0.02	2.25 ± 0.02	1.40 ± 0.02	_	_			
2.50	7.13 ± 0.22	18.89 ± 1.37	2.65 ± 0.21	_	_	_		

Table 4 Separation factors (SF_{Am/Cm}) for Am(III) and Cm(III) as a function of pH in HNO₃ with DHDECMP and HP or HTTA

pH value (HNO ₃)	0.05 M DHDECMP							
	0.05 M HTTA			0.05 M HP				
	$D_{\mathrm{Cm}} \pm \sigma_{\mathrm{Cm}}$	$D_{ m Am} \pm \sigma_{ m Am}$	$\mathrm{SF_{Am/Cm}} \pm \sigma_{\mathrm{SF_{Am/Cm}}}$	$D_{\mathrm{Cm}} \pm \sigma_{\mathrm{Cm}}$	$D_{ m Am}\pm\sigma_{ m Am}$	$\mathrm{SF_{Am/Cm}} \pm \sigma_{\mathrm{SF_{Am/Cm}}}$		
1.00	_	_	_	0.47 ± 0.00	0.68 ± 0.04	1.45 ± 0.08		
1.25	_	_	_	2.31 ± 0.15	2.57 ± 0.08	1.11 ± 0.08		
1.50	0.24 ± 0.03	$\boldsymbol{0.16 \pm 0.01}$	0.69 ± 0.11	9.03 ± 0.66	11.83 ± 0.28	$\boldsymbol{1.31 \pm 0.10}$		
2.00	1.77 ± 0.06	$\boldsymbol{1.78 \pm 0.16}$	1.00 ± 0.10	_	_	_		
2.50	15.67 ± 0.84	14.60 ± 3.50	0.93 ± 0.23	_	_	_		

Table 5 Optimized separation factors ($SF_{Am/Cm}$) for Am(III) and Cm(III) extraction with DBDECMP and HTTA at pH 2.50 HNO₃, as a function of DBDECMP or HTTA concentration

	DBDECMP and HTTA extraction at pH = 2.50							
[Extractant] M	0.05 M DBDECMP HTTA screen			0.05 M HTTA DBDECMP screen				
	$D_{\mathrm{Cm}} \pm \sigma_{\mathrm{Cm}}$	$D_{ m Am} \pm \sigma_{ m Am}$	$\mathrm{SF_{Am/Cm}} \pm \sigma_{\mathrm{SF_{Am/Cm}}}$	$D_{\mathrm{Cm}} \pm \sigma_{\mathrm{Cm}}$	$D_{ m Am}\pm\sigma_{ m Am}$	$\mathrm{SF_{Am/Cm}} \pm \sigma_{\mathrm{SF_{Am/Cm}}}$		
0.0125	0.37 ± 0.02	0.85 ± 0.06	2.27 ± 0.21	3.35 ± 0.13	6.39 ± 0.19	1.91 ± 0.09		
0.0250	1.65 ± 0.04	4.05 ± 0.05	2.45 ± 0.06	6.64 ± 0.39	12.51 ± 0.31	$\boldsymbol{1.88 \pm 0.12}$		
0.0750	32.09 ± 0.52	49.30 ± 2.88	1.54 ± 0.09	19.53 ± 0.40	30.15 ± 0.83	1.54 ± 0.05		
0.1500	185.0 ± 13.19	106.25 ± 7.85	0.57 ± 0.06	23.72 ± 2.10	41.12 ± 0.40	1.73 ± 0.15		

improvements in $SF_{Am/Cm}$ compared to existing systems, it revealed that dual ligands, even those with minimal individual extraction efficiency, can be combined to improve both extraction and separation under previously challenging conditions. Previous research on the Lanthaniden und Curium Americum Trennung (LUCA) process—a synergic system combining bis (chlorophenyl)dithiophosphinic acid and tris(2-ethylhexyl) phosphate—achieved an $SF_{Am/Cm}$ value greater than 7, suggesting that synergic systems hold significant promise for enhancing separation. This highlights the potential for newer systems, like those investigated in this work, to offer further advancements.

3.3. Preparation of HTTA and/or DBDECMP extraction chromatographic resins

Table 1 describes the combined and individual extractant-loaded resins that were prepared. The unrecovered resin losses (average 22%) are due to transfer loss and material bumping while under vacuum. The possibility of residual 1,2-dichloroethane and methanol in the final resins cannot be eliminated but is unlikely. The final dry density of resin 2 was $\rho_{\rm dry}$ = 0.3 g mL⁻¹, whereas for a wet resin it was $\rho_{\rm wet}$ = 0.7 g mL⁻¹.

3.4. Synergism in solid phase extraction

A corresponding solid-phase SEC was computed for the resin 2 using resin 4 (DBDECMP only) and resin 5 (HTTA only). The solid-phase SEC values of 1.42 \pm 0.02 at pH 2.00 and 1.19 \pm 0.02 at pH 2.50 demonstrate that synergism is retained upon conversion to an EXC resin. However, as pH increases, resin 5 begins to exhibit a higher affinity for ²⁴¹Am³⁺, thereby decreasing the observed synergic effect. While synergism is main-

tained at higher pH values, a single extractant may become increasingly dominant in the extraction process.

3.5. Weight distribution ratio (D_w) and $SF_{Am/Cm}$ of HTTA-DBDECMP resins

The $D_{\rm w}$ for $^{241}{\rm Am}^{3+}$ for the two studied mass ratios and two solvents are shown in Table 6. Resin 1 had a lower amount of DBDECMP compared to resin 2. Higher $D_{\rm w}$ for resin 1 suggest that smaller ratios of DBDECMP: HTTA yield higher $D_{\rm w}$ values. This agrees with the behavior observed in the SX systems. If similar extraction mechanisms are at play in both SX and EXC resins, then increasing the mass of DBDECMP while reducing the amount of HTTA is likely to result in decreased metal extraction. Comparing resin 2 and resin 3 showed that EXC resin made with 1,2-dichloroethane and methanol have similar $D_{\rm w}$ values with no observed impact on synergism. Overall, the $D_{\rm w}$ values for all resins showed an upward trend with increasing pH. The observed discrepancies between pH 2.00 and 2.25 are likely due to variations in ionic strength and

Table 6 Weight distribution ratios ($D_{\rm w}$) for Am(III) extraction with DBDECMP and HTTA resins as a function of pH in HNO₃ at varying ionic strengths

	HTTA-DBDECMP resins					
pH value (HNO ₃)	Resin 1 $D_{\rm w, Am} \pm \sigma_{\rm Am}$	Resin 2 $D_{\rm w, \ Am} \pm \sigma_{\rm Am}$	Resin 3 $D_{\rm w, Am} \pm \sigma_{\rm Am}$			
1.75	12.7 ± 0.3	9.0 ± 0.2	9.0 ± 0.5			
2.00	317.7 ± 6.0	219.4 ± 5.1	184.4 ± 1.5			
2.25	203.4 ± 1.6	138.0 ± 2.1	122.2 ± 1.7			
2.50	1546.4 ± 50.9	1143.1 ± 22.4	927.1 ± 18.2			

daily environmental variables that can impact pH, not captured in daily calibrations.

Table 7 shows extraction results done at constant ionic strength and in one day to avoid any effects of temperature on the pH calibrations. The highest separation factor achieved was 1.38 ± 0.02 . Curium exhibited higher $D_{\rm w}$ values at pH 1.75 and 2.00, while americium displayed higher $D_{\rm w}$ and at pH 2.25 and 2.50. Further studies are needed to determine if the switch in apparent preference is a flip in affinity or if other experimental conditions are at play. Overall, the separation factors between Am and Cm appears to increase with increasing pH. Further increasing pH might result in increasing separation factors, but at reduced utility with high $D_{\rm w}$ values. Evaluating $D_{\rm w}$ values of EXC resins at higher metal loading might reveal renewed possibilities.

3.6. Thermodynamics, kinetic, stability, and ionic strength experiments

3.6.1. Adsorption thermodynamics. The $D_{\rm w}$ values were multiplied by the $\rho_{\rm wet}$ to calculate $K_{\rm eq}$. Illustrated in Fig. 3 is a linear relationship between $\ln(D_{\rm w} \times \rho_{\rm wet})$ *versus* 1/T (K) for HTTA-DBDECMP resin 2 at pH 2.00 HNO₃. A strong but revers-

Table 7 Weight distribution ratios (D_w) and separation factors ($SF_{Am/Cm}$) for Am(III) and Cm(III) extraction with DBDECMP and HTTA resin 2, as a function of pH (HNO₃) at a constant ionic strength of 0.1 M

	HTTA-DBDECMP resin 2				
pH value (HNO ₃)	$D_{ m w, \ Cm} \pm \sigma_{ m Cm}$	$D_{ m w,\ Am}\pm\sigma_{ m Am}$	$SF_{Am/Cm} \pm \sigma_{Am}$		
1.75 2.00 2.25 2.50	131.4 ± 1.3 163.7 ± 5.1 341.5 ± 6.3 878.8 ± 13.1	94.9 ± 1.1 142.7 ± 1.4 396.5 ± 3.2 1204.1 ± 17.0	0.72 ± 0.01 0.87 ± 0.03 1.16 ± 0.02 1.37 ± 0.03		

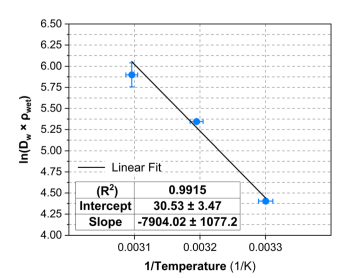


Fig. 3 Ln($D_{\rm w}$ × $\rho_{\rm wet}$) versus 1/T (K) for HTTA-DBDECMP resin 2 at pH 2.00 HNO₃.

ible dependence on temperature was observed, where increased extraction with increasing temperature was seen. Table 8 shows the derived thermodynamic properties for HTTA-DBDECMP resin 2 at pH 2.00 HNO₃. The positive value of ΔS indicates an increase in entropy, whereas the positive value of ΔH indicates an endothermic progress. Positive values of ΔH and ΔS suggest that the adsorption process is only spontaneous at high temperatures.

Extraction increases in synergic SX systems are driven by a gain in configurational entropy.⁵⁶ This entropy increase arises because synergic systems expand the number of possible extraction configurations for a metal ion. Whether a similar phenomenon occurs in synergic EXC resins remains unclear without further investigation. However, the observed trend of increasing extraction with temperature suggests that additional conformational states become energetically favorable at higher temperatures. If this theory holds, future experiments should explore less sterically hindered yet sufficiently hydrophobic neutral donors to access additional extraction configurations. Reduced steric hindrance could enhance binding at lower temperatures, which may be advantageous for high-loading applications.

3.6.2. Uptake kinetics. Adsorption data as a function of time (Fig. 4) revealed a rapid initial uptake from in the first two minutes, followed by slight desorption to the apparent equilibrium at 10 minutes. As the resin was not pre-conditioned with each acid prior to spiking in the metal, the decrease in $D_{\rm w}$ may be from the rapid initial leaching of extractant from the EXC resin as the metal solution diffuses. Future experiments could investigate this by pre-equilibrating the resin in solution before the addition of metal to disentangle resin wetting time and sorption kinetics. As all other samples were equilibrate for 60 minutes, they are expected to be in equilibrium.

3.6.3. γ -Irradiation and acid stability. Table 9 presents the $D_{\rm w}$ for γ -irradiated resin alongside experimental controls. The resulting $D_{\rm w}$ values for the γ -irradiated resin showed remarkably higher adsorption compared to the experimental controls. The acid contact control suggests that extractant leaching was likely responsible for the decreased adsorption observed. Ligand leaching is a well-known issue with EXC resins. Additionally, the inert macroporous support (pre-filter resin) with no ligand loading showed no adsorption.

3.6.4. Ionic strength dependence. Table 10 shows the $D_{\rm w}$ as a function of increasing ionic strength (Na,H(NO₃)). Decreased adsorption was observed with increasing ionic strength. Adsorption is sensitive to both pH and ionic strength, so control of both is necessary. When correcting pH value, introducing ions complicates comparing two separate metal adsorption experiments at identical ionic strengths. Column experiments were determined as a means of investigating separation while minimizing the impact of ionic strength.

3.7. Preliminary column separation studies

In both column types, americium was more strongly retained on the column compared to curium. This is opposite as what

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Table 8 Thermodynamic parameters (ΔS , ΔH , and ΔG) of Am(III) adsorption at pH 2.00 HNO₃ for HTTA-DBDECMP resin 2

				ΔG (kJ mol ⁻¹)		
				Temperature (K)		_
Resin	pH, HNO ₃	ΔS (kJ mol ⁻¹ K ⁻¹)	$\Delta H \left(\text{kJ mol}^{-1} \right)$	303 ± 1	318 ± 1	333 ± 1
HTTA-DBDECMP Resin 2	2.00	0.25 ± 0.03	65.72 ± 8.96	-11.19 ± 0.29	-14.99 ± 0.30	-18.80 ± 0.61

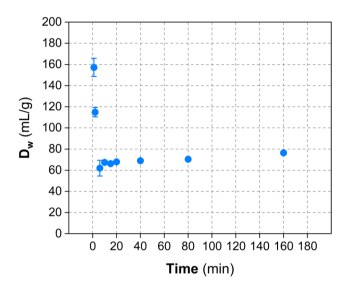


Fig. 4 Kinetic Uptake of ²⁴¹Am³⁺ adsorption on HTTA-DBDECMP Resin 2 (20-50 μm) at pH 2.00 HNO₃

Table 9 Comparison of weight distribution ratios (D_w) for Am(III) adsorption with DBDECMP-HTTA resin 2 at pH 2.00 HNO₃ between v-irradiated resin and experimental controls

	HTTA-DBDECMP resin 2						
pH value (HNO ₃)	Dry resin $D_{ m w, \ Am} \pm \sigma_{ m Am}$	γ -Irradiation resin $D_{ m w,\ Am}\pm\sigma_{ m Am}$	Acid contact $D_{ m w, \ Am} \pm \sigma_{ m Am}$	Pre-filter support $D_{ m w, \ Am} \pm \sigma_{ m Am}$			
2.00	71.90 ± 0.78	511.90 ± 9.65	14.48 ± 1.48	≅MDA ₉₅			

Table 10 Weight distribution ratios (D_w) for Am(III) extraction with HTTA-DBDECMP resin 2, as a function of ionic strength at pH 2.00

	HTTA-DBDECMP resin 2						
pH valu e (HNO ₃)	0.05 M $D_{ ext{w, Am}} \pm \sigma_{ ext{Am}}$	$\begin{array}{c} 0.10 \; \mathrm{M} \\ D_{\mathrm{w, \; Am}} \pm \sigma_{\mathrm{Am}} \end{array}$	$\begin{array}{l} 0.50 \; \mathrm{M} \\ D_{\mathrm{w, \; Am}} \pm \sigma_{\mathrm{Am}} \end{array}$	$1.00~\mathrm{M}$ $D_{\mathrm{w, \ Am}} \pm \sigma_{\mathrm{Am}}$			
2.00	82.94 ± 4.67	69.89 ± 3.69	58.62 ± 0.25	61.11 ± 1.36			

was observed in the batch $D_{\rm w}$ studies. $D_{\rm w}$ may differ between batch and column studies due to dynamic flow conditions, contact time variations, resin bed packing, concentration gradients, and changes in ionic strength and pH.

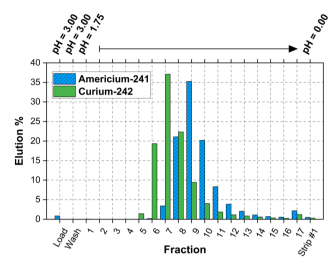


Fig. 5 Elution profile for ²⁴¹Am³⁺ and ²⁴²Cm³⁺ using column type 1 (inner diameter: 4.5 mm, length: 45 mm, column volume: ≈800 µL, and fraction volume: \approx 500 µL).

The elution 3.7.1. Column type 1. profile HTTA-DBDECMP Resin 2 in column type 1 is shown in Fig. 5. A maximum DF_{1,6}, for fractions 1 through 6 show a value of 10.

3.7.2. Column type 2. In attempts to further improve the separation, a longer and thinner column was employed. The elution profile for HTTA-DBDECMP resin 2 in column type 2 is shown in Fig. 6. Using a longer and thinner column showed an increase in the separation of americium and curium. Calculating a $DF_{8,10}$, for fractions 8, 9, and 10 using MDA_{95} for ²⁴¹Am³⁺ shows a value of 88 corresponding to a 27% curium recovery with no detectable americium. One notable difference between column type 1 and 2 is the observed elution profile. Column type 1 shows more uniform peak shapes, while type 2 shows significant tailing from ²⁴²Cm³⁺. This tailing likely results from two main contributions: (1) the column was stopped overnight at fraction 6 and resumed 12 hours later and (2) at fraction 11 gentle pressure was applied to increase the drip rate of the column. The next day after resuming the column and determining the time needed for collection of the remaining fractions, it was determined to apply gently pressure to increase the drip rate to (\approx 1.2 mL per hour). Given the large mesh size of the resin, length of the column, and fine chromatographic control required, we hypothesize that use of a peristaltic pump for this separation will improve the resolution of the peaks.

The raffinates from UNF reprocessing schemes present challenging matrices for chromatographic separations, which

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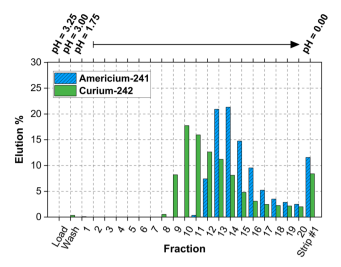


Fig. 6 Elution profile for ²⁴¹Am³⁺ and ²⁴²Cm³⁺ using column type 2 (inner diameter: 2.5 mm, length: 60 mm, column volume: \approx 400 μ L, and fraction volume: \approx 300 µL).

explains the limited literature on Am/Cm separations and a stronger focus on solvent extraction. 16 Most column chromatography efforts have centered on ion exchange, 57,58 with extraction chromatography being studied less frequently.⁵⁹⁻⁶¹ However, Usuda et al. investigated the use of single lipophilic extractant-impregnated adsorbents eluted with a hydrophilic complexant in nitric acid. In their study involving a TODGAbased adsorbent system with N,N,N',N'-tetraethyl-3,6-dioxaoctanediamide (DOODA(C2)) in dilute HNO3, they achieved relatively high-purity fractions, with the curium fraction containing 97.8% Cm and only 2.62% Am.62 In contrast, our preliminary column system yielded a lower overall Cm recovery of 27%, with no detectable Am. Another popular system that avoids the use of complexing agents altogether is a tertiary pyridine impregnated silica resin and a nitric acid/methanol elution mixture. This is a simple method to separate nearly all Am from Cm but poses safety concerns over the use of nitric acid and methanol. Suzuki et al. do not report a DF or % recovery/contamination in their work, but column separation factor of 2.54.63

These proof of principle column separations served as preliminary attempts to demonstrate that synergic EXC resins can facilitate effective separations. Despite facing chromatographic challenges, future investigations could enhance chromatographic performance and achieve more refined separations. This work establishes a foundational approach for synergic EXC resins, highlighting the potential for identifying synergic ligand combinations that could lead to improved separations of americium and curium.

Conclusions

The study presented here focused on identifying synergic extractant combinations to improve separation efficiency of americium and curium, aiming to fill a gap in the existing EXC resin compendium. Enhanced metal extraction was observed for all SX ligand combinations, though most showed limited selectivity between Am(III) and Cm(III). Although establishing clear relationships between synergism and selectivity is challenging, increased synergism generally correlates with decreased selectivity. An important exception was observed with DBDECMP and HTTA, which showed an increase in selectivity with increasing synergism. Subsequently, this ligand combination was translated into synergic EXC resins and evaluated under several experimental conditions. Once converted, the resulting resins exhibited acid dependency comparable to that of the SX experiments, while maintaining synergic behavior. The resulting $D_{\rm w}$ ratio for Am(III) and Cm(III) decreased compared to SX, suggesting reduced selectivity when translated and evaluated in EXC form. In preliminary column evaluations, the EXC resins exhibited promising metal retention and separation under mildly acidic conditions. Despite a low $D_{\rm w}$ ratio, the resin demonstrated the potential to separate Am(III) and Cm(III) with a DF_{8,10} = 88. However, further systematic investigations are necessary to enhance chromatographic performance and evaluate the technical readiness of this separation method.

This work demonstrates that a synergic EXC resin improves the extraction and separation of Am(III) and Cm(III) compared to individual ligand EXC resins; however, its performance does not surpass that of state-of-the-art ion-exchange chromatography. While not suitable for large-scale industrial applications, synergic EXC resins could enable the production of small-scale, high-purity Am and Cm samples for research and development. Overall, this study advances the development of EXC materials by demonstrating that synergic ligand combinations enhance the extraction and separation efficiency of trivalent actinides. Additionally, it highlights the feasibility of converting SX systems into EXC resins. These findings underscore the importance of exploring dual-ligand systems for the effective separation of Am(III) and Cm(III).

Author contributions

C. H. writing - original draft, writing - review & editing, visualization, validation, methodology, investigation, formal analysis, data curation, conceptualization, funding acquisition. T. M. Writing - review & editing, validation, supervision, resources, methodology, investigation formal analysis, conceptualization. J. S. writing - review & editing, validation, supervision, resources, project administration, methodology, investigation, funding acquisition, formal analysis, data curation, conceptualization.

Data availability

The authors confirm that the processed data supporting the findings of this study are included in the article. Raw counting data are available from the corresponding author upon reasonable request.

Conflicts of interest

Paper

There are no conflicts to declare.

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