Solvent extraction and fluorescence spectroscopic investigation of the selective Am(III) complexation with TS-BTPhen

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Abstract
An americium(III) selective separation procedure was developed based on the co-extraction of trivalent actinides (An(III)) and lanthanides (Ln(III)) by TODGA (N,N,N′,N′-tetraoctyl-diglycolamide), followed by Am(III) selective stripping using the hydrophilic complexing agent TS-BTPhen (3,3′,3″,3‴-[3-(1,10-phenanthroline-2,9-diyl)-1,2,4-triazine-5,5,6,6-tetrayl]tetrabenzenesulfonic acid). Distribution ratios were found at an acidity of 0.65 mol L⁻¹ nitric acid that allowed for the separation of Am(III) from Cm(III) (D_Cm > 1, D_Am < 1), giving a separation factor between curium and americium of SF_Cm/Am = 3.6 within the stripping step. Furthermore, Am(III) was readily separated from the lanthanides with the lowest selectivity for the Ln(III)/Am(III) separation being lanthanum with a separation factor of SF_La/Am = 20. The influence of the TS-BTPhen concentration on Am(III) distribution ratios was studied, giving a slope (logD vs. log[TS-BTPhen]) of approximately −1 for the stripping of An(III) with TS-BTPhen from the TODGA-based organic phase. Time-resolved laser fluorescence spectroscopy (TRLFS) measurements of curium(III) were used to analyze the speciation of Cm(III)-TS-BTPhen complexes. Both 1:1 and 1:2 complexes were identified in single-phase experiments.
The formation of the 1:1 complex was suppressed in 0.5 mol L\(^{-1}\) nitric acid but it was significantly present in HClO\(_4\) at pH 3. Conditional stability constants of the complex species were calculated from the TRLFS experiments.

Keywords: Am(III) separation, TODGA, selective stripping, curium, americium, lanthanides, partitioning, TS-BTPhen, americium-curium separation.

Introduction

The long-term radiotoxicity and heat load of used nuclear fuel is mainly due to the presence of the transuranium elements (TRU: Np - Cm). In the industrially applied PUREX process, recycling of the uranium and the most abundant TRU plutonium is accomplished. The minor actinides (MA) Np, Am, and Cm contribute to less than 0.1% of the initial spent fuel mass, but dominate the long-term radiotoxicity and heat load of the residual high active waste (e.g. vitrified waste) after the PUREX process.\(^{[1]}\) Therefore, the MA content in the disposed waste is a decisive factor in determining the size of a final geological repository for high active wastes. Neptunium can be separated in a modified PUREX process,\(^{[2]}\) leaving americium and curium as the principal contributors to the long-term radiotoxicity of the remaining waste. The partitioning and transmutation (P&T) concept strives for the multi-recycling of long lived MA in generation IV reactor systems by fast neutron irradiation. To achieve high americium concentrations and high purity in the transmutation targets, a separation of the actinides from lanthanides and fission products has to be achieved. Due to its high specific activity and the high share of spontaneous fission in the decay of curium, any dry or wet fabrication of Cm containing transmutation targets would require special shielding and handling. With respect to the half-life of the major curium isotope \(^{244}\)Cm (\(~18\) a), a separation of americium from curium and the fission and corrosion products is desirable. However, due to the chemical similarity between americium and curium, it is perhaps the most difficult separation in the P&T strategy.\(^{[3]}\) Within international research communities such as the SACSESS project of the European Commission\(^{[4]}\) and the US Department of Energy “Sigma-Team”,\(^{[5]}\) the development of processes for the selective removal of Am is being actively pursued. Different approaches have been investigated to carry out the difficult separation of americium from curium. In addition to solvent extraction methods where the separation of Am(III) is carried out by combining lipophilic and hydrophilic ligands, the
selective oxidation of americium to Am(IV) and Am(VI) followed by extraction of the oxidized species has also been investigated, but is not the topic of this paper.\textsuperscript{[6-7]} Several liquid-liquid extraction processes employing different strategies for MA handling in the fuel cycle have been established based on the high active PUREX- raffinate.\textsuperscript{[8-9]} On the one hand, there is a multi-cycle option making use of different consecutive processes with individual solvent compositions for each process. On the other hand, single cycle processes have been developed, opening the possibility to separate An(III) simultaneously from the light fission and corrosion products and Ln(III).\textsuperscript{[10]}

An important single cycle process is the innovative-SANEX process recently demonstrated at Forschungszentrum Jülich. The process follows the concept of an An(III) + Ln(III) co-extraction using TODGA (Figure 1) in a mixture of TPH containing 5 vol.-% 1-octanol.\textsuperscript{[11]} After scrubbing some undesired co-extracted fission products, the An(III) are then selectively stripped using a water-soluble tetrathedronated BTP (SO$_3$Ph-BTP).\textsuperscript{[12-14]} The remaining Ln(III) are then stripped using a buffer solution. The co-extraction and scrubbing steps of the innovative SANEX process were taken as the basis for this study. Changing the actinide-selective stripping of the innovative SANEX process to an americium(III) selective stripping step would lead to an americium selective process where americium is separated from curium. The hydrophilic actinide-selective N-donor complexing agent TS-BTPhen (3,3′,3″,3‴-[3-(1,10-phenanthroline-2,9-diyl)-1,2,4-triazine-5,5,6,6-tetrayl]tetrabenzenesulfonic acid, Figure 1) was studied for this purpose. The high selectivity of TS-BTPhen towards americium over curium has been previously described in the literature for the chromatographic separation of Am(III) from Cm(III) using a TODGA-PAN based chromatographic column.\textsuperscript{[15]} Preliminary studies on liquid-liquid extraction using TS-BTPhen\textsuperscript{[16]} in nitric acid and DGA-based extractants in the organic phase have been published recently by our group.\textsuperscript{[17]} In this paper the influence of the hydrophilic complexant TS-BTPhen on the extraction of actinides and lanthanides using TODGA was investigated with respect to its ability to carry out the selective back-extraction of americium from the organic phase.

Furthermore, the complexation of Cm(III) with TS-BTPhen was investigated herein in single-phase TRLFS titration experiments.

The concept of separating solely Am(III) from Cm(III) and all the FP in a single process was first realized by the EXAm process developed by the CEA in France.\textsuperscript{[18]} A separation factor between Am(III) and Cm(III) of $S_{\text{Am/Cm}} = 2.5$ was achieved within this process by using a combination of
DMDOHEMA and HDEHP (di(2-ethylhexyl)phosphoric acid) for extracting Am(III) and light Ln(III), and using TEDGA ($N,N,N',N'$-tetraethyldiglycolamide) as a hydrophilic complexant in the aqueous feed for complexing Cm(III) and heavy Ln(III) during the extraction.\cite{19} Co-extracted fission and corrosion products were removed by several pH-controlled scrubbing steps. Am(III) was selectively stripped from the solvent using buffered HEDTA solution, resulting in a 68-stage process. Another approach to the selective separation of Am(III) was developed using TODGA for co-extraction of An(III) and Ln(II) in combination with a water-soluble tetrasulfonated BTBP (SO$_3$-Ph-BTBP) for Am(III) selective stripping. The combination of TODGA and SO$_3$-Ph-BTBP used in the AmSel process provides separation factors of $S_{Cm(III)/Am(III)} = 2.5$ and $S_{Eu(III)/Am(III)} = 200$.\cite{16, 20-21} The TODGA–TS-BTPhen based concept investigated in this work is a less complex alternative approach to the EXAm process and provides higher separation factors for Cm(III) over Am(III) compared to the AmSel process.

**Figure 1:** Chemical structures of TODGA and TS-BTPhen.
Materials and Methods

TODGA was purchased from TechnoComm Ltd, UK. Hydrogenated tetrapropene (TPH) was received from the CEA Marcoule, France. Exxsol D80 (ExxonMobil) was used for biphasic TRLFS experiments. 1-Octanol was purchased from J.T. Baker ≥99%. The TS-BTPhen complexant was synthesized at the University of Reading according to the literature procedure.[16] Nitric acid solutions (Merck AG) were prepared by dilution from a 65% nitric acid solution EMSURE® for analysis.

Extraction experiments were carried out as forward extraction experiments using equal volumes of 500 µL of each phase. A combined aqueous tracer containing 300 kBq/mL $^{152}$Eu, 150 kBq/mL of $^{241}$Am and $^{244}$Cm each (10 µL) was used. A lanthanide solution containing all lanthanides except promethium in a concentration of $10^{-5}$ mol L$^{-1}$ of each lanthanide was used for the lanthanide experiments. An organic phase consisting of 0.2 mol L$^{-1}$ TODGA in TPH + 5 vol.-% 1-octanol was used. Aqueous phases were prepared by dissolving weighed amounts of TS-BTPhen in the appropriate aqueous phases. The shaking experiments were carried out in a temperature-controlled test tube shaker. Previous kinetic experiments showed that the applied contact time is sufficient. In batch kinetic experiments equilibrium D values were reached within contact times of less than 5 minutes. Afterwards, centrifugation was applied to maintain a complete separation of the phases, and the phases were separated using pipettes. Gamma measurements of $^{152}$Eu (122 keV) and $^{241}$Am (60 keV) were carried out using a Eurisys EGC35-195-R germanium coaxial N-type detector. Alpha measurements were carried out for $^{241}$Am (5486 keV) and $^{244}$Cm (5805 keV) using an Ortec Octète-pc eight chamber alpha measurement system equipped with PIPS detectors. Sample preparation was done by homogenizing the 10 µL alpha-spectroscopy sample in 100 µL of a mixture of Zapon varnish and acetone (1:100 v/v). This mixture was distributed over a stainless steel plate obtained from Berthold, Bad Wildbad, Germany. The sample was dried under a heating lamp and annealed into the stainless steel plate by a gas-flame burner. For stable elements, inductively coupled plasma mass spectrometry (ICP-MS) was applied using a Perkin Elmer SCIEX Elan 6100 DRC. Aqueous samples were measured after dilution in 1% v/v nitric acid solution without further treatment. Organic samples were measured directly in a tenside matrix (Triton-X-100) in 1% v/v nitric acid after dilution. The distribution ratios D were calculated as the quotient of the concentration or activity of the metal ion in the organic phase over the concentration or activity of the metal ion in the aqueous phase.
(D = c_{M(\text{org})}/c_{M(\text{aq})}). The separation between two metal ions is expressed using the separation factor, being the quotient of the distribution ratios of two metal ions (SF = D_{M1}/D_{M2}). The uncertainties of the data achieved from the method described above is < 5% for 0.01 < D < 100. Below and above these limits the uncertainties are lower than 20% for all data shown.

TRLFS measurements were carried out using a setup of a Continuum SURELITE Nd:YAG laser with a repetition rate of 10 Hz in combination with a NARROWscan D-R dye laser containing an Exalite 398 dye. For the excitation of Cm(III), a wavelength of 396.6 nm was used. The fluorescence was detected at a 90° angle to the exciting laser pulse using a Shamrock 303i spectrographic analyser with an ANDOR iStar generation III ICCD camera. Short-lived fluorescence was discriminated by starting the measurement at a time delay of 1 µs after laser excitation. Samples for single-phase titration experiments were prepared by adding 30 µl of a 3.34·10^{-6} mol L^{-1} Cm(III) stock solution to 970 µl of the respective medium. The Cm(III) stock solution had an isotopic mass distribution of 89.7% Cm-248, 9.4% Cm-246, ≤ 1% Cm-243, Cm-244, Cm-245 and Cm-247. With each step, a defined amount of complexant was added to the solution, increasing the total ligand concentration in the solution. Before starting the measurements, mixtures were allowed to equilibrate for 15 minutes.

A ligand stock solution was prepared by dissolving 2.63 mg of TS-BTPhen in 250 µl of the applied medium. This procedure gave a 10^{-2} mol/L TS-BTPhen solution. Solutions used for the titration experiment containing 10^{-3} -10^{-6} mol/L TS-BTPhen were prepared by subsequent dilution of the stock solution.

**Results and Discussion**

**Influence of the nitric acid concentration**

Batch extraction experiments were carried out to analyse the influence of the nitric acid concentration on the co-extraction of Am(III), Cm(III) and Eu(III), and the selective stripping of Am(III) using a 10 mmol L^{-1} solution of TS-BTPhen in different nitric acid concentrations spiked with $^{152}$Eu(III), $^{241}$Am(III), and $^{244}$Cm(III). The results are shown in Figure 2. Open symbols represent the distribution ratios of americium, curium and europium using TODGA + 5 vol.-% 1-octanol without the addition of TS-BTPhen to the aqueous phase. The distribution ratios of Am(III), Cm(III) and Eu(III) increase with increasing nitric acid concentrations which is known
to be the classical behavior of neutral extractants such as TODGA. The influence of TS-BTPhen addition to the aqueous phase is shown by the filled symbols. The addition of the hydrophilic ligand decreased the distribution ratios of An(III) by ca three orders of magnitude, while the Eu(III) distribution ratios were only decreased by one order of magnitude, resulting in a high Eu/Am separation factor of $SF_{Eu/Am} = 300$ at 0.65 mol L$^{-1}$ nitric acid. However, lanthanides lighter than Eu may have a significantly lower separation factor $SF_{Ln/Am}$. The distribution ratios of Am(III) were affected slightly more than those of Cm(III). The separation factor for Cm(III) over Am(III) increased from $SF_{Cm/Am} = 1.6$ for a TODGA-nitric acid system to $SF_{Cm/Am} = 2.4$ to 4.5 using TS-BTPhen in the nitric acid concentration range covered by Figure 2 (0.01 – 1 mol L$^{-1}$). The separation factors for Eu(III) over Am(III) using TS-BTPhen ($SF_{Eu/Am} = 70$-1000) in the aqueous phase are similar to the $SF_{Eu/Am}$ values reached using SO$_3$-Ph-BTP (250-1000), and are in agreement with the separation factors previously reported for TS-BTPhen. A high nitric acid concentration in the stripping step is favorable, as it simplifies scrubbing and gives robustness to the system. The region of interest for Am(III) selective separation using 10 mmol L$^{-1}$ TS-BTPhen is around 0.65 mol L$^{-1}$ initial nitric acid concentration, providing separation factors of $SF_{Eu/Am} \approx 300$ and $SF_{Cm/Am} = 3.6$. The distribution ratios of Am(III), Cm(III) and Eu(III) all show a linear increase with increasing TODGA concentration in the log(D) vs. log([TODGA]) plot. The results were already previously published.
Figure 2: Distribution ratios of Eu(III), Am(III) and Cm(III) and separation factors of Eu(III) and Cm(III) over Am(III) as a function of the initial nitric acid concentration in the radiotracer experiment. Organic phase: 0.2 mol L\(^{-1}\) TODGA in TPH + 5 vol.-% 1-octanol; aqueous phase: filled symbols: 10 mmol L\(^{-1}\) TS-BTPhen in nitric acid, open symbols: no addition of TS-BTPhen.

Behavior of the lanthanides

For a feasible selective separation of Am(III) from a loaded organic phase, not only is the separation of the trivalent actinides Am(III) and Cm(III) important, but so too is the separation of Am(III) from co-extracted lanthanides. The distribution ratios of Ln(III) are known to be high for TODGA-based organic phases at the relevant nitric acid concentrations.[25] Within the lanthanide group, the distribution ratios increase with decreasing ionic radius of the lanthanide.[26] Figure 3 shows the comparison of the distribution ratios of Ln(III) and An(III) using TODGA against nitric acid (Figure 3 A) and TODGA against nitric acid containing 10 mmol L\(^{-1}\) TS-BTPhen (Figure 3 B). Without using the hydrophilic complexing agent, the distribution ratios of Am(III) and Cm(III) are located between neodymium and samarium within the lanthanide group. The actinides are selectively removed from all of the lanthanides by using TS-BTPhen in nitric acid solution. The separation factor between the least extracted lanthanide La(III) and Am(III) is \(SF_{La/Am} \approx 20\) at 0.25 mol L\(^{-1}\) nitric acid. The selectivity for stripping of An(III) from Ln(III) is
lower compared to SO$_3$-Ph-BTP used for the An(III) stripping step in the innovative-SANEX test (SF$_{\text{La/An}} \approx 50$), but the selectivity for curium over americium (SF$_{\text{Cm/Am}}$) is significantly higher.[24] The initial concentration of the Ln(III) in the aqueous phase was $10^{-5}$ mol·L$^{-1}$ of each lanthanide.

![Figure 3](image.png)

Figure 3: Distribution ratios of Ln(III), Am(III) and Cm(III) with and without TS-BTPhen in the aqueous phase. Left side (A) without addition of TS-BTPhen, right side (B) with 10 mmol L$^{-1}$ TS-BTPhen. Organic phase: 0.2 mol L$^{-1}$ TODGA in TPH + 5 vol.-% 1-octanol. Aqueous phase: A: $10^{-5}$ mol L$^{-1}$ Ln(III) in nitric acid without addition of TS-BTPhen; B: $10^{-5}$ mol L$^{-1}$ Ln(III) and 10 mmol L$^{-1}$ TS-BTPhen in nitric acid; 30 minutes phase contact at 22 °C.

**Effect of the TS-BTPhen-concentration**

The influence of the TS-BTPhen concentration on the distribution ratios of Am(III), Cm(III) and Eu(III) was tested in the concentration range from 1 to 50 mmol L$^{-1}$. Figure 4 shows the distribution ratios as a function of the TS-BTPhen concentration on a logarithmic scale at constant nitric acid and TODGA concentrations.
A slope of $-0.8$ was observed for the linear regression of the Am(III) and Cm(III) data with increasing TS-BTPhen concentration. The Eu(III) data did not follow a linear trend in the double-logarithmic plot. In solution, the structurally similar lipophilic extractant CyMe₄BTPhen preferentially forms 1:2 complexes with An(III), as demonstrated by TRLFS and NMR studies. Therefore, a slope of $-2$ would also have been expected if the hydrophilic TS-BTPhen forms 1:2 complexes with An(III) in the aqueous phase. Equation 1 shows the simplified possible complexation reactions with An(III) in the TODGA/TS-BTPhen system. Similar deviations are also observed in the TODGA/SO₃-Ph-BTP and TODGA/SO₃-Ph-BTBP systems. The differences between a slope of $-0.8$ in the slope analysis and the expected slope of $-2$ may be related to the formation of mixed complexes. Time resolved laser fluorescence (TRLFS) studies were subsequently carried out to analyse the deviation between the slopes ($n = 1$ or 2) of the hydrophilic BTPhen and the literature stoichiometry of the lipophilic BTPhen, and to obtain comprehensive data on the complex properties.
**Equation 1:** Possible complexation reactions of An(III) in the biphasic TS-BTPhen-TODGA system.

\[ \text{An(III)}_{\text{aq}} + n \text{ TS-BTPhen}_{\text{aq}} \rightarrow [\text{An(III)(TS-BTPhen)}_n]_{\text{aq}} \]

\[ \text{An(III)}_{\text{aq}} + 3 \text{ TODGA} \rightarrow [\text{An(III)(TODGA)}_3] \]
Cm(III) TRLFS studies of TS-BTPhen

Complexometric titrations were carried out by adding TS-BTPhen to a $10^{-7}$ mol L$^{-1}$ Cm(III) solution in $10^{-3}$ mol L$^{-1}$ HClO$_4$ as well as in 0.5 mol L$^{-1}$ HNO$_3$. As ClO$_4^-$ is a very weakly coordinating anion, $10^{-3}$ mol/L HClO$_4$ was used as an idealized system with little influence of ionic strength, concurring anions or protonation of TS-BTPhen. However, biphasic extraction experiments were performed with the aqueous phase consisting of nitric acid (0.01 - 1.5 mol L$^{-1}$). Therefore, 0.5 mol L$^{-1}$ nitric acid was used as medium in further investigations to study the complexation of Cm(III) with TS-BTPhen under extraction conditions. Figure 5 shows the Cm(III) fluorescence emission spectra of the $^6D_{2} \rightarrow ^8S_{7/2}$ transition as a function of the TS-BTPhen concentration in $10^{-3}$ mol L$^{-1}$ HClO$_4$.

![Figure 5: Normalized fluorescence spectra of Cm(III) at increasing TS-BTPhen concentration (0 – 7.07x10$^{-5}$ mol L$^{-1}$) in $10^{-3}$ mol L$^{-1}$ HClO$_4$ ([Cm(III)] = 1x10$^{-7}$ mol L$^{-1}$)](image)

The fluorescence spectra in Figure 5 show three emission bands, the Cm(III) aquo ion (Cm(III) solvent species) at 593.9 nm and the emission bands of two complex species at 606.5 nm and 617 nm. The observed bathochromic shifts of the emission bands of the complex species are
characteristic of the formation of a 1:1 and a 1:2 complex with a strong tetradeinate N-donor ligand like TS-BTPhen\[^{20, 27, 29}\]. Peak deconvolution was carried out, providing the spectra of the pure components shown in Figure 6. The spectra of both complex species exhibit a hot band caused by population of a higher energy level by thermal effects. The hot bands are visible as shoulders at higher energy at 597 nm and 607 nm, respectively.\[^{30}\]

![Normalized fluorescence spectra of the [Cm(TS-BTPhen)]\(_n\) complexes in 10\(^{-3}\) mol L\(^{-1}\) HClO\(_4\) (n= 0, 1, 2).](image)

**Figure 6** Normalized fluorescence spectra of the [Cm(TS-BTPhen)]\(_n\) complexes in 10\(^{-3}\) mol L\(^{-1}\) HClO\(_4\) (n= 0, 1, 2).

The relative distribution of the different Cm(III) species after each titration step was obtained by peak deconvolution of the fluorescence spectra using the emission bands of the individual species (cf. Figure 6). As the fluorescence intensity increases strongly upon complexation as a result of intramolecular energy transfer processes,\[^{31}\] fluorescence intensity factors (FI) of the Cm-TS-BTPhen complexes have to be considered when calculating the concentrations of the formed Cm(III) species from the peak areas obtained by peak deconvolution. The FI factors were determined by fitting the increase of the Cm(III) fluorescence intensity as a function of TS-BTPhen concentration (FI[Cm(TS-BTPhen)] = 5, FI[Cm(TS-BTPhen)]\(_2\) = 560). The species distribution as a function of the TS-BTPhen concentration is shown in Figure 7.
Figure 7: Species distribution of Cm(III) as a function of the TS-BTPhen concentration in $10^{-3}$ mol L$^{-1}$ HClO$_4$. Symbols represent experimental data, lines are calculated with $\log \beta_{01} = 6.2$ and $\log \beta_{02} = 10.7$.

The complexation of Cm(III) started at a TS-BTPhen concentration of $3.48 \times 10^{-8}$ mol L$^{-1}$ with a maximum fraction of species 1 ($[\text{Cm(TS-BTPhen)}]$) of 78% at $4.48 \times 10^{-6}$ mol L$^{-1}$ TS-BTPhen. At concentrations above $3.45 \times 10^{-5}$ mol L$^{-1}$ complex species 2 ($[\text{Cm(TS-BTPhen)}_2]$) was the major species in solution.

A plot of the concentration ratios of the different species vs. the free ligand concentration in double logarithmic scale and subsequent linear regression yields slopes of 1.1 for the stepwise formation of the 1:1 and 1:2 complexes. The slopes depict the stoichiometric factor of TS-BTPhen added to the Cm(III) during transition from one species to another species. The overall transition from Cm(III) solvent species to the final complex yields a slope of 2.2, depicting a final 1:2 Cm(III) to TS-BTPhen complex ($[\text{Cm(TS-BTPhen)}_2]$). Therefore, the species with emission bands at 606.5 nm and 617.0 nm were identified as 1:1 and 1:2 complexes, respectively (Figure 8). Furthermore, conditional stability constants were derived from the speciation yielding $\log \beta_{01} = 6.2$ for the 1:1 complex, and $\log \beta_{02} = 10.7$ for the 1:2 complex.
Figure 8: Linear regression of the logarithm of the species ratios of the solvent species and the two complex species observed in titration experiments in $10^{-3}$ mol L$^{-1}$ HClO$_4$ vs. the logarithm of the free TS-BTPhen concentration.

The determination of the number of water molecules in the inner coordination shell by measurement of the fluorescence lifetime of the two complex species was not possible, due to the fluorescence quenching properties of TS-BTPhen.
Further experiments were performed in 0.5 mol L\(^{-1}\) HNO\(_3\) to study the complexation of Cm(III) under extraction conditions. Figure 9 shows the normalized fluorescence emission spectra of Cm(III) at increasing TS-BTPhen concentration in 0.5 mol L\(^{-1}\) nitric acid. The Cm(III) solvent species displays two emission maxima at 595 nm and 597 nm. These emission bands correspond to the formation of a mixture of Cm(III) aquo ion and Cm(III) mononitrato complex and are in excellent agreement with the literature.\(^{[32]}\) An additional species at 618.8 nm was observed at higher TS-BTPhen concentrations with a similar bathochromic shift as the [Cm(TS-BTPhen)\(_2\)] complex observed in 10\(^{-3}\) mol L\(^{-1}\) HClO\(_4\). The peak intensity of an additional 1:1 species at 606 nm was very low in nitric acid.

![Normalized fluorescence spectra of Cm(III) at increasing TS-BTPhen concentration (0 – 7.10x10\(^{-4}\) mol L\(^{-1}\)) in 0.5 mol/L HNO\(_3\) ([Cm(III)] = 1x10\(^{-7}\) mol L\(^{-1}\))](image)

**Figure 9:** Normalized fluorescence spectra of Cm(III) at increasing TS-BTPhen concentration (0 – 7.10x10\(^{-4}\) mol L\(^{-1}\)) in 0.5 mol/L HNO\(_3\) ([Cm(III)] = 1x10\(^{-7}\) mol L\(^{-1}\))

The pure component spectra of the Cm(III) solvent species and the [Cm(TS-BTPhen)\(_2\)] complex were obtained from the titration experiment. In addition, the spectrum of the [Cm(TS-BTPhen)] complex derived in 10\(^{-3}\) mol L\(^{-1}\) HClO\(_4\) was used for peak deconvolution. Considering the
fluorescence intensity factors \( \text{FI}[^{246}\text{Cm}(\text{TS-BrPhen})] = 5, \text{FI}[^{246}\text{Cm}(\text{TS-BrPhen})_2] = 620 \), the species distribution as a function of the free TS-BrPhen concentration was derived (Figure 10).

**Figure 10:** Species distribution of Cm(III) as a function of the TS-BTPhen concentration in 0.5 mol L\(^{-1}\) nitric acid. Symbols represent experimental data, lines are calculated with log\(\beta_{01}\) = 4.2 and log\(\beta_{02}\) = 8.8.

The complexation of Cm(III) with TS-BTPhen starts at \(2 \times 10^{-7}\) mol L\(^{-1}\) TS-BTPhen and exhibits a maximum for species 1 \([^{246}\text{Cm}(\text{TS-BrPhen})]\) of 24% at \(4.05 \times 10^{-5}\) mol L\(^{-1}\). At concentrations above \(4.1 \times 10^{-5}\) mol L\(^{-1}\) species 2 \([^{246}\text{Cm}(\text{TS-BrPhen})_2]\) is the major species present in the system.

The slope analyses calculated from the species distribution are given in Figure 11. A slope of 1.0 for species 1/solvent, 0.8 for species 2/species 1 depict the addition of 1 TS-BTPhen molecule. A slope of 1.8 for species 2/solvent depicts an overall complexation stoichiometry from the solvated curium ion to the 1:2 complex. The results confirm the assignment of the observed emission bands. The results show that a 1:1 complex is formed under extraction conditions as an intermediate minor species whereas the 1:2 complex predominates at high ligand concentrations.

The conditional stability constants were calculated to be log\(\beta_{01}\) = 4.2 for the 1:1 complex, and
logβ_{02} = 8.8 for the 1:2 complex. A summary of the obtained conditional stability constants in different media is given in Table 1.

![Figure 11: Linear regression of the logarithm of the peak ratios of the solvent species and the two complex species observed in the titration experiments in 0.5 mol L^{-1} HNO_{3} vs. the logarithm of the free ligand concentration.](image)

The minor amounts of the 1:1 complex obtained above in HNO_{3} compared to the titration experiment performed in 10^{-3} mol L^{-1} HClO_{4} is explained by the effects of the applied medium. As already observed by Wagner et al{sup}{20}, the increase of ionic strength destabilizes the mono anionic [Cm(TS-BTPhen)]^{-} complex relative to the penta anionic [Cm(TS-BTPhen){sub}2]^{5-} complex. Furthermore, the decrease of logβ_{02} resulting from the change of the medium from 10^{-3} mol L^{-1} HClO_{4} to 0.5 mol L^{-1} HNO_{3} (10.7 → 8.8) is explained by three different impacts: a change of the activity of TS-BTPhen and Cm(III) due to the increased ionic strength, and the competing reaction between protonated TS-BTPhen ligands and nitrate ions caused by the protonation of the N-atoms of TS-BTPhen. These effects result in a decreased concentration of free TS-BTPhen.
Previous studies have shown the protonation of the free ligand to be the major contribution to the decrease of the conditional stability constants.\textsuperscript{[20]}

Table 1 shows the conditional stability constants of TS-BTPhen in comparison to the stability constants of the SO\textsubscript{3}-Ph-BTBP complexant used in the AmSel process. Both complexants form an intermediate 1:1 Cm(III) to complexant and a final 1:2 Cm(III) to complexant complex. TS-BTPhen shows a slightly stronger complexation with Cm(III) compared to SO\textsubscript{3}-Ph-BTBP of the final 1:2 Cm(III) to complexant complex in both diluents. This effect was also observed for lipophilic BTBP and BTPhen ligands\textsuperscript{[27]} and is attributed to the greater degree of preorganization of the BTPhen ligand. The formation of the 1:1 Cm(III) to TS-BTPhen complex is stronger in the HClO\textsubscript{4} system which can be explained by a concurrency of the [Cm(III)(NO\textsubscript{3})\textsubscript{3}] complexes in nitric acid. Furthermore, the change of the solvent from 10\textsuperscript{-3} mol L\textsuperscript{-1} HClO\textsubscript{4} to 0.5 mol L\textsuperscript{-1} HNO\textsubscript{3} has a smaller effect on the final stability constant of the TS-BTPhen complex compared to that of SO\textsubscript{3}-Ph-BTBP.
Table 1: Conditional stability constants of \([\text{Cm(TS-BTPhen)}_n]\) (n = 1,2) and \([\text{Cm(SO}_3\text{-Ph-BTBP)}_n]\) (n = 1,2) complexes in \(10^{-3}\) mol L\(^{-1}\) HClO\(_4\) and 0.5 mol L\(^{-1}\) HNO\(_3\).

<table>
<thead>
<tr>
<th>Solvent</th>
<th>log(\beta_{01})</th>
<th>log(K_{12})</th>
<th>log(\beta_{02})</th>
</tr>
</thead>
<tbody>
<tr>
<td>TS-BTPhen (10^{-3}) mol L(^{-1}) HClO(_4)</td>
<td>6.2</td>
<td>4.5</td>
<td>10.7</td>
</tr>
<tr>
<td>TS-BTPhen (0.5) mol L(^{-1}) HNO(_3)</td>
<td>4.2</td>
<td>4.6</td>
<td>8.8</td>
</tr>
<tr>
<td>SO(_3)-Ph-BTBP (10^{-3}) mol L(^{-1}) HClO(_4)</td>
<td>5.3</td>
<td>5.1</td>
<td>10.4</td>
</tr>
<tr>
<td>SO(_3)-Ph-BTBP (0.5) mol L(^{-1}) HNO(_3)</td>
<td>-</td>
<td>-</td>
<td>7.3</td>
</tr>
</tbody>
</table>

**Biphasic TRLFS Experiments**

To analyze the complexes formed during the solvent extraction experiments, a two phase TRLFS experiment was performed. An aqueous phase containing Cm(III) and TS-BTPhen dissolved in 0.13 mol/L nitric acid was contacted with an organic phase containing TODGA in Exxsol D80 (a diluent very similar to TPH) + 5 vol.-% 1-octanol. After extraction, the phases were analyzed using TRLFS.

Figure 12 shows the spectra of the organic (left) and aqueous phases (right). The spectra of \([\text{Cm(TS-BTPhen)}_2]\) and \([\text{Cm(TODGA)}_3](\text{NO}_3)_3\)\(^{[26]}\) were added for comparison.
Figure 12: TRLFS-Spectra of the organic phase (left) and the aqueous phase (right) of a solvent extraction experiment. Organic phase: 0.2 mol L⁻¹ TODGA in Exxsol D80 + 5 vol.-% 1-octanol; aqueous phase: 10⁻⁷ mol L⁻¹ Cm(III), 1 mmol L⁻¹ TS-BTPhen in 0.13 mol L⁻¹ nitric acid. 30 minutes of extraction at RT, 5 minutes of centrifugation. For comparison, spectra of the [Cm(TODGA)₃](NO₃)₃ and [Cm(TS-BTPhen)₂] complexes from single phase experiments are added.

Figure 12 confirms the formation of a 1:2 Cm(III) TS-BTPhen complex in the aqueous phase of the extraction experiment. Furthermore, a comparison with the [Cm(TODGA)₃](NO₃)₃ spectrum shows the exclusive formation of [Cm(TODGA)₃](NO₃)₃ in the organic phase. The formation of mixed [Cm(TS-BTPhen)ₓ] complexes in the organic phase can be excluded within the sensitivity of the TRLFS measurement. Due to the low Cm(III) distribution ratios under the experimental conditions, the concentration of Cm(III) in the organic phase was low leading to a broadened peak with a low signal to noise ratio.

Conclusions
The novel complexing agent TS-BTPhen was investigated for its selectivity for Am(III) over Cm(III) and Ln(III) with a view to its application as a complexing agent for the selective recovery of Am(III) from PUREX raffinate. In solvent extraction experiments, the distribution ratios of the trivalent actinides were significantly reduced using TS-BTPhen in the aqueous phase in combination with TODGA in the organic phase. A separation factor between Cm(III) and Am(III) of SFₐm/Cm = 3.6 was achieved at nitric acid concentrations which allowed for a feasible separation of Am(III) from Cm(III) (D_Cm > 1, D_Am < 1 using 10 mmol L⁻¹ TS-BTPhen at 0.65 mol L⁻¹ nitric acid). The lanthanides were less affected by the presence of TS-BTPhen in the aqueous
phase resulting in a minimum separation factor of $S_{\text{FLa/Am}} = 20$ at 0.25 mol L$^{-1}$ nitric acid. The separation factor $S_{\text{FLn/An}}$ is lower compared to SO$_3$-Ph-BTP used in the i-SANEX test ($S_{\text{FLa/Am}} \approx 50$), but the $S_{\text{FCm/Am}}$ is higher, making the system attractive for a selective Am(III) separation process. A slope analysis of TS-BTPhen gave a slope of $-0.8$ for the actinides. To further investigate this result monophasic Cm(III) TRLFS studies were performed in 10$^{-3}$ mol L$^{-1}$ HClO$_4$ and 0.5 mol L$^{-1}$ HNO$_3$ identifying and quantifying the [Cm(TS-BTPhen)] and [Cm(TS-BTPhen)$_2$] complexes. Furthermore, TRLFS investigations of the aqueous phase of an extraction experiment revealed the exclusive formation of [Cm(TS-BTPhen)$_2$] during the extraction. The deviation between the slope analysis and the TRLFS results may be caused by the formation of mixed complexes influencing the slope in the extraction experiments. An i-SANEX based process with a selective Am(III) back extraction using TS-BTPhen thus seems to be possible and will be further investigated. This research contributes to the selective separation of only americium from PUREX raffinate.

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