

1 Lanthanide Speciation in Potential SANEX and GANEX Actinide/ 2 Lanthanide Separations Using Tetra-N-Donor Extractants

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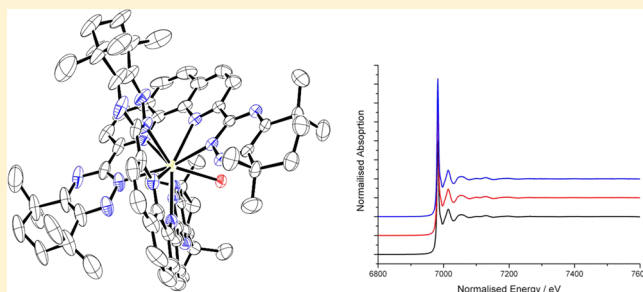
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13 **S** Supporting Information

14 **ABSTRACT:** Lanthanide(III) complexes with N-donor ex-
15 tractants, which exhibit the potential for the separation of
16 minor actinides from lanthanides in the management of spent
17 nuclear fuel, have been directly synthesized and characterized
18 in both solution and solid states. Crystal structures of the Pr³⁺,
19 Eu³⁺, Tb³⁺, and Yb³⁺ complexes of 6,6'-bis(5,5,8,8-tetramethyl-
20 5,6,7,8-tetrahydro-1,2,4-benzotriazin-3-yl)-1,10-phenanthroline
21 (CyMe₄-BTPhen) and the Pr³⁺, Eu³⁺, and Tb³⁺ complexes of
22 2,9-bis(5,5,8,8-tetramethyl-5,6,7,8-tetrahydro-1,2,4-benzotriazin-
23 3-yl)-2,2'-bipyridine (CyMe₄-BTBP) were obtained. The
24 majority of these structures displayed coordination of two of
25 the tetra-N-donor ligands to each Ln³⁺ ion, even when in some cases the complexations were performed with equimolar amounts
26 of lanthanide and N-donor ligand. The structures showed that generally the lighter lanthanides had their coordination spheres
27 completed by a bidentate nitrate ion, giving a 2+ charged complex cation, whereas the structures of the heavier lanthanides
28 displayed tricationic complex species with a single water molecule completing their coordination environments. Electronic
29 absorption spectroscopic titrations showed formation of the 1:2 Ln³⁺/L_{N-donor} species (Ln = Pr³⁺, Eu³⁺, Tb³⁺) in methanol when
30 the N-donor ligand was in excess. When the Ln³⁺ ion was in excess, evidence for formation of a 1:1 Ln³⁺/L_{N-donor} complex
31 species was observed. Luminescent lifetime studies of mixtures of Eu³⁺ with excess CyMe₄-BTBP and CyMe₄-BTPhen in
32 methanol indicated that the nitrate-coordinated species is dominant in solution. X-ray absorption spectra of Eu³⁺ and Tb³⁺
33 species, formed by extraction from an acidic aqueous phase into an organic solution consisting of excess N-donor extractant in
34 pure cyclohexanone or 30% tri-*n*-butyl phosphate (TBP) in cyclohexanone, were obtained. The presence of TBP in the organic
35 phase did not alter lanthanide speciation. Extended X-ray absorption fine structure data from these spectra were fitted using
36 chemical models established by crystallography and solution spectroscopy and showed the dominant lanthanide species in the
37 bulk organic phase was a 1:2 Ln³⁺/L_{N-donor} species.



38 **I** INTRODUCTION

39 The reprocessing of irradiated spent nuclear fuel (SNF) has
40 been performed since the 1940s, with the initial motivation to
41 isolate plutonium for military purposes but more recently with
42 the purpose to separate and recover both uranium and
43 plutonium in order to maximize the resources available to
44 generate civil nuclear energy.^{1,2} Reprocessing can also reduce
45 the volume of nuclear waste generated with high levels of
46 radioactivity due to the presence of long-lived radionuclides.^{1,2}
47 This separation is most commonly performed by PUREX
48 (Plutonium URanium EXtraction, also known as Plutonium

Uranium Reduction EXtraction), which is a biphasic solvent
49 extraction process whereby {UO₂}²⁺ and Pu⁴⁺, from SNF
50 dissolved in nitric acid (3–4 M), are extracted into an organic
51 phase containing tri-*n*-butyl phosphate (TBP; Figure 1) in a
52 hydrocarbon diluent (e.g., *n*-dodecane or odorless kero-
53 sene).^{1–4} The uranium and plutonium are transferred into
54

Special Issue: Inorganic Chemistry Related to Nuclear Energy

Received: July 23, 2012

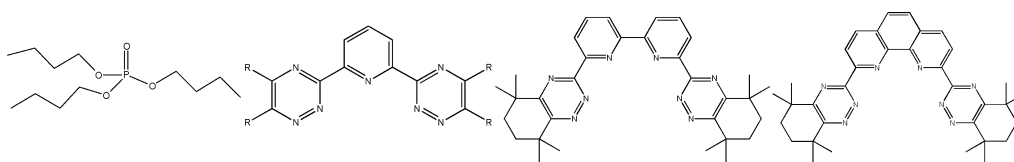


Figure 1. Structures of TBP (far left), BTP (center left), CyMe₄-BTBP (center right), and CyMe₄-BTPhen (far right).

55 the organic phase by forming charge-neutral complexes with
 56 TBP (i.e., [UO₂(TBP)₂(NO₃)₂] and [Pu(TBP)₂(NO₃)₄]).^{1,3–5}
 57 The plutonium, after reduction to Pu³⁺, and uranium are then
 58 back-extracted into an aqueous phase for reuse. The aqueous
 59 phase remaining after the initial separation, known as highly
 60 active raffinate (HAR), contains over 99.9% of the fission
 61 products (e.g., lanthanide isotopes, ¹³⁷Cs, ⁹⁰Sr, ⁹⁹Tc) and the
 62 minor actinide activation products (neptunium, americium, and
 63 curium) with decontamination factors of 10⁶–10⁸ achieved by a
 64 multistage separation process.¹ The long-term management of
 65 HAR, after conversion into an appropriate wasteform, can be
 66 extremely problematic, in part due to the presence of
 67 americium and curium, which are highly radioactive and have
 68 very long half-lives (up to 10⁵ years).^{1,2}

69 Considerable efforts have been made recently to develop
 70 advanced separation methodologies in order to maximize fuel
 71 resources and reduce the impact of nuclear waste while
 72 providing a proliferation-resistant fuel cycle (i.e., no pure
 73 plutonium is isolated).^{1,2,4,6–11} This forms part of the
 74 “Partitioning and Transmutation” strategy, where it is proposed
 75 that all of the actinides in SNF, including the minor actinides,
 76 can be separated and recycled as nuclear fuel. Another option is
 77 to “burn” the separated actinides, which will also result in
 78 conversion to short-lived fission product nuclides but without
 79 nuclear energy production for public consumption. This
 80 provides the added benefit of converting most of the long-
 81 lived actinides in SNF to shorter-lived fission product nuclides
 82 compared to current spent fuel management options. As a
 83 result, the “Partitioning and Transmutation” strategy can
 84 significantly reduce the time it takes for SNF to decay to
 85 radioactivity levels of natural uranium and therefore the
 86 necessary design lifetime of any nuclear waste repository.^{7–11}

87 One of the major separation challenges that need to be
 88 overcome for this strategy to be successful is the separation of
 89 americium and curium from the lanthanide fission products.
 90 This is because the high neutron absorption cross sections of
 91 some of the lanthanide ions present in SNF both decrease the
 92 flux in a reactor and create more activation products, thereby
 93 making transmutation a less attractive option if the lanthanides
 94 cannot be separated from the actinides.¹⁰ Achieving this
 95 separation is extremely difficult because of the chemical
 96 similarities between americium, curium, and the lanthanides,
 97 which all most commonly exist in the III+ oxidation state in
 98 solution.¹² Consequently, organic molecules that can selectively
 99 extract actinides, in particular Am³⁺ and Cm³⁺, over the Ln³⁺
 100 ions are of great interest, as is evident by the number of
 101 different ligand systems and processes that have been
 102 developed by various groups in the field of partition-
 103 ing.^{2,4,6–10,12–22} Examples include the TALSPEAK (Trivalent
 104 Actinide Lanthanide Separation by Phosphorus reagent
 105 Extraction from Aqueous Komplexes) process, which uses
 106 diethylenetriaminepentaacetic acid in a lactic acid solution to
 107 hold back Am³⁺ and Cm³⁺ in the aqueous phase while the
 108 lanthanide ions are extracted into the organic phase containing
 109 di(2-ethylhexyl)phosphoric acid,^{13,14} and the TRUEX (TRANS-

Uranic EXtraction) process, where the addition of octyl-
 (phenyl)-*N,N*-diisobutylcarboylmethylphosphineoxide to the
 organic phase in the core PUREX process allows Am³⁺ and
 Cm³⁺ to be extracted alongside {UO₂}²⁺ and Pu⁴⁺, leaving the
 lanthanide ions and other fission products in the aqueous
 phase.^{14,15}

The SANEX (Selective ActiNide EXtraction) solvent
 extraction process^{8,9} aims to separate the minor actinides
 americium and curium from the lanthanide fission products
 remaining after plutonium and uranium removal by PUREX
 and fission product separation (except the lanthanides) by
 DIAMEX (DIAMide EXtraction)¹⁶ using only carbon-, hydro-
 gen-, oxygen-, and nitrogen-containing compounds as extrac-
 tants, diluents, or phase modifiers. A class of molecules that
 showed early promise for the selective extraction of An³⁺ over
 Ln³⁺ in a SANEX process were the tridentate 2,6-bis(5,6-
 dialkyl-1,2,4-triazin-3-yl)pyridines (BTPs; Figure 1).^{7,17} How-
 ever, many of these extractant molecules suffered problems that
 precluded them from use in plant-scale extractions including
 poor stability, slow extraction kinetics, and inefficient back-
 extraction due to high An^{III} affinities.⁷ Further developments in
 the use of triazinyl-based N-donor extractants for actinide/
 lanthanide separations have led to the tetradentate ligand 2,9-
 bis(5,5,8,8-tetramethyl-5,6,7,8-tetrahydro-1,2,4-benzotriazin-3-
 yl)-2,2'-bipyridine (CyMe₄-BTBP; Figure 1), which exhibits
 significant potential for use in SANEX separations, with
 separation factors for Am³⁺ over Eu³⁺ found to be
 ~150.^{7,18,19} The CyMe₄-BTBP extractant has been successfully
 tested for the extraction of genuine actinide/lanthanide feed
 through a 16-stage centrifugal contactor setup with excellent
 recoveries for americium and curium (>99.9%) but has been
 shown to undergo radiolytic degradation at doses that will be
 encountered at the high minor actinide loadings obtained in the
 reprocessing of, for example, fast reactor fuels.¹⁹ The kinetics
 for actinide extraction with CyMe₄-BTBP are still relatively
 slow, so the addition of a phase-transfer catalyst is necessary
 [e.g., *N,N'*-dimethyl-*N,N'*-dioctylethylethoxymalonamide
 (DMDOHEMA)] if this extractant is to be used for large-
 scale partitioning.¹⁹ In an attempt to improve the kinetics of
 extraction with these tetradentate N-donor extractants, greater
 conformational rigidity was enforced in the ligand backbone
 with the synthesis of 2,9-bis(5,5,8,8-tetramethyl-5,6,7,8-tetrahy-
 dro-1,2,4-benzotriazin-3-yl)-1,10-phenanthroline (CyMe₄-
 BTPhen; Figure 1).²⁰ This rigid ligand displays very high
 separation factors for Am³⁺ over Eu³⁺ (up to 400) with
 significantly faster kinetics of extraction compared to those
 found for CyMe₄-BTBP, thereby eliminating the need for a
 phase-transfer catalyst.²⁰ These high separation factors even at
 low acidities for the aqueous phase may prove problematic
 during back-extractions,⁷ but the use of alternative diluents has
 shown that efficient back-extractions may be achievable when
 using the CyMe₄-BTPhen extractant.²⁰

An alternative concept being considered in Europe for the
 recovery of actinides from SNF is the GANEX (Group
 ActiNide EXtraction) process, which is proposed to consist 164

165 of two cycles.^{16,21,22} Most of the uranium is removed in the first
 166 cycle, while the second cycle recovers all of the remaining
 167 actinides, mainly the transuranics neptunium through curium,
 168 concurrently in varying oxidation states (III–VI) from the
 169 fission products found in spent fuel, including the lanthanides.
 170 The GANEX process is aimed for generation IV nuclear fuel
 171 cycles, where plutonium is likely to exist in higher
 172 concentrations during partitioning processes compared to
 173 those found in the processing of SNF in current cycles.²¹
 174 The major novelty with GANEX compared to most other more
 175 technologically mature separation processes is that the
 176 plutonium is routed with the minor actinides rather than with
 177 the majority of the uranium. The separation of Am³⁺ and Cm³⁺
 178 from the lanthanide ions in a SANEX process is already
 179 considered extremely challenging, so performing the same
 180 separation in addition to partitioning neptunium, plutonium,
 181 and any remaining uranium from all of the fission products in
 182 the second stage of the GANEX process is even more difficult.
 183 A single extractant in the organic phase is unlikely to achieve
 184 the group separation of multiple actinides in variable oxidation
 185 states with appropriate efficiencies. Consequently, the perform-
 186 ance of multiple extractants in the organic phase, typically
 187 already established from other separation processes, has been
 188 explored for use in a GANEX process.^{16,21,22} A number of
 189 different extractant combinations have been shown to have
 190 potential including *N,N,N',N'*-tetraoctyldiglycolamide
 191 (TODGA; used in DIAMEX) with DMDOHEMA, TODGA
 192 with TBP, and CyMe₄-BTBP with TBP.^{16,21,22}
 193 The N-donor extractants CyMe₄-BTPhen and CyMe₄-BTBP
 194 have already demonstrated potential as extractants for
 195 partitioning SNF mixtures, in particular the separation of
 196 minor actinides from the lanthanides.^{7,18–20} However, the
 197 mode of action of these ligands with these metal ions in
 198 extraction conditions has not been definitively established.
 199 Here, we have produced numerous Ln³⁺ complexes across the
 200 lanthanide series with both CyMe₄-BTPhen and CyMe₄-BTBP
 201 ligands using a direct synthetic approach. These complexes
 202 have been fully characterized in both solution and solid states
 203 using multiple techniques including electronic absorption
 204 spectroscopy, luminescence spectroscopy, and single-crystal
 205 X-ray diffraction (XRD). We have then used X-ray absorption
 206 spectroscopy (XAS) to probe the lanthanide (europium and
 207 terbium) species, which have been extracted into the organic
 208 phase using conditions similar to those proposed for SANEX
 209 and GANEX separation processes that use CyMe₄-BTPhen and
 210 CyMe₄-BTBP. The extended X-ray absorption fine structure
 211 (EXAFS) of the Ln L_{III}-edge XAS spectra obtained from each
 212 of these systems has been fitted to structural models established
 213 by characterization of the directly synthesized Ln³⁺ complexes
 214 with these N-donor extractants, thus providing definitive
 215 evidence for Ln³⁺ speciation in the bulk organic phase during
 216 extraction processes.

217 ■ RESULTS AND DISCUSSION

218 **Synthesis.** Ln^{III} complexes of the extractant CyMe₄-
 219 BTPhen (see Table 1 for the list) were readily synthesized by
 220 the addition of Ln(NO₃)₃ (Ln = Pr, Eu, Tb, Yb) in acetonitrile
 221 to 1 mol equiv of CyMe₄-BTPhen in dichloromethane (DCM).
 222 The reaction solution was allowed to evaporate to dryness,
 223 leaving a powder that could be crystallized from a mixture of
 224 CH₃CN, DCM, and ethanol in a volume ratio of ~2:2:1, where
 225 CH₃CN readily dissolves the complex, DCM acts to reduce the
 226 solubility of the complex in solution, and ethanol improves the

Table 1. List of Synthesized Complexes

formula	compound number
[Pr(CyMe ₄ -BTPhen) ₂ (NO ₃)](NO ₃) ₂ ·10H ₂ O	1
[Pr(CyMe ₄ -BTPhen) ₂ (NO ₃)] [Pr(NO ₃) ₅] ²⁻ ·1.63EtOH·0.75H ₂ O	2
[Eu(CyMe ₄ -BTPhen) ₂ (H ₂ O)](NO ₃) ₃ ·9H ₂ O	3
[Tb(CyMe ₄ -BTPhen) ₂ (H ₂ O)](NO ₃) ₃ ·9H ₂ O	4
[Yb(CyMe ₄ -BTPhen) ₂ (H ₂ O)](NO ₃) ₃ ·9H ₂ O	5
[Pr(CyMe ₄ -BTBP) ₂ (NO ₃)](NO ₃) ₂ ·4EtOH·H ₂ O	6
[Pr(CyMe ₄ -BTBP) ₂ (NO ₃) ₂][Pr(NO ₃) ₆](NO ₃) ₃ ·6CH ₃ CN	7
[Eu(CyMe ₄ -BTBP) ₂ (NO ₃)](NO ₃) ₂ ·4EtOH·2H ₂ O	8
[Eu(CyMe ₄ -BTBP)(NO ₃) ₃] ²⁻ ·toluene	9
[Tb(CyMe ₄ -BTBP) ₂ (H ₂ O)](NO ₃) ₃ ·4EtOH	10

227 miscibility of the solvent mixture. In all examples, yellow
 228 crystals were obtained. Elemental analysis, single-crystal XRD
 229 (see the Solid-State Structure section), and electrospray
 230 ionization mass spectrometry (ESI-MS, positive ion) indicated
 231 that, in the majority of cases, complex cations of stoichiometry
 232 1:2 Ln³⁺/CyMe₄-BTPhen with nitrate counterions were
 233 obtained even though the syntheses were conducted with
 234 equimolar amounts of Ln(NO₃)₃ and CyMe₄-BTPhen. The
 235 only exception was found during the synthesis of the Pr³⁺
 236 complex of CyMe₄-BTPhen, where the major product consisted
 237 of a 1:2 Pr/CyMe₄-BTPhen complex cation but with a
 238 [Pr(NO₃)₅]²⁻ counterion present per cationic unit. The initial
 239 crystallization of this mixture led to isolation of a small amount
 240 of this cationic species with only nitrate present as counterions,
 241 as determined by XRD (see the Solid-State Structure section).
 242 The structural determinations show that the Ln³⁺ coordination
 243 sphere is completed by a single nitrate anion for the Pr³⁺
 244 complexes (1 and 2), while for the Eu³⁺, Tb³⁺, and Yb³⁺
 245 complexes (3–5), a single molecule of water completes the
 246 coordination sphere (see the Solid-State Structure section).
 247 However, ESI-MS spectrometry of all the studied Ln³⁺
 248 complexes with CyMe₄-BTPhen from a methanol (MeOH)
 249 solution indicates that a nitrate ion is coordinated, and there
 250 was no evidence to suggest that a water molecule was present in
 251 the coordination sphere.

252 The synthesis of Ln³⁺ complexes (Ln = Pr, Eu, Tb) of
 253 CyMe₄-BTBP (see Table 1 for the list) was also attempted by
 254 adding a DCM solution of the ligand to 0.5 equiv of Ln(NO₃)₃
 255 in MeOH. The powder obtained upon evaporation of the
 256 reaction mixture was best crystallized by slow evaporation from
 257 a 1:1:1:1 by volume mixture of toluene, isopropyl alcohol,
 258 ethanol, and DCM. The alcohols dissolve the complexes
 259 reasonably well, while the use of toluene and DCM reduces the
 260 solubility of the complexes, assists in controlling the rate of
 261 evaporation, and provides reasonable miscibility in these
 262 solvent mixtures. Characterization of the bulk crystallized
 263 material obtained from all of the attempted Ln³⁺ complexations
 264 of CyMe₄-BTBP indicated that a mixture of products was
 265 present, which is likely to be due to the formation of products
 266 with different combinations of Ln³⁺/CyMe₄-BTBP ratios and
 267 anionic molecular ions (i.e., NO₃⁻, [Ln(NO₃)₆]³⁻, [Ln-
 268 (NO₃)₅]²⁻). However, the selection of individual crystals
 269 obtained from these reactions was able to afford the structural
 270 determination of a number of products by XRD. The vast
 271 majority of these structures indicated complex cations of 1:2
 272 Ln³⁺/CyMe₄-BTBP stoichiometry (6–8 and 10) with nitrates
 273 (6–8 and 10) and metallonitrates (7) present as counterions.
 274 The first structures of Ln-BTBP complexes to be isolated were 274

275 with the ligand 6,6'-bis(5,6-diethyl-1,2,4-triazin-3-yl)-2,2'-bipyridine (C2-BTBP), and these had a single C2-BTBP molecule
 276 coordinated to the Ln^{3+} ion.²³ It was noted that in solution
 277 both 1:1 and 1:2 $\text{Ln}^{3+}/\text{C2-BTBP}$ complexes were observed.²³
 278 More recently, crystals of $[\text{Eu}(\text{CyMe}_4\text{-BTBP})_2(\text{NO}_3)]^{2+}$ with a
 279 metallonitrate counterion and the charge-neutral species
 280 $[\text{Eu}(\text{CyMe}_4\text{-BTBP})(\text{NO}_3)_3]$ were isolated by slow evaporation
 281 from a mixture of DCM and CH_3CN .²⁴ Our attempts to form
 282 the Eu^{3+} complex of $\text{CyMe}_4\text{-BTBP}$ produced a 1:1 $\text{Eu}^{3+}/$
 283 $\text{CyMe}_4\text{-BTBP}$ molecular species with a toluene molecule
 284 present as a solvent of crystallization (9) in addition to the
 285 1:2 $\text{Eu}^{3+}/\text{CyMe}_4\text{-BTBP}$ complex cation containing species but
 286 with only nitrate counterions present in the lattice. The Pr^{3+}
 287 and Eu^{3+} complexes isolated in the solid state (6–9) have one
 288 or more nitrate ions completing the coordination sphere, while
 289 only the Tb^{3+} complex of $\text{CyMe}_4\text{-BTBP}$ has a water molecule in
 290 its coordination environment. The ESI-MS spectra of all of the
 291 $\text{CyMe}_4\text{-BTBP}$ complexes obtained from MeOH indicated that
 292 the only intact molecular species present was $[\text{Ln}(\text{CyMe}_4\text{-}$
 293 $\text{BTBP})_2(\text{NO}_3)]^{2+}$. The ESI-MS spectra of the $\text{CyMe}_4\text{-BTBP}$
 294 complexes provide comparable results and are in agreement
 295 with similar ESI-MS studies previously performed on extracted
 296 solutions of Eu^{3+} with BTBP extractants.²⁵ This suggests that
 297 the 1:2:1 $\text{Ln}^{3+}/\text{CyMe}_4\text{-BTBP}/\text{NO}_3^-$ complex is dominant in
 298 solution, while other compositions were only present in
 299 solution in minor quantities, if at all.

301 **Solution Spectroscopy.** The UV–visible absorption
 302 spectra of complexes 2–4, isolated in a pure bulk form,
 303 dissolved in MeOH are dominated by charge-transfer
 304 transitions in the UV region of the spectra (see the Supporting
 305 Information). These transitions are most likely due to $\pi-\pi^*$
 306 transitions from the aromatic nature of the $\text{CyMe}_4\text{-BTBP}$
 307 ligand. A clear difference in the spectral profile is observed
 308 between the free $\text{CyMe}_4\text{-BTBP}$ ligand and Ln^{3+} complexes,
 309 indicating that the electronic structure of the $\text{CyMe}_4\text{-BTBP}$
 310 molecule is perturbed upon Ln^{III} coordination. Essentially no
 311 difference is observed between the spectroscopic profiles for 2–
 312 4, indicating that there is little or no influence by the type of
 313 coordinating lanthanide ion on the electronic structure of the
 314 $\text{CyMe}_4\text{-BTBP}$ ligand. The limited solubility of these
 315 complexes in most common solvents precluded the study of
 316 the typically weakly absorbing $f-f$ transitions of the lanthanides
 317 in 1-cm-path-length cells.

318 Titrations of $\text{CyMe}_4\text{-BTBP}$ and $\text{CyMe}_4\text{-BTBP}$ with the
 319 lanthanide ions, Pr^{3+} , Eu^{3+} , and Tb^{3+} in MeOH were performed
 320 to study the lanthanide speciation behavior of these extractant
 321 molecules, in particular the equilibrium between 1:1 and 1:2
 322 $\text{Ln}^{3+}/\text{L}_{\text{N}_4\text{-donor}}$ species. The titrations of $\text{CyMe}_4\text{-BTBP}$ with
 323 each of the lanthanides studied show that there is essentially no
 324 difference in the titration profiles with different lanthanide ions
 325 (see Figure 2 for Pr^{3+} and Supporting Information). Sharp
 326 decreases in the intensity of the absorption maxima for free $\text{CyMe}_4\text{-BTBP}$
 327 at 261 and 295 nm with the addition of up to 0.5
 328 equiv of $\text{Ln}(\text{NO}_3)_3$ are observed. The absorption maximum at
 329 261 nm also shifts to ~ 266 nm with the addition of $\text{Ln}(\text{NO}_3)_3$.
 330 Isosbestic points are observed at 229 and 279 nm. Further
 331 additions of $\text{Ln}(\text{NO}_3)_3$, up to 3 equiv, result in a subtle decrease
 332 in the absorption intensity for most of the spectrum but with
 333 no changes in the shape of the spectral profile. This indicates
 334 that the 1:2 $\text{Ln}^{3+}/\text{CyMe}_4\text{-BTBP}$ complex forms with the
 335 initial addition of $\text{Ln}(\text{NO}_3)_3$, as expected.^{23,26} The subtle
 336 changes in the spectra when more than 0.5 equiv of $\text{Ln}(\text{NO}_3)_3$

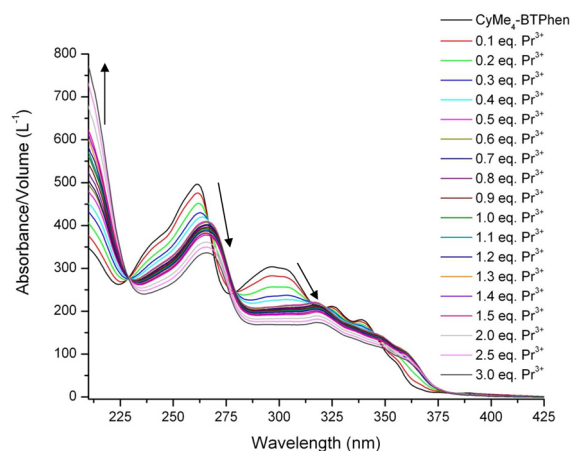


Figure 2. UV–visible absorption spectroscopic titration of $\text{CyMe}_4\text{-BTPhen}$ with $\text{Pr}(\text{NO}_3)_3$ in MeOH (initial conditions, $[\text{CyMe}_4\text{-BTPhen}] = 2.0 \times 10^{-5}$ M, volume = 2.0 mL; titrant conditions, $[\text{Pr}(\text{NO}_3)_3] = 4.0 \times 10^{-4}$ M).

is present in solution are most likely explained by an
 equilibrium being established between 1:1 and 1:2 $\text{Ln}^{3+}/$
 $\text{CyMe}_4\text{-BTPhen}$ species, where more 1:1 complex is likely to
 form with increasing additions of $\text{Ln}(\text{NO}_3)_3$. Similar behavior is
 observed for the titrations of $\text{CyMe}_4\text{-BTBP}$ with $\text{Ln}(\text{NO}_3)_3$
 (see Figure 3 for Eu^{3+} and Supporting Information).

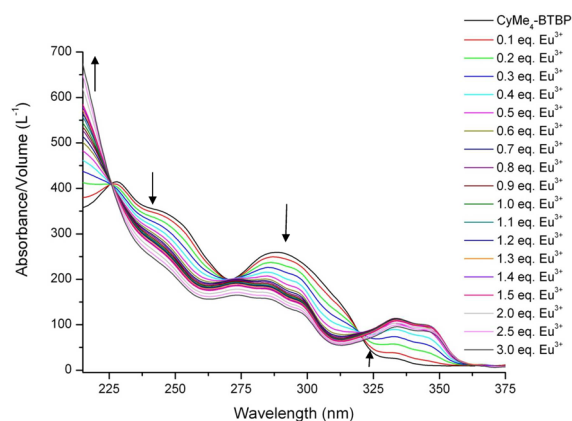


Figure 3. UV–visible absorption spectroscopic titration of $\text{CyMe}_4\text{-BTBP}$ with $\text{Eu}(\text{NO}_3)_3$ in MeOH (initial conditions, $[\text{CyMe}_4\text{-BTBP}] = 2.0 \times 10^{-5}$ M, volume = 2.0 mL; titrant conditions, $[\text{Eu}(\text{NO}_3)_3] = 4.0 \times 10^{-4}$ M).

Absorption maxima at 228 and 289 nm sharply decrease in
 intensity with the initial addition of $\text{Ln}(\text{NO}_3)_3$ up to 0.5 equiv.
 Two absorption maxima are seen to emerge at 334 and 346 nm
 with the initial addition of $\text{Ln}(\text{NO}_3)_3$. Further additions of
 $\text{Ln}(\text{NO}_3)_3$ also result in a subtle decrease in the absorption
 intensity for most of the spectrum. Therefore, it can be
 deduced that the 1:2 $\text{Ln}^{3+}/\text{CyMe}_4\text{-BTBP}$ complex is probably most favored to form, but the 1:1 species
 can be forced to form in solution with excess Ln^{3+} ion present.
 Similar results have been previously observed for Ln^{3+}
 complexation behavior with analogous BTBP ligands.²⁶

The overall stability constants for both 1:1 and 1:2 $\text{Ln}^{3+}/$
 $\text{CyMe}_4\text{-BTBP}$ – $\text{CyMe}_4\text{-BTBP}$ species were determined by
 fitting the appropriate spectrophotometric titration data (Table
 2). These fits confirm that the formation of both ML and ML_2
 (where L is the N-donor ligand) species does occur over the

Table 2. Fitted Metal–Ligand Overall Stability Constants Determined from UV–Visible Spectroscopic Data Using Hyperquad²⁸ ($I = 0$ M in MeOH; $T = 25$ °C)

N-donor ligand	Ln ³⁺	overall stability constant ($\log \beta_{ML}$)				σ^b
		$\log \beta_{11}$	standard deviation ^a	$\log \beta_{12}$	standard deviation ^a	
CyMe ₄ -BTPPhen	Pr ³⁺	4.7	0.5	11.8	0.1	0.0032
	Eu ³⁺	7.9	0.5	15.6	1.0	0.0028
	Tb ³⁺	8.1	0.5	13.2	0.5	0.011
CyMe ₄ -BTBP	La ³⁺	4.4 ^c	0.2 ^c	8.8 ^c	0.1 ^c	
	Pr ³⁺	10.9	0.7	18.9	1.1	0.0042
	Eu ³⁺	9.5	0.6	16.9	1.1	0.0067
		6.5 ^c	0.2 ^c	11.9 ^c	0.5 ^c	
	Tb ³⁺	8.8	0.2	15.9	0.4	0.0037
	Yb ³⁺	5.9 ^c	0.1 ^c			

^aStandard deviations determined by the fitting process. ^bGoodness-of-fit parameter. ^cReference 27 ($I = 0.01$ M Et₄NNO₃; $T = 25$ °C; in MeOH; determined by UV–visible absorption spectroscopy).

359 conditions used in these titrations, as has been observed
360 previously in similar titrations of CyMe₄-BTBP with La³⁺ and
361 Eu³⁺.²⁷ The speciation plots corresponding to the titrations
362 with CyMe₄-BTBP (see the Supporting Information) show the
363 initial emergence of the ML₂ species when less than 0.5 mol
364 equiv of lanthanide is present (relative to L), with further
365 additions of lanthanide showing the increasing formation of the
366 ML species. The magnitude of the lanthanide stability constants
367 for the CyMe₄-BTBP species indicates the greatest affinity for
368 the mid-lanthanides with lower stability constants obtained for
369 the lanthanides at either end of the series, which is in
370 agreement with previous work and the corresponding
371 distribution ratios for Ln³⁺ extractions using CyMe₄-BTBP
372 and DMDOHEMA into *n*-octanol.^{12,27} The speciation plots for
373 the CyMe₄-BTPPhen titrations (see the Supporting Informa-
374 tion) indicate behavior different from that observed for CyMe₄-
375 BTBP. For Pr³⁺, the 1:2 Ln/CyMe₄-BTPPhen species is favored
376 to form, compared to the 1:1 M/L species even at relatively
377 high metal concentrations due to a highly positive cooperative
378 effect for the formation of the ML₂ species. However, this
379 strong cooperative effect diminishes substantially with progress
380 along the lanthanide series where the ML species is
381 predominantly favored for Tb³⁺ even at reasonably low metal
382 concentrations. The stability constants for 1:1 Ln/CyMe₄-
383 BTPPhen increases as the lanthanide series is traversed. The
384 differences observed between the lanthanide stability behaviors
385 for complexes of CyMe₄-BTBP and CyMe₄-BTPPhen are most
386 likely due to the lack of flexibility in the BTPPhen backbone,
387 resulting in the greater likelihood of a mismatch between the
388 lanthanide ionic radius and the CyMe₄-BTPPhen binding cavity
389 as the lanthanide series is traversed.

390 The absorption spectroscopic profiles showed little difference
391 between the light and heavy lanthanides, but XRD studies (see
392 the Solid-State Structure section) indicate that the heavy
393 lanthanides in the 1:2 Ln³⁺/CyMe₄-BTPPhen–CyMe₄-BTBP
394 complexes prefer to have their coordination sphere completed
395 by water, whereas the lighter lanthanide complexes generally
396 prefer to have nitrate in their coordination environment, a
397 consequence of the lanthanide contraction. This is commonly
398 observed in a series of lanthanide complexes of a given
399 multidentate ligand.²⁹ Luminescence studies were therefore
400 undertaken in an attempt to assess the involvement of nitrate

and water in the coordination sphere of these lanthanide
401 species, as has been performed previously to investigate the
402 coordination behavior of other extractant molecules.³⁰
403 Excitation and emission spectra of the Eu³⁺ and Tb³⁺ complexes
404 with CyMe₄-BTPPhen and CyMe₄-BTBP are displayed in Figure
405 4 and in the Supporting Information. Excitation into the
406 4

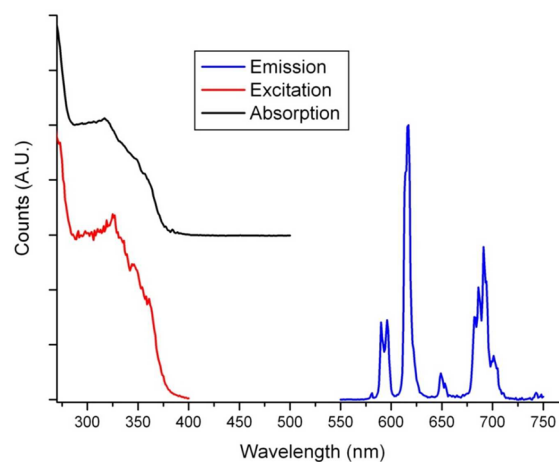


Figure 4. Emission (following excitation at 320 nm), excitation (monitoring emission at 616 nm), and absorption spectra of [Eu(CyMe₄-BTPPhen)₂(X)]ⁿ⁺ in MeOH ($X = \text{H}_2\text{O}/\text{NO}_3^-$; $n = 3$ and 2).

intraligand absorption bands (280–330 nm) of the Eu³⁺ and
407 Tb³⁺ complexes produced characteristic f-centered emission
408 spectra with resolvable bands due to the ⁵D₀ to ⁷F_J and ⁵D₄ to
409 ⁷F_J ($J = 0–6$) transitions, respectively. The emission spectrum
410 of the Eu³⁺ complexes are dominated by the electric-dipole-
411 allowed $\Delta J = 2$ transition, which is hypersensitive to the site
412 symmetry. The absence of a hyperfine structure in this band
413 indicates that the complexes exist as a single emissive species on
414 the experimental time scale.³¹ The emission profiles for the
415 Eu³⁺ complexes are similar to those observed with other BTBP
416 ligands, but in our examples, the splitting of the ⁵D₀ to ⁷F₂
417 transition at ~617 nm upon complexation with the N-donor
418 ligands is not resolved, which has been observed previously in
419 some examples.^{24,32} The respective excitation spectra recorded
420 at the emission maxima (545 nm for Tb³⁺ and 616 nm for
421 Eu³⁺) display ligand-centered absorption bands that overlap
422 well with the absorption spectra, indicating that sensitized
423 emission is occurring in all of the systems under study.
424

In order to assess the inner coordination sphere of the
425 complexes, lifetime data were recorded in MeOH and MeOH-
426 *d*₄ following 320 nm excitation (e.g., see Figure 5) and the
427 number of coordinated MeOH molecules determined accord-
428 ing to Horrock's equation (eq 1)³³
429

$$q_{\text{bound MeOH}} = A \left[\left(\frac{1}{\tau_{\text{MeOH}}} \right) - \left(\frac{1}{\tau_{\text{CD}_3\text{OD}}} \right) \right] \quad (1)$$

where A is a proportionality constant; $A = 2.1$ ms for Eu³⁺ and
431 $A = 8.4$ ms for Tb³⁺.
432

For solutions of Eu³⁺ and CyMe₄-BTPPhen in a 2:1 molar
433 ratio, this gave a q value of 0.3; an identical q value was
434 obtained for the analogous complex with CyMe₄-BTBP of 0.3
435 (Table 3). This strongly suggests that the first coordination
436 sphere of the complexes is completed by ligation of nitrate
437

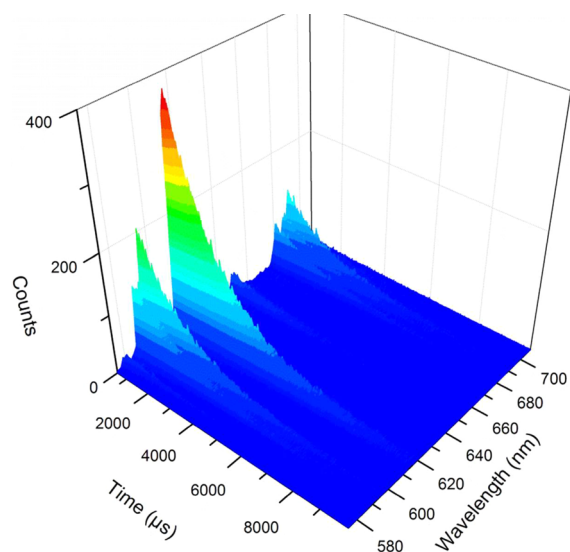


Figure 5. Time-resolved emission spectrum of $[\text{Eu}(\text{CyMe}_4\text{-BTPhen})_2(\text{X})]^{n+}$ in MeOH following excitation at 320 nm ($\text{X} = \text{H}_2\text{O}/\text{NO}_3^-$; $n = 3$ and 2).

Table 3. Photophysical Properties of Solutions of $\text{Ln}(\text{NO}_3)_3$ with Tetra-N-Donor Ligands in a 1:2 Molar Ratio at 298 K^a

complex	λ_{em} (nm)	τ_{MeOH} (ms)	τ_{MeOD} (ms)	q_{MeOH}
$[\text{Eu}(\text{BTBP})_2(\text{X})]^{n+}$	617	1.94	2.61	0.3
$[\text{Eu}(\text{BTPhen})_2(\text{X})]^{n+}$	617	1.49	1.87	0.3

^aAll lifetimes were recorded by TCSPC at 320 nm excitation using a 5 W xenon flashlamp and are subject to a $\pm 10\%$ error. Identical data within error were obtained for 1:3 and 1:5 solutions of $\text{Eu}^{3+}/\text{L}_{\text{N}_4\text{-donor}}$ and the crystalline complexes **3** and **8**.

438 anions rather than exchangeable solvent molecules, and there
439 may be a minor species that exists, with either water or MeOH
440 occupying this coordination site for these Eu^{3+} complexes.
441 Because the emissive quantum yield of a solvated species would
442 be much lower, the contribution to the initial emission intensity
443 will be low, perhaps precluding observation of a second species
444 in solution, and/or the rate of solvent and nitrate anion
445 exchange is much faster than the luminescence time scale, so a
446 noninteger value of q is determined. Similar data were obtained
447 for 1:3 and 1:5 molar ratios of Eu^{3+} with both N_4 -donor ligands
448 and the isolated complexes **3** and **8**, suggesting that the 1:2
449 $\text{Ln}^{3+}/\text{L}_{\text{N}_4\text{-donor}}$ complex is the only emissive species formed
450 under these conditions.

451 In the case of the Tb^{3+} complexes of both ligands, excitation
452 into the ligand absorption bands resulted in comparatively weak
453 emission spectra. This is unsurprising given the estimated
454 triplet energies of the ligands and the high-energy emissive $^5\text{D}_4$
455 excited state and suggests that back-energy transfer from the
456 Tb^{3+} excited-state manifold to the ligand triplet state is a
457 competitive nonradiative decay process.³⁴ This is corroborated
458 by the fact that the radiative lifetimes for the Tb^{3+} emission are
459 extremely short; the kinetic traces could be satisfactorily fitted
460 with two exponential functions, giving lifetime values of
461 approximately 18 and 6 μs (for solutions of BTBP in
462 MeOH). Moreover, the kinetic traces recorded without a
463 time gate and delay additionally exhibit a short-lived
464 component of nanosecond order, which we attribute to
465 ligand-centered emission.

Solid-State Structure. Single-crystal XRD studies of
466 complexes of Tb^{3+} , Eu^{3+} , and Pr^{3+} with ligands $\text{CyMe}_4\text{-BTBP}$
467 and $\text{CyMe}_4\text{-BTPhen}$ were obtained (**1–4** and **6–10**,
468 respectively). The complex of Yb^{3+} with $\text{CyMe}_4\text{-BTPhen}$
469 was also studied (**5**). Complexes **3–5** are isostructural crystallizing
470 in the orthorhombic space group $Fdd2$. Plots of these structures
471 are displayed in Figures 6–11 (complexes **1**, **3**, **6**, and **8–10**)
472 f6 – f11

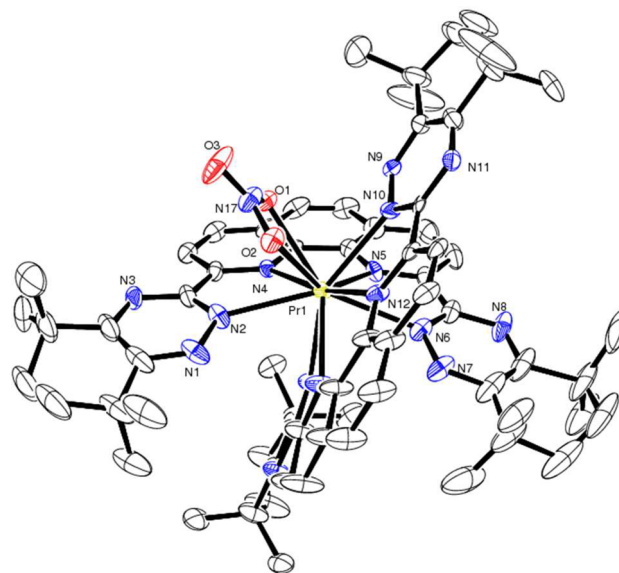


Figure 6. ORTEP plot of the complex cation of **1**, with crystallographic numbering (H atoms omitted). Probability ellipsoids of 50% displayed.

and the Supporting Information (complexes **2**, **4**, **5**, and **7**) with
473 crystal data given in Tables 4 and 5. In the vast majority of
474 cases (**1–8** and **10**), two of the N-donor ligands (either
475 $\text{CyMe}_4\text{-BTBP}$ or $\text{CyMe}_4\text{-BTPhen}$) were found to coordinate to
476 the metal center occupying four coordination sites each, with
477 another ligand (water or nitrate) occupying a cavity between 478

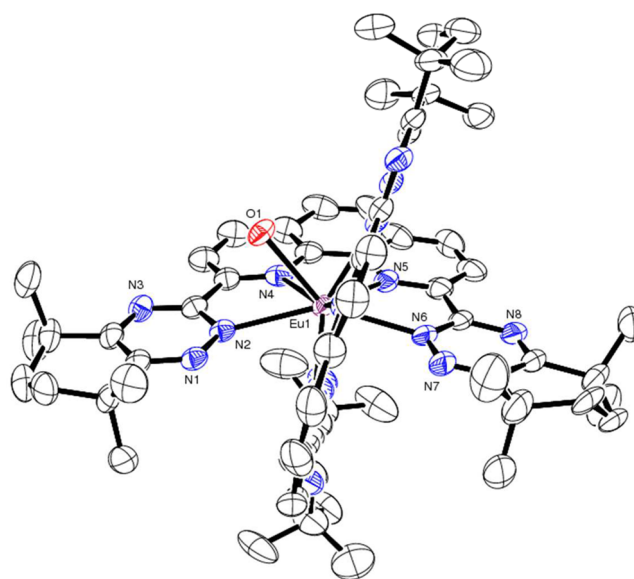


Figure 7. ORTEP plot of the complex cation of **3**, with crystallographic numbering (H atoms omitted). Probability ellipsoids of 50% displayed.

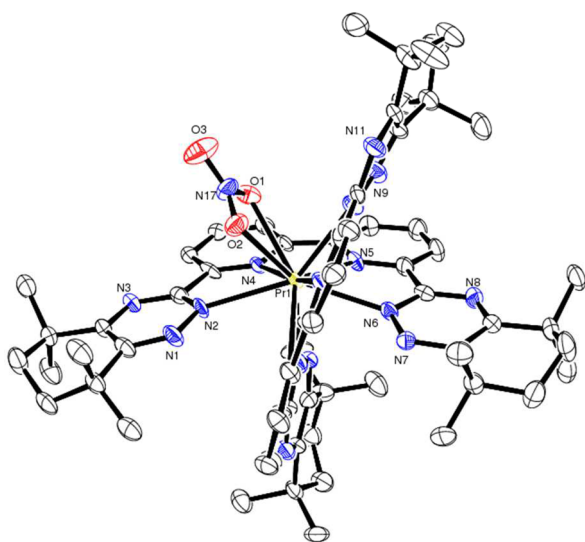


Figure 8. ORTEP plot of the complex cation of **6**, with crystallographic numbering (H atoms omitted). Probability ellipsoids of 50% displayed.

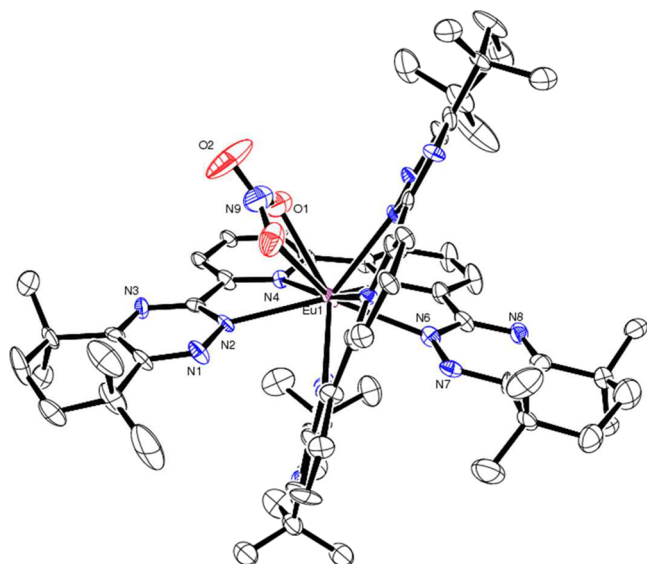


Figure 9. ORTEP plot of the complex cation of **8**, with crystallographic numbering (H atoms omitted). Probability ellipsoids of 50% displayed.

479 the two bound N-donor ligands, giving a distorted capped
480 square-antiprismatic geometry about the Ln^{3+} center. This leads
481 to a total coordination number of 9 for water-coordinated
482 complexes (**3–5** and **10**) and 10 for the bidentate nitrate-
483 coordinated complexes (**1**, **2**, and **6–8**).

484 For the Ln^{3+} complexes with $\text{CyMe}_4\text{-BTPhen}$, only 1:2 $\text{Ln}^{3+}/$
485 $\text{L}_{\text{N-donor}}$ coordination stoichiometries have been isolated and
486 structurally characterized in the solid state. The nitrate ion is
487 found to occupy the remaining coordination sites in the Pr^{3+}
488 complexes isolated, while a single water molecule completes the
489 coordination sphere for the $\text{CyMe}_4\text{-BTPhen}$ complexes of the
490 heavier Ln^{3+} ions investigated in this study (Eu^{3+} , Tb^{3+} , and
491 Yb^{3+} in **3–5**). This is likely to be due to a combined effect of
492 the lanthanide contraction and the structural rigidity of the
493 $\text{CyMe}_4\text{-BTPhen}$ ligand sterically hindering the remaining
494 coordination sites in the more contracted structures of Eu^{3+} ,

Tb^{3+} , and Yb^{3+} such that only water can access this binding
495 cavity in these solid-state systems. However, previous work has
496 shown that the 1:2 complex of $\text{Eu}^{3+}/\text{CyMe}_4\text{-BTPhen}$ can be
497 obtained with a nitrate ion completing the coordination sphere
498 in the solid state where MeOH was used as the reaction
499 solvent,²⁰ thus indicating that the position of the equilibrium
500 between bound nitrate and bound water in these Ln^{3+}
501 complexes may be influenced by the choice of solvent. The
502 nitrate-coordinated complexes form 2+ charged complex
503 cations, while the water-coordinated complexes form tricationic
504 complex cations, where charge balance is achieved with
505 nonbinding nitrate anions in the crystal lattice (**1** and **3–5**)
506 or with an anionic metallonitrate species (**2**). The previously
507 obtained $[\text{Eu}(\text{CyMe}_4\text{-BTPhen})_2(\text{NO}_3)]^{2+}$ solid-state complex
508 was also charge-balanced with a pentanitratolanthanide anionic
509 species.²⁰

510 All of the M–N bond lengths in the $\text{CyMe}_4\text{-BTPhen}$ -
511 containing structures decrease as the lanthanide series is
512 traversed from left to right (Table 6), as expected due to the
513 lanthanide contraction. In all cases, the lanthanide ion sits
514 outside of the plane of the N-donor ligand cavity. The out-of-
515 plane displacement of the Ln^{3+} ion from the average plane
516 defined by the four coordinating N atoms for each N-donor
517 ligand follows a trend similar to that of the bond lengths by
518 decreasing across the lanthanide series: $\sim 0.80/0.71$, $0.77/0.62$,
519 0.56 , 0.55 , and 0.51 Å for species **1–5**, respectively. The
520 average M–N_{triazinyl} bonds lengths are consistently longer than
521 those for the M–N_{phen} bonds in the Eu^{3+} , Tb^{3+} , and Yb^{3+}
522 complexes (**3–5**). This may imply that a greater degree of
523 interaction exists between the Ln^{3+} ion and the phenanthroline
524 N-donor atoms than that with the triazinyl N-donor atoms.
525 However, the same cannot be said for the structures of the Pr^{3+}
526 complexes obtained (**1** and **2**), where in some instances the
527 M–N_{phen} bond lengths are, in fact, longer than the M–N_{triazinyl}
528 bond distances. The previously obtained structure of $[\text{Eu}-$
529 $(\text{CyMe}_4\text{-BTPhen})_2(\text{NO}_3)]^{2+}$ shows little difference between
530 the Eu–N_{triazinyl} and Eu–N_{phen} bond distances.²⁰ Therefore, it is
531 most likely the triazinyl groups that are restrained to be further
532 away from the Ln^{3+} center relative to the phenanthroline
533 backbone as the Ln center approaches the plane of the $\text{CyMe}_4\text{-}$
534 BTPhen binding cavity, as this is only evident for the latter
535 lanthanides. The Ln–O_{water} bond distances also decrease as the
536 lanthanide series is traversed from left to right because of
537 lanthanide contraction (Table 6). The Pr–O_{nitrate} bond
538 distances for **1** and **2** [2.592(7) and 2.544(7) Å for **1**;
539 2.581(5) and 2.605(5) Å for **2**] are typical for Pr^{3+} complexes
540 with coordinated nitrates (2.5–2.8 Å).^{23,35,36}

541 Where $\text{CyMe}_4\text{-BTBP}$ is the ligand, both 1:1 (**9**) and 1:2
542 $\text{Ln}^{3+}/\text{CyMe}_4\text{-BTBP}$ (**6–8** and **10**) coordination structures were
543 isolated. Structures of metal complexes with $\text{CyMe}_4\text{-BTBP}$ have
544 only been previously obtained for Eu^{3+} , U^{4+} , and $\{\text{UO}_2\}^{2+}$.^{36,37}
545 Previous studies of the complexation of Eu^{3+} with $\text{CyMe}_4\text{-}$
546 BTBP , using a preparation similar to that described in this
547 work, isolated structures consisting of the same 1:2 and 1:1
548 $\text{Eu}^{3+}/\text{CyMe}_4\text{-BTBP}$ complexes found (structures **8** and **9**,
549 respectively). However, these structures exhibit different crystal
550 forms due to either different counterions or alternate solvent
551 molecules of crystallization present in the lattices.²⁴ Further
552 structural information has been obtained for Ln^{3+} complexes
553 with C2-BTBP, where only 1:1 $\text{Ln}/\text{C2-BTBP}$ complexes were
554 isolated, essentially for the entire lanthanide series.²³ The
555 remaining coordination sites were occupied by three nitrate
556 anions to give charge-neutral species.²³ The structure of the
557

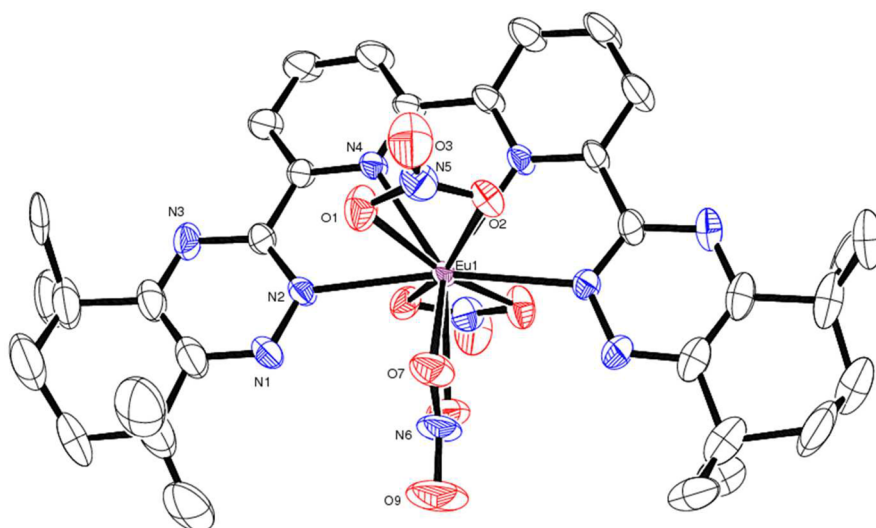


Figure 10. ORTEP plot of the complex molecule of **9**, with crystallographic numbering (H atoms omitted). Probability ellipsoids of 50% displayed.

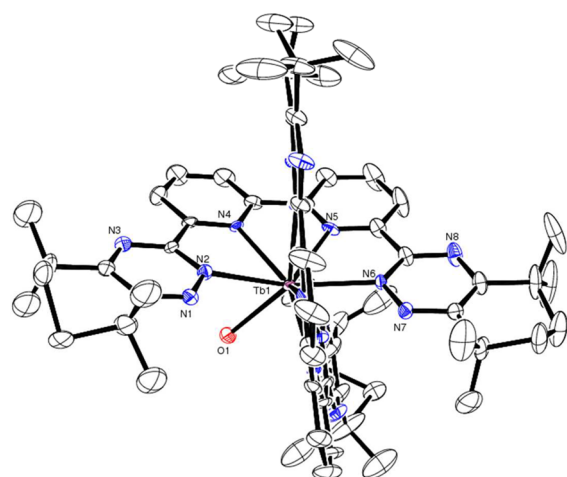


Figure 11. ORTEP plot of the complex cation of **10**, with crystallographic numbering (H atoms omitted). Probability ellipsoids of 50% displayed.

Eu³⁺ complex, **9**, is analogous to the Ln³⁺ complexes of C2-
BTBP. For the cationic Ln³⁺ complexes of CyMe₄-BTBP,
charge balance was achieved either with extra lattice nitrate
anions (**6**, **8**, and **10**) or in combination with a
hexanitratometallo anion (**7**). The two crystalline forms
obtained from the complexation of Eu³⁺ with CyMe₄-BTBP
offers further insight into the equilibrium between 1:1 and 1:2
Ln/BTBP-BTPhen complex stoichiometries. Although it may
be possible for both of these stoichiometries to be isolated, the
vast majority of the structural evidence indicates that the
lanthanides preferentially coordinate to two of these tetra N-
donor ligands from this class of extractant molecules. In
contrast to the CyMe₄-BTPhen structures, metal-bound nitrate
ions are observed with all CyMe₄-BTBP species except Tb³⁺.
This is presumably due to the greater flexibility afforded from
the bipyridine, compared to the “locked” phenanthroline,
permitting the sterically larger bidentate nitrate anion, relative
to water, to bind to the Ln³⁺ center.

For all of the complexes of CyMe₄-BTBP (**6**–**10**), the Ln–N
bond distances (Table 7) decrease as the lanthanide series is
traversed from left to right, similar to the CyMe₄-BTPhen and

C2-BTBP-containing structures.²³ The Ln–O_{nitrate} bond
lengths also clearly decrease across the series, demonstrating
lanthanide contraction again. The 1:2 Ln³⁺/CyMe₄-BTBP
complexes bear further similarity to those of CyMe₄-BTPhen
with the Ln³⁺ ion located outside of the average plane of the
tetra-N-donor cavity and this displacement following the same
trend as that of the bond lengths, decreasing across the series:
~0.73/0.78, 0.72/0.76, 0.69, and 0.56 Å for **6**–**8** and **10**,
respectively. However, the 1:1 Eu³⁺/CyMe₄-BTBP complex (**9**)
does effectively sit in the plane average plane of the four N-
donor atoms (out-of-plane displacement ~0 Å). The
coordination bond lengths and motifs observed in the
structures of the 1:1 and 1:2 Eu³⁺/CyMe₄-BTBP complexes
(**8** and **9**) are similar to those observed for the structures
obtained previously for the same complex molecules but in
different crystal forms.²⁴ There is little difference observed
in the Ln–N bond lengths between the 1:1 and 1:2 Ln³⁺/BTBP
complex molecular species obtained here and elsewhere,²³
suggesting that if there are indeed any cooperative or
destructive effects for 1:2 Ln³⁺/Cy-Me₄-BTBP binding over
the 1:1 Ln³⁺/Cy-Me₄-BTBP complex, they do not significantly
alter the N-donor coordination environment. In contrast to the
CyMe₄-BTPhen structures, there is no clearly identifiable trend
between the M–N_{bipy} and M–N_{triazinyl} bond lengths for all of
the CyMe₄-BTBP complexes. This suggests that the greater
flexibility of the bipyridyl group, relative to the phenanthroline
group, allows minimal distinction between the triazinyl and
bipyridyl N atoms when coordinated to a Ln³⁺ ion.

XAS of Lanthanide-Extracted Species. XAS spectra were
obtained for Eu³⁺ and Tb³⁺ species formed by extraction from
an acidic aqueous phase into an organic phase containing an
excess of either CyMe₄-BTBP or CyMe₄-BTPhen in cyclo-
hexanone as a guide for speciation in a potential SANEX
process. Studies were also performed for potential GANEX-like
systems where the organic phase also included 30% TBP. XAS
spectra were obtained for the crystallographically characterized
solids [Eu(CyMe₄-BTPhen)₂(H₂O)]³⁺ (**3**) and [Tb(CyMe₄-
BTPhen)₂(H₂O)]³⁺ (**4**) for comparative purposes. The spectra
obtained show little difference between the extracted species
with or without the presence of TBP (Figures 11 and 12 and
the Supporting Information). This indicates that the presence
of TBP does not influence lanthanide speciation when used in a

Table 4. Crystal Data for Complexes 1–5

	[Pr(CyMe ₄ -BTPhen) ₂ (NO ₃) ₂ ·10H ₂ O (1)]	[Pr(CyMe ₄ -BTPhen) ₂ (NO ₃) ₂ ·1.63EtOH·0.75H ₂ O (2)]	[Eu(CyMe ₄ -BTPhen) ₂ (H ₂ O) ₂ (NO ₃) ₃ ·9H ₂ O (3)]	[Tb(CyMe ₄ -BTPhen) ₂ (H ₂ O)(NO ₃) ₃ ·9H ₂ O (4)]	[Yb(CyMe ₄ -BTPhen) ₂ (H ₂ O) ₂ (NO ₃) ₃ ·9H ₂ O (5)]
formula	C ₆₈ H ₉₆ N ₁₉ O ₁₉ Pr	C _{71.25} H _{87.25} N ₂₂ O _{20.38} Pr ₂	C ₆₈ H ₉₆ N ₁₉ O ₁₉ Eu	C ₆₈ H ₉₆ N ₁₉ O ₁₉ Tb	C ₆₈ H ₉₆ N ₁₉ O ₁₉ Yb
<i>M</i>	1624.55	1859.70	1615.44	1642.56	1656.68
cryst syst	monoclinic	triclinic	orthorhombic	orthorhombic	orthorhombic
<i>a</i> (Å)	31.654(5)	13.716(5)	31.172(3)	31.3486(7)	31.3257(13)
<i>b</i> (Å)	26.271(5)	15.221(5)	38.128(3)	38.0261(9)	37.709(2)
<i>c</i> (Å)	19.501(5)	20.359(5)	14.8296(13)	14.8414(3)	14.8783(7)
<i>α</i> (deg)	90	107.225(5)	90	90	90
<i>β</i> (deg)	109.504(5)	99.422(5)	90	90	90
<i>γ</i> (deg)	90	97.083(5)	90	90	90
space group	<i>C2/c</i>	<i>P</i> $\bar{1}$	<i>Fdd2</i>	<i>Fdd2</i>	<i>Fdd2</i>
<i>Z</i>	8	2	8	8	8
<i>T</i> (K)	100(2)	100(2)	100(2)	100(2)	100(2)
<i>μ</i> (mm ⁻¹)	0.719	1.309	0.782	0.870	1.135
reflms measd	20981	25422	32709	49820	8497
reflms obsd	6002	14178	8290	9044	5110
<i>R</i> ₁ (obsd)	0.0551	0.0547	0.0512	0.0513	0.0658
<i>wR</i> ₂ (all data)	0.1257	0.1393	0.1364	0.1423	0.1984

Table 5. Crystal Data for Complexes 6–10

	[Pr(CyMe ₄ -BTBP) ₂ (NO ₃) ₂ ·4EtOH·H ₂ O (6)]	[Pr(CyMe ₄ -BTBP) ₂ (NO ₃) ₂][Pr(NO ₃) ₆](NO ₃) ₆ ·6CH ₃ CN (7)]	[Eu(CyMe ₄ -BTBP) ₂ (NO ₃) ₂ (NO ₃) ₂ ·4EtOH·2H ₂ O (8)]	[Eu(CyMe ₄ -BTBP)(NO ₃) ₃ ·toluene (9)]	[Tb(CyMe ₄ -BTBP) ₂ (H ₂ O) ₂ (NO ₃) ₃ ·4EtOH (10)]
formula	C ₇₂ H ₁₀₂ N ₁₉ O ₁₄ Pr	C ₁₄₀ H ₁₆₆ N ₄₇ O ₂₇ Pr ₃	C ₇₂ H ₁₀₄ N ₁₉ O ₁₅ Eu	C ₄₆ H ₅₄ N ₁₁ O ₉ Eu	C ₇₂ H ₁₀₂ N ₁₉ O ₁₄ Tb
<i>M</i>	1598.64	3361.93	1627.70	1056.96	1614.63
cryst syst	monoclinic	monoclinic	orthorhombic	monoclinic	monoclinic
<i>a</i> (Å)	24.2790(7)	16.604(2)	16.4128(6)	26.385(2)	30.5621(7)
<i>b</i> (Å)	16.5467(4)	28.1161(19)	23.8916(6)	11.6674(11)	14.8217(4)
<i>c</i> (Å)	19.4601(5)	17.7385(14)	19.7838(6)	15.7469(14)	23.9083(6)
<i>α</i> (deg)	90	90	90	90	90
<i>β</i> (deg)	90.355(3)	106.609(10)	90	90.6730(10)	129.4280(10)
<i>γ</i> (deg)	90	90	90	90	90
space group	<i>P2</i> ₁ / <i>c</i>	<i>P2</i> ₁ / <i>n</i>	<i>Pccn</i>	<i>C2/c</i>	<i>C2/c</i>
<i>Z</i>	4	2	4	4	4
<i>T</i> (K)	100(2)	100(2)	100(2)	100(2)	100(2)
<i>μ</i> (mm ⁻¹)	0.697	0.988	0.884	1.359	4.723
reflms measd	77969	17453	47979	17887	25718
reflms obsd	13767	17454	6854	4618	7271
<i>R</i> ₁ (obsd)	0.0562	0.0766	0.1158	0.0430	0.0915
<i>wR</i> ₂ (all data)	0.1435	0.2209	0.2628	0.1164	0.2381

621 potential GANEX process with CyMe₄-BTBP or CyMe₄-
 622 BTPhen. The XAS profiles for the directly synthesized solid
 623 species (3 and 4) also correlate well with the corresponding
 624 extracted species (Figures 12 and 13), suggesting that the
 625 [Ln(CyMe₄-BTPhen)₂(H₂O)]²⁺ coordination species found in
 626 the solid state also exists in the bulk organic-phase
 627 postextraction.

628 The shells used to fit the EXAFS data for all samples were
 629 derived from the corresponding Ln^{III} structures that have two
 630 CyMe₄-BTX (X = BP, Phen) ligands bound to the metal. The

dominant scatter paths include a shell corresponding to the 8 N
 atoms from the CyMe₄-BTX ligands that are coordinated to the
 metal (~2.51 Å) and two shells from the 16 C/N and 16 C
 atoms located at the ortho and meta positions relative to the
 coordinating N atoms, respectively (Figure 14). The initial
 positions of these modeled shells, relative to the central Ln
 atom, are averaged from the atomic positions obtained from the
 crystal structures determined by XRD and are located at ~2.51,
 3.42, and 4.75 Å from the Ln atom for the 8 N, 16 C/N, and 16
 C shells, respectively (Figure 4 and Tables 8 and 9). It was

Table 6. Selected Interatomic Distances (\AA) for $\text{CyMe}_4\text{-BTPhen-Containing Complexes 1-5}^a$

bond	origin	1 (Pr)	2 (Pr)	3 (Eu)	4 (Tb)	5 (Yb)
N2-M	N _{triazinyl}	2.636(9)	2.644(6)	2.539(5)	2.527(5)	2.475(9)
N6-M		2.623(9)	2.635(6)	2.542(5)	2.516(5)	2.507(11)
N10-M		2.623(9)	2.568(6)	N/A	N/A	N/A
N14-M		2.618(8)	2.592(6)	N/A	N/A	N/A
N4-M	N _{phen}	2.668(8)	2.632(6)	2.507(5)	2.485(5)	2.422(11)
N5-M		2.644(9)	2.587(6)	2.523(5)	2.499(5)	2.442(10)
N12-M		2.675(8)	2.617(6)	N/A	N/A	N/A
N13-M		2.638(9)	2.583(6)	N/A	N/A	N/A
O1-M	O _{water}	N/A	N/A	2.414(6)	2.398(6)	2.373(11)
O1-M	O _{nitrate}	2.592(7)	2.581(5)	N/A	N/A	N/A
O2-M		2.544(7)	2.605(5)	N/A	N/A	N/A

^aN/A = not applicable. The designated bond length does not exist or is symmetry-related to another bond length.

Table 7. Selected Interatomic Distances (\AA) for $\text{CyMe}_4\text{-BTBP-Containing Complexes 6-10}^a$

bond	origin	6 (Pr)	7 (Pr)	8 (Eu)	9 (Eu)	10 (Tb)
N2-M	N _{triazinyl}	2.637(5)	2.654(9)	2.565(9)	2.533(4)	2.516(6)
N6-M		2.597(5)	2.595(9)	2.578(10)	N/A	2.512(6)
N10-M		2.611(5)	2.579(9)	N/A	N/A	N/A
N14-M		2.634(6)	2.597(9)	N/A	N/A	N/A
N4-M	N _{bipy}	2.638(5)	2.645(9)	2.569(8)	2.545(4)	2.485(5)
N5-M		2.623(5)	2.663(10)	2.562(9)	N/A	2.504(6)
N12-M		2.615(5)	2.650(9)	N/A	N/A	N/A
N13-M		2.633(5)	2.684(9)	N/A	N/A	N/A
O1-M	O _{water}	N/A	N/A	N/A	N/A	2.407(8)
O1-M	O _{nitrate}	2.596(5)	2.606(8)	2.564(10)	2.548(4)	N/A
O2-M		2.607(5)	2.625(8)	N/A	2.487(3)	N/A
O7-M		N/A	N/A	N/A	2.455(4)	N/A

^aN/A = not applicable. The designated bond length does not exist or is symmetry-related to another bond length.

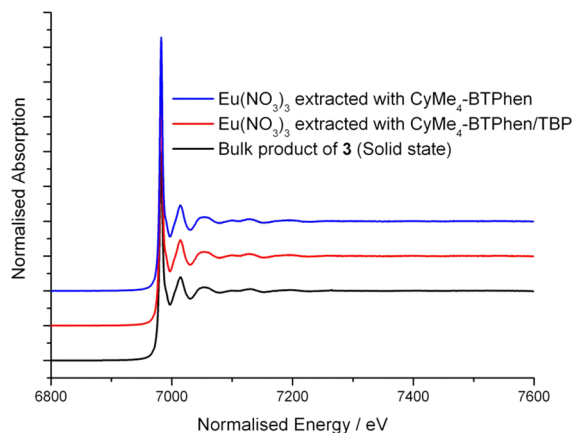


Figure 12. Eu L_{III} -edge XAS spectra of $\text{CyMe}_4\text{-BTPhen-Containing species}$.

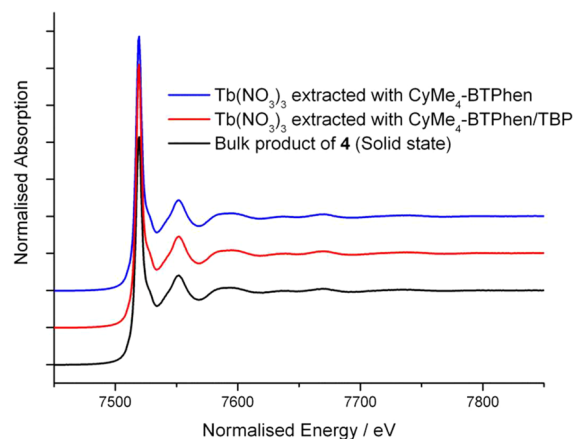


Figure 13. Tb L_{III} -edge XAS spectra of $\text{CyMe}_4\text{-BTPhen-Containing species}$.

641 found necessary to include an extra shell assigned to 32 C/N
 642 atoms, initially located at $\sim 3.40 \text{ \AA}$ from the Ln atom, due to
 643 multiple scattering from the planar aromatic rings in the N-
 644 donor ligands in order to obtain appropriate fits (Figure 14). A
 645 shell corresponding to oxygen coordination at the ninth site
 646 was included in all fits, initially located at $2.4\text{--}2.6 \text{ \AA}$ from the
 647 Ln atom. Two sets of models corresponding to nitrate
 648 coordination (i.e., O shell occupancy = 2) and water
 649 coordination (i.e., O shell occupancy = 1) to the lanthanide
 650 ion were used to fit all EXAFS data in order to ascertain
 651 whether XAS can be used to distinguish between nitrate and
 652 water binding in these systems (Tables 8 and 9; see the

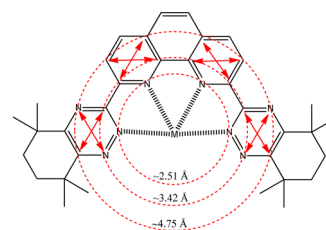


Figure 14. Depiction of the shell occupancy of complexes formed with $\text{CyMe}_4\text{-BTPhen}$ (or $\text{CyMe}_4\text{-BTBP}$). Also depicted with arrows are some of the multiple scatter paths within the complex.

Table 8. Eu L_{III}-Edge EXAFS Data^a

physical state	aqueous phase	organic-phase extractants	chemical composition used in fitted models	occupancy ^b	interatomic distances from XRD (Å)	fitted interatomic distances ^c (Å)	σ^2 (Å ²) ^d	r^e
solution	Eu(NO ₃) ₃	CyMe ₄ -BTBP	[Eu(CyMe ₄ -BTBP) ₂ (NO ₃) ₂] ²⁺	Eu–O2	2.56	2.46	0.00200	0.0159
				Eu–N8	2.57	2.59	0.00391	
				Eu–C/ N16	3.44	3.44	0.00731	
				Eu–C/ N32 ^f	3.67	3.68	0.00791	
				Eu–C/ N16	4.78	4.87	0.00374	
solution	Eu(NO ₃) ₃	CyMe ₄ -BTBP + TBP	[Eu(CyMe ₄ -BTBP) ₂ (NO ₃) ₂] ²⁺	Eu–O2	2.56	2.55	0.00567	0.0193
				Eu–N8	2.57	2.55	0.00594	
				Eu–C/ N16	3.44	3.45	0.00632	
				Eu–C/ N32 ^f	3.67	3.66	0.00796	
				Eu–C/ N16	4.78	4.87	0.00554	
solution	Eu(NO ₃) ₃	CyMe ₄ -BTPhen	[Eu(CyMe ₄ -BTPhen) ₂ (NO ₃) ₂] ²⁺	Eu–O2	2.56	2.60	0.00200	0.0206
				Eu–N8	2.51	2.55	0.00493	
				Eu–C/ N16	3.41	3.45	0.00496	
				Eu–C/ N32 ^f	3.65	3.69	0.00800	
				Eu–C/ N16	4.75	4.91	0.00303	
solution	Eu(NO ₃) ₃	CyMe ₄ -BTPhen + TBP	[Eu(CyMe ₄ -BTPhen) ₂ (NO ₃) ₂] ²⁺	Eu–O2	2.56	2.58	0.00432	0.0222
				Eu–N8	2.51	2.56	0.00486	
				Eu–C/ N16	3.41	3.45	0.00484	
				Eu–C/ N32 ^f	3.65	3.69	0.00800	
				Eu–C/ N16	4.75	4.91	0.00282	
solid	N/A	N/A	[Eu(CyMe ₄ -BTPhen) ₂ (H ₂ O)] ³⁺	Eu–O1	2.41	2.57	0.00121	0.0273
				Eu–N8	2.51	2.56	0.00504	
				Eu–C/ N16	3.41	3.44	0.00663	
				Eu–C/ N32 ^f	3.65	3.68	0.00800	
				Eu–C/ N16	4.75	4.92	0.00944	
			[Eu(CyMe ₄ -BTPhen) ₂ (NO ₃) ₂] ²⁺	Eu–O2	2.56	2.56	0.00193	0.0231
				Eu–N8	2.51	2.56	0.00667	
				Eu–C/ N16	3.41	3.43	0.00605	
				Eu–C/ N32 ^f	3.65	3.68	0.00800	
				Eu–C/ N16	4.75	4.91	0.00763	

^aS₀² is fitted but constrained to be within the range of 0.8–1.0 and the same value for all shells. ^bOccupancy numbers, held constant at given values. ^c±0.02 Å. ^dDebye–Waller factors. ^eParameter describing goodness of fit = weighted sum of squares of residuals divided by the degree of freedom. ^fShell due to multiple scattering.

Supporting Information). The EXAFS data fits were obtained by allowing the shell distances to be refined, while the shell occupancies were fixed at chosen integer values. Attempts were made to fit the EXAFS data to a model corresponding to the coordination of one molecule of CyMe₄-BTX with three nitrate molecules occupying the remaining coordination sites, but these did not give any statistically justifiable fits.

Fits for all of the EXAFS data obtained, using a model relating to the coordination of two CyMe₄-BTX molecules gave very good statistical correlations (Figure 15 and Tables 8 and 9; see the Supporting Information), indicating that the predominant lanthanide species in the bulk organic phase formed by extraction with these tetra-N-donor molecules is a 1:2 Ln³⁺/CyMe₄-BTX complex. The refined radial distances for the three closest N-donor ligand-based shells to the Ln center

Table 9. Tb L_{III}-Edge EXAFS Data^a

physical state	aqueous phase	organic-phase extractants	chemical composition used in fitted models	occupancy ^b	interatomic distances from XRD (Å)	fitted interatomic distances (Å) ^c	σ^2 (Å ²) ^d	r^e
solution	Tb(NO ₃) ₃	CyMe ₄ -BTBP	[Tb(CyMe ₄ -BTBP) ₂ (NO ₃) ₂] ²⁺	Tb–O2		2.53	0.00347	0.0140
				Tb–N8	2.50	2.51	0.00654	
				Tb–C/ N16	3.38	3.39	0.00530	
				Tb–C/ N32 ^f	3.61	3.65	0.00800	
				Tb–C/ N16	4.72	4.83	0.00165	
solution	Tb(NO ₃) ₃	CyMe ₄ -BTBP + TBP	[Tb(CyMe ₄ -BTBP) ₂ (NO ₃) ₂] ²⁺	Tb–O2		2.52	0.00365	0.0146
				Tb–N8	2.50	2.51	0.00675	
				Tb–C/ N16	3.38	3.40	0.00576	
				Tb–C/ N32 ^f	3.61	3.64	0.00800	
				Tb–C/ N16	4.72	4.83	0.00160	
solution	Tb(NO ₃) ₃	CyMe ₄ -BTPhen	[Tb(CyMe ₄ -BTPhen) ₂ (NO ₃) ₂] ²⁺	Tb–O2		2.52	0.00210	0.0133
				Tb–N8	2.52	2.52	0.00840	
				Tb–C/ N16	3.40	3.40	0.00570	
				Tb–C/ N32 ^f	3.62	3.66	0.00800	
				Tb–C/ N16	4.73	4.84	0.00190	
solution	Tb(NO ₃) ₃	CyMe ₄ -BTPhen + TBP	[Tb(CyMe ₄ -BTPhen) ₂ (NO ₃) ₂] ²⁺	Tb–O2		2.53	0.00204	0.0134
				Tb–N8	2.52	2.51	0.00835	
				Tb–C/ N16	3.40	3.40	0.00537	
				Tb–C/ N32 ^f	3.62	3.66	0.00800	
				Tb–C/ N16	4.73	4.84	0.00185	
solid	N/A	N/A	[Tb(CyMe ₄ -BTPhen) ₂ (H ₂ O)] ³⁺	Tb–O1	2.40	2.54	0.00196	0.0166
				Tb–N8	2.52	2.52	0.00620	
				Tb–C/ N16	3.40	3.41	0.00601	
				Tb–C/ N32 ^f	3.62	3.66	0.00800	
				Tb–C/ N16	4.73	4.85	0.00532	
			[Tb(CyMe ₄ -BTPhen) ₂ (NO ₃) ₂] ²⁺	Tb–O2		2.53	0.00199	0.0141
				Tb–N8	2.52	2.52	0.00772	
				Tb–C/ N16	3.40	3.40	0.00529	
				Tb–C/ N32 ^f	3.62	3.66	0.00800	
				Tb–C/ N16	4.73	4.84	0.00324	

^aS₀² is fitted but constrained to be within the range of 0.8–1.0 and the same value for all shells. ^bOccupancy numbers, held constant at given values. ^c±0.02 Å. ^dDebye–Waller factors. ^eParameter describing goodness of fit = weighted sum of squares of residuals divided by the degree of freedom. ^fShell due to multiple scattering.

668 (i.e., 8 N, 16 C/N, and 32 C/N shells) generally only show
669 minimal shifts from the initial input values derived from the
670 structural information obtained by XRD (Tables 8 and 9; see
671 the Supporting Information). The radial distance of the 16 C
672 shell does typically refine to a slightly larger value (4.80–4.95
673 Å) relative to the initial input value (4.70–4.80 Å) for the
674 extracted solutions and solid-state samples. This suggests that
675 either this outer C shell is influenced by multiple scattering

676 effects or some fluctuation of the N₄-donor ligand occurs at the
677 outer regions of these lanthanide complexes. No significant
678 differences in the refined radial distances are observed in the
679 extracted samples when TBP is present or not, providing
680 further proof that TBP that does not influence Ln³⁺ speciation
681 in a GANEX process with CyMe₄-BTX and TBP in the organic
682 phase.

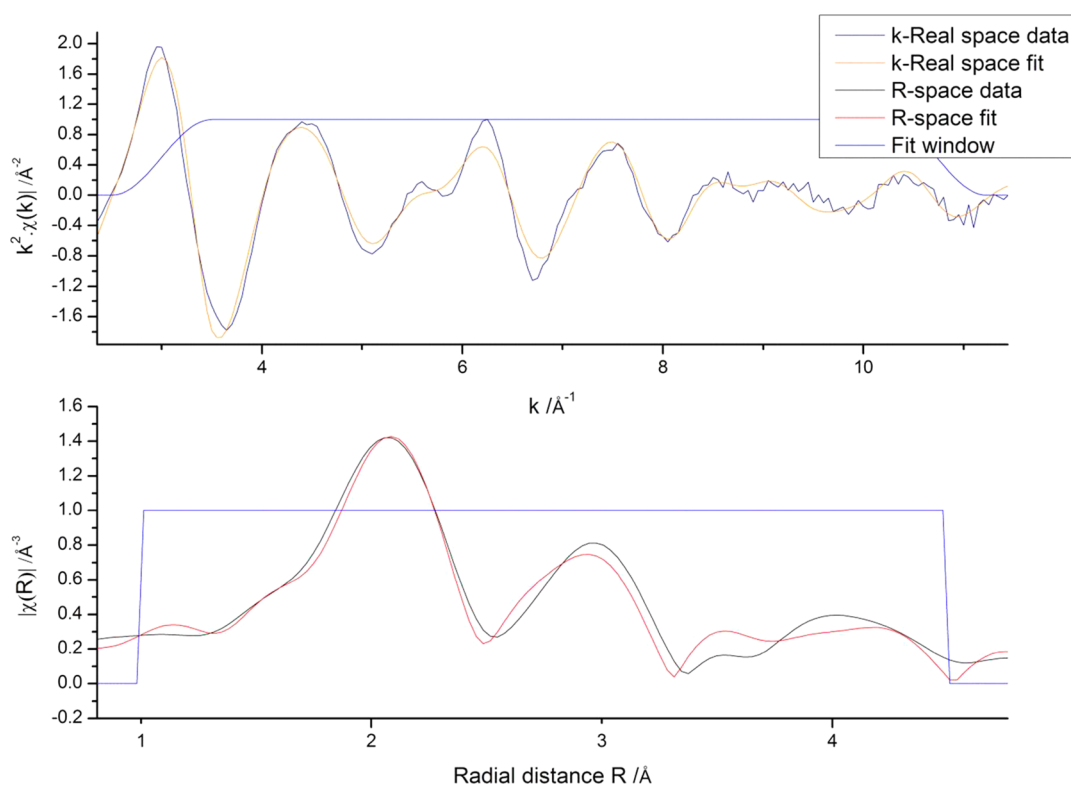


Figure 15. Eu L_{III}-edge EXAFS spectrum in *k* space (upper plot) and its Fourier transform in *R* space (lower plot) of the extraction of Eu(NO₃)₃ (10 mM) from an aqueous solution (1 M HNO₃ and 3 M NaNO₃) into cyclohexanone with CyMe₄-BTPPhen (50 mM). The data are fitted to the model complex [Eu(CyMe₄-BTPPhen)₂(NO₃)₂]²⁺.

683 The identity of the ninth coordination site species cannot be
 684 unambiguously assigned from the EXAFS data because the fits
 685 are unable to resolve the relatively small change between nitrate
 686 and water coordination at this site. Both sets of models, either
 687 with water or nitrate bound at the ninth coordination site,
 688 provided fits with very good statistical correlations (Tables 8
 689 and 9; see the Supporting Information). The fits for all of the
 690 Eu L_{III}-edge data (Table 8; see the Supporting Information)
 691 show that the first O shell refines to give Eu–O distances
 692 between 2.46 and 2.60 Å. There is little distinction between the
 693 refined Eu–O distances when the shell occupancy is fixed at 1
 694 (for water coordination) or 2 (for nitrate coordination). The
 695 refined Eu–O_{nitrate} distances agree with those bond lengths
 696 determined for structures 8 [2.56(1) Å] and 9 [2.548(4) Å]
 697 and fall within the range of all known Eu–O_{nitrate} distances
 698 (2.31–2.82 Å) established by crystallography.^{35,38} The refined
 699 Eu–O_{water} distances also fall within the wide range of Eu–O_{water}
 700 bond lengths from previously reported structures (2.27–2.72
 701 Å)^{35,39} but are larger than this distance in complex 3 [2.414(6)
 702 Å]. The O shells for all of the Tb L_{III}-edge EXAFS spectra
 703 modeled with either water or nitrate coordination (Table 9; see
 704 the Supporting Information) all refine to within a narrow range
 705 of 2.52–2.54 Å from the Tb center and fall within the relatively
 706 wide range of known Tb–O_{nitrate} (2.19–2.85 Å)^{35,40} and Tb–
 707 O_{water} bond lengths (2.27–2.70 Å).^{35,39} As is similarly observed
 708 in the equivalent europium studies, all of the Tb–O_{water}
 709 distances refined from the EXAFS data are larger than that
 710 observed in complex 4 [2.397(6) Å] by XRD. The refined
 711 radial distances for these low-occupancy O shells generally
 712 match the refined location for the dominant 8 N shell from the
 713 coordinating N-donor ligands even when these distances are
 714 expected to be different. This is particularly evident in the

EXAFS fits of complexes 3 and 4 in the solid state, which have
 also been characterized by XRD. The refined distances for the
 O shells obtained from the fits of the EXAFS data in these
 solid-state samples (Eu–O for 3 = 2.57 Å; Tb–O for 4 = 2.54
 Å) are distinctly longer than those observed by XRD [Eu–O
 for 3 = 2.414(6) Å; Tb–O for 4 = 2.397(6) Å] and are similar
 to the refined radial distances of the 8 N shell (Eu–N for 3 =
 2.56 Å; Tb–N for 4 = 2.52 Å). It may be that the high-
 occupancy 8 N shell is masking the contribution of the lower-
 occupancy (1 or 2) O shell.

EXPERIMENTAL SECTION

General Procedures. Elemental analyses were performed with a
 Carlo Erba Instruments CHNS-O EA1108 elemental analyzer for
 carbon, hydrogen, and nitrogen and a Fisons Horizon elemental
 analysis ICP-OES spectrometer for praseodymium, europium, and
 terbium. ESI-MS (positive ion) was performed using a Micromass
 Platform spectrometer. Solution UV–visible spectra were recorded on
 a PG Instruments T60U spectrophotometer with a fixed spectral
 bandwidth of 2 nm. Typical scan ranges were 200–500 nm at a scan
 rate of ~390 nm min⁻¹. Excitation and emission spectra were recorded
 with Edinburgh Instrument FP920 phosphorescence lifetime spec-
 trometer equipped with a 5 W microsecond-pulsed xenon flashlamp
 (with single 300 mm focal length excitation and emission
 monochromators in a Czerny Turner configuration) and a red-
 sensitive photomultiplier in Peltier (air-cooled) housing (Hamamatsu
 R928P) using a gate time of 0.05 ms and a delay time of 0.5 ms.
 Excitation spectra were obtained using the following emission
 wavelengths: Eu³⁺, 616 nm; Tb³⁺, 545 nm. Lifetime data were
 recorded following 320 nm excitation with a microsecond-pulsed
 xenon flashlamp (Edinburgh Instruments) using the multichannel
 scaling method. Lifetimes were obtained by a tail fit on the data
 obtained, and the quality of the fit was judged by minimization of
 reduced χ^2 and residuals squared. Where the decay profiles are

748 reported as monoexponential, fitting to a double-exponential decay
749 yielded no improvement in the fit, as judged by minimization of
750 residuals squared and reduced χ^2 .

751 **Syntheses and Solution Preparations.** All chemicals were
752 purchased from Sigma Aldrich and were used as supplied. CyMe₄-
753 BTPPhen and CyMe₄-BTBP were synthesized as previously de-
754 scribed^{20,23} but using an improved purification methodology.

755 **Purification of CyMe₄-BTPPhen and CyMe₄-BTBP.** The crude
756 product of CyMe₄-BTPPhen or CyMe₄-BTBP was dissolved in DCM
757 and loaded onto a column of silica resin. The column was washed with
758 neat DCM, quickly eluting a yellow solution and leaving a dark-
759 orange/brown band at the top of the column. The solvent was
760 removed from the yellow eluent by rotary evaporation, yielding a
761 vibrant-yellow powder, which was found to be pure by NMR
762 spectroscopy. The purified product was found to be soluble in DCM,
763 cyclohexanone, and 1-octanol up to concentrations of 5 mM. Further
764 purified CyMe₄-BTPPhen/CyMe₄-BTBP could be obtained by elution
765 with 1–5% (v/v) MeOH in DCM from the silica column.

766 **Synthesis of Pr³⁺ Complexes with CyMe₄-BTPPhen.** A solution
767 of Pr(NO₃)₃·6H₂O (23 mg, 54 μmol) in CH₃CN (5 mL) was added
768 to a solution of CyMe₄-BTPPhen (30 mg, 54 μmol) in DCM (5 mL)
769 and left standing to evaporate to dryness. The resultant powder was
770 dissolved in a mixture of CH₃CN (2 mL), DCM (2 mL), and EtOH
771 (0.5 mL) and again allowed to evaporate slowly in order to crystallize.
772 A yellow platelike crystal was selected from the isolated material, and
773 XRD analysis indicated that the composition of the crystal was of the
774 formulation [Pr(CyMe₄-BTPPhen)₂(NO₃)₂](NO₃)₂·10H₂O (1·10H₂O).
775 Elemental analysis of the isolated material indicated that the
776 composition of the bulk product was of the formulation [Pr(CyMe₄-
777 BTPPhen)₂(NO₃)₂][Pr(NO₃)₃·2H₂O (2·2H₂O)]. Elem anal. Calcd for
778 [(C₃₄H₃₈N₈)₂(NO₃)₂Pr][Pr(NO₃)₃·2H₂O]: C, 45.19; H, 4.46; N,
779 17.05; Pr, 15.59. Found: C, 45.01; H, 4.08; N, 16.90; Pr, 15.23. The
780 bulk material was dissolved in EtOH (1 mL) and allowed to slowly
781 evaporate over 1 week, yielding yellow blocklike crystals suitable for
782 single-crystal XRD analysis (yield = 0.03 g). ESI-MS (positive ion): *m/z*
783 *z* 659 ([C₃₄H₃₈N₈)₂(NO₃)₂Pr]²⁺). UV–visible spectrum (MeOH)
784 [λ_{max}/nm (ε_{max}/L mol⁻¹ cm⁻¹): 266 (71000), 321 (38000)].

785 **Synthesis of [Eu(CyMe₄-BTPPhen)₂(H₂O)](NO₃)₃·2H₂O
786 (3·2H₂O).** A solution of Eu(NO₃)₃·6H₂O (24 mg, 54 μmol) in
787 CH₃CN (5 mL) was added to a solution of CyMe₄-BTPPhen (30 mg,
788 54 μmol) in DCM (5 mL) and left standing to evaporate to dryness.
789 The resultant powder was dissolved in a mixture of CH₃CN (2 mL),
790 DCM (2 mL), and EtOH (0.5 mL) and allowed to evaporate slowly,
791 yielding yellow blocklike crystals suitable for single-crystal XRD
792 analysis (yield = 0.02 g). Elem anal. Calcd for [(C₃₄H₃₈N₈)₂(H₂O)-
793 Eu](NO₃)₃·2H₂O: C, 54.11; H, 5.48; N, 17.63; Eu, 10.07. Found: C,
794 54.18; H, 5.07; N, 17.61; Eu, 10.51. ESI-MS (positive ion): *m/z* 666
795 ([C₃₄H₃₈N₈)₂(NO₃)₂Eu]²⁺). UV–visible spectrum (MeOH) [λ_{max}/
796 nm (ε_{max}/L mol⁻¹ cm⁻¹): 266 (99000), 321 (52000)].

797 **Synthesis of [Tb(CyMe₄-BTPPhen)₂(H₂O)](NO₃)₃·H₂O (4·H₂O).**
798 The synthesis was performed as described for 2 except using
799 Tb(NO₃)₃·5H₂O (17 mg, 38 μmol) and CyMe₄-BTPPhen (21 mg,
800 38 μmol) as the initial reagents. Yellow platelike crystals suitable for
801 single-crystal XRD analysis were obtained (yield = 0.02 g). Elem anal.
802 Calcd for [(C₃₄H₃₈N₈)₂(H₂O)Tb](NO₃)₃·H₂O: C, 54.51; H, 5.38; N,
803 17.76; Tb, 10.61. Found: C, 54.69; H, 5.17; N, 17.73; Tb, 9.82. ESI-
804 MS (positive ion): *m/z* 669 ([C₃₄H₃₈N₈)₂(NO₃)₂Tb]²⁺). UV–visible
805 spectrum (MeOH) [λ_{max}/nm (ε_{max}/L mol⁻¹ cm⁻¹): 265 (96000), 322
806 (51000)].

807 **Synthesis of [Yb(CyMe₄-BTPPhen)₂(H₂O)](NO₃)₃·3H₂O
808 (5·3H₂O).** The synthesis was performed as described for 2 except
809 using Yb(NO₃)₃·5H₂O (24 mg, 54 μmol) and CyMe₄-BTPPhen (30
810 mg, 54 μmol) as the initial reagents. Yellow rhombohedron-like
811 crystals suitable for single-crystal XRD analysis were obtained (yield <
812 0.01 g). ESI-MS (positive ion): *m/z* 677 ([C₃₄H₃₈N₈)₂(NO₃)₂Yb]²⁺).
813

814 **Synthesis of Ln³⁺ Complexes with CyMe₄-BTBP.** A solution of
815 CyMe₄-BTBP (30 mg, 56 μmol) in DCM (1 mL) was added to a
816 solution of Ln(NO₃)₃·xH₂O [Pr(NO₃)₃·6H₂O, 12 mg, 28 μmol;
817 Eu(NO₃)₃·6H₂O, 13 mg, 28 μmol; Tb(NO₃)₃·5H₂O, 12 mg, 28 μmol]
818 in MeOH (1 mL). CH₃CN (1.5 mL) was added to the reaction

819 mixture, and the solution was allowed to evaporate to dryness. Once
820 dry, toluene (1.25 mL), EtOH (1.25 mL), ⁱPrOH (1.25 mL), and
821 DCM (1.25 mL) were added to dissolve the residues, and the
822 solutions were allowed to evaporate slowly. Crystals suitable for single-
823 crystal XRD were obtained over several weeks. The mixtures afforded a
824 variety of crystals of varying compositions determined by single-crystal
825 XRD analysis to be [Pr(CyMe₄-BTBP)₂(NO₃)₂](NO₃)₂·4EtOH·H₂O
826 (6·4EtOH·H₂O), [Pr(CyMe₄-BTBP)₂(NO₃)₂][Pr(NO₃)₃·6H₂O]
827 (NO₃)₂·6CH₃CN (7·6CH₃CN), [Eu(CyMe₄-BTBP)₂(NO₃)₂]-
828 (NO₃)₂·4EtOH·2H₂O (8·4EtOH·2H₂O), [Eu(CyMe₄-BTBP)-
829 (NO₃)₃]₂·toluene (9·toluene), and [Tb(CyMe₄-BTBP)₂(H₂O)]-
830 (NO₃)₂·4EtOH (10·4EtOH). Bulk analysis of the crystallized
831 samples by ESI-MS provided the following data:-

832 ESI-MS (positive ion): Pr³⁺ complexation, *m/z* 635
833 ([C₃₂H₃₈N₈)₂(NO₃)₂Pr]²⁺); Eu³⁺ complexation, *m/z* 641
834 ([C₃₂H₃₈N₈)₂(NO₃)₂Eu]²⁺); Tb³⁺ complexation, *m/z* 643
835 ([C₃₂H₃₈N₈)₂(NO₃)₂Tb]²⁺).
836

837 **Solution Preparation for UV–visible Spectroscopic Studies
838 of Ln³⁺ Complexation with CyMe₄-BTPPhen and CyMe₄-BTBP.**
839 Methanolic solutions of the ligands CyMe₄-BTBP and CyMe₄-BTPPhen
840 (1.0 × 10⁻⁴ M, 0.4 mL) were added to a quartz cuvette of 1 cm path
841 length, and the solutions were diluted to 2 mL with MeOH (2.0 ×
842 10⁻⁵ M). At this point, an initial spectrum of the ligand was recorded.
843 Metal solutions of Eu(NO₃)₃·6H₂O, Pr(NO₃)₃·6H₂O, and Tb-
844 (NO₃)₃·5H₂O (4.0 × 10⁻⁴ M) in MeOH were used. For each
845 titration, the metal solution was added into the cuvette in 10 μL (4.0 ×
846 10⁻⁹ mol, 0.10 equiv) aliquots and shaken, and spectra were recorded
847 after each addition up to a ratio of 1.5:1 metal/ligand. At this point,
848 the aliquot size was increased to 50 μL (0.50 equiv) to a final ratio of
849 3:1 metal/ligand.
850

851 **Solution Preparation for Luminescence Studies of Ln³⁺ (Ln
852 = Pr, Tb, Eu) Complexation with CyMe₄-BTPPhen and CyMe₄-
853 BTBP.** A solution of CyMe₄-BTPPhen/CyMe₄-BTBP in MeOH (120
854 μL, 1 × 10⁻⁴ M) was added to a 1.2 mL quartz cuvette followed by the
855 addition of a solution of Ln(NO₃)₃ in MeOH (20 μL, 3 × 10⁻⁴ M).
856 The solution was diluted to ~1 mL with MeOH, and spectra were
857 obtained.
858

859 Solution samples in MeOH-*d*₄ were prepared in the same manner as
860 that for the MeOH samples but using a 6 × 10⁻⁴ M solution of
861 CyMe₄-BTPPhen/CyMe₄-BTBP (20 μL) in MeOH-*d*₄, and solutions
862 were diluted using MeOH-*d*₄.
863

864 **Extracted Sample Preparation for XAS Measurements.**
865 Predistilled cyclohexanone and a 30% (v/v) solution of TBP in
866 cyclohexanone were “washed” before use according to previously
867 outlined procedures.⁴¹ The washing of the organic solvent took place
868 days before lanthanide extractions were performed. The extractants
869 CyMe₄-BTBP and CyMe₄-BTPPhen were dissolved in either solvent
870 system by gentle warming and sonication to a final extractant
871 concentration of 50 mM. Aqueous stock solutions of Ln(NO₃)₃ (Ln =
872 Pr, Eu, Tb; 10 mM) were prepared by dissolution of the relevant salt
873 in 4 M HNO₃ in deionized H₂O for extractions with 30% TBP/
874 cyclohexanone, while an aqueous mixture of 1 M HNO₃ and 3 M
875 NaNO₃ in deionized water was used for extractions with pure
876 cyclohexanone because of previously reported miscibility issues.²²
877

878 The extractions were performed using 1.0 mL of each phase
879 (organic and aqueous) contained in a 2.5 mL sample vial. The phases
880 were mixed using a Labinco L46 shaker for 5 min each. Once
881 contacted, each sample had the (lower) aqueous layer syringed out of
882 the vial and then the (upper) organic layer pipetted into another vial
883 for storage before XAS measurements were performed.
884

885 **Solid Sample Preparation for XAS Measurements.** Solid
886 samples of 2–4 were prepared for XAS measurements by crushing
887 ~5–6 mg of the crystalline material in a mortar and pestle and mixed
888 thoroughly with ~90 mg of BN. The homogeneous material was then
889 pressed into flat disks (~2 cm diameter).
890

891 **X-ray Crystallography.** Diffraction data for 1·10H₂O,
892 2·1.63EtOH·0.75H₂O, 3·9H₂O, 4·9H₂O, 5·9H₂O, 6·4EtOH·H₂O,
893 7·6CH₃CN, 8·4EtOH·2H₂O, 9·toluene, and 10·4EtOH were meas-
894 ured at 100 K with either a Bruker APEX SMART platform CCD area
895 Mo Kα diffractometer (2, 3, and 9), an Oxford Diffraction XCalibur2
896

888 Mo $K\alpha$ diffractometer (1 and 4–8), or a Bruker APEX2 Cu $K\alpha$
889 diffractometer (10). All were equipped with a low-temperature device,
890 and collections were performed at 100 K. *CryAlisPro* was used to guide
891 the Oxford diffractometer for collection of a full set of diffraction
892 images and perform unit cell determination and data reduction. These
893 data were corrected for Lorenz and polarization factors, and analytical,
894 multiscan, and absorption corrections were applied. Bruker SMART
895 (Mo $K\alpha$) or APEX2 (Cu $K\alpha$) was used to guide the Bruker
896 diffractometers and perform unit cell determinations.⁴² Reduction of
897 the Bruker collected data was performed using *SAINTE PLUS* (Mo $K\alpha$)
898 or APEX2 (Cu $K\alpha$), and a multiscan absorption correction was
899 performed using *SADABS*.^{43,44} For all crystal data, the structures were
900 solved by direct methods using *SIR92*.⁴⁵ Structure refinement was
901 achieved via full-matrix least squares based on F^2 using *SHELXL97*.⁴⁶
902 All non-H atoms not exhibiting disorder were refined anisotropically,
903 while H atoms were included in calculated positions. Molecular
904 graphics were generated using *ORTEP*, and all displayed plots show
905 probability ellipsoids of 50%.⁴⁷ In the case of structure 10, modeling of
906 residual solvent molecules was not possible. As such, the *SQUEEZE*
907 procedure in *PLATON* was used to obtain solvent-free reflection data,
908 and subsequent refinement was performed on these data. The *PART*
909 command was used to model disorder over multiple sites, where
910 appropriate, and is detailed in the relevant CIF (crystallographic
911 information) files (see the Supporting Information).

912 **General XAS Measurements.** Ln (Eu and Tb) L_{III} -edge XAS
913 spectra of extracted solutions and crystalline solids were recorded in
914 transmission and fluorescence modes on Beamline B18 at the
915 Diamond Light Source operating in a 10 min top-up mode for a
916 ring current of 300 mA and an energy of 3 GeV. The radiation was
917 monochromated with a Si(111) double crystal, and harmonic rejection
918 was achieved through the use of two platinum-coated mirrors
919 operating at an incidence angle of 8.3 mrad. The monochromator
920 was calibrated using the K-edge of an iron foil, taking the first
921 inflection point in the Fe-edge as 7112 eV. Spectra obtained in
922 fluorescence mode utilized a nine-element germanium detector. The
923 spectra were summed and background-subtracted using the software
924 package *Athena*.⁴⁸ The spectra were simulated using the software
925 package *Artemis*, which utilizes the Feff database in its simulations.^{48,49}

926 ■ CONCLUSIONS

927 The successful characterization of a series of directly
928 synthesized Ln^{III} complexes of the tetra-N-donor extractants
929 CyMe₄-BTPhen and CyMe₄-BTBP using XRD for solid-state
930 studies and solution electronic spectroscopy has provided
931 robust chemical models, which can be used to assist in the
932 determination of lanthanide species formed under proposed
933 conditions for the partitioning of SNF. Fits of the EXAFS
934 region from XAS spectra showed that the dominant species
935 extracted into the organic phase were complex, where two N₄-
936 donor extractant ligands were coordinated to the Ln³⁺ center, as
937 is mainly observed in the direct synthesis studies. XAS was
938 unable to elucidate the bound ligand at the ninth coordination
939 site in these Ln³⁺ complexes, but luminescence spectroscopy
940 indicates that nitrate coordination is preferred over water
941 binding in organic solvents. The presence of TBP in the organic
942 phase, which may be used in a potential GANEX separation,
943 clearly showed no influence with regards to lanthanide
944 speciation. Further work will assess the source of the high
945 separation factors that these N-donor ligands exhibit for minor
946 actinide/lanthanide partitioning. Similar speciation studies for
947 extracted Am³⁺ and Cm³⁺ in the bulk organic phase will be
948 performed to determine if minor actinide complexes analogous
949 to those observed in the lanthanide studies are formed or
950 whether separation is achieved by the formation of minor
951 actinide species that are substantially different [e.g., charge-
952 neutral tris(nitrate) complex molecules] from those of the

lanthanides. Such studies have been performed for BTP-derived
extractants and indicate little difference between Eu³⁺ and Cm³⁺
speciation,⁵⁰ but this needs to be confirmed for the N₄-donor
extractants particularly with respect to the role of nitrate ions as
the lanthanide series is traversed in minor actinide/lanthanide
coordination. Studies investigating metal speciation at the
interfacial region in these liquid–liquid separations will also be
conducted to assess the mechanism by which the minor
actinides preferentially cross from the aqueous phase into the
organic phase using these organic-soluble N-donor extractants
and whether actinide/lanthanide speciation in the bulk organic
phase is different from that at the liquid–liquid interface.
Understanding the molecular-scale processes that underpin
techniques for the partitioning of SNF will provide improved
development of advanced separation methodologies like
SANEX and GANEX.

■ ASSOCIATED CONTENT

Supporting Information

Plots of single-crystal XRD structures, crystallographic
information files (CIF), UV–visible absorption, excitation
and emission spectra, ORTEP plots, and XAS spectra with
corresponding fits and parameters for the EXAFS data region.
This material is available free of charge via the Internet at
<http://pubs.acs.org>.

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Notes

The authors declare no competing financial interest.

■ ACKNOWLEDGMENTS

This work is funded by the RCUK Energy Programme through
its support of the MBASE consortium and the Nuclear Fission
Safety Program of the European Union through ACSEPT
Contract FP7-CP-2007-211 267. The studentship for D.M.W.
was provided by the EPSRC-funded Nuclear FiRST Doctoral
Training Centre. We thank Diamond Light Source for access to
Beamline B18 (SP7226), which contributed to the results
presented here. We acknowledge use of the EPSRC-funded
Chemical Database Service at Daresbury. We also thank Dr.
John Charnock for his assistance with EXAFS simulations. Use
of the Chemical Analysis Facility at the University of Reading
and access to the Mass Spectrometry Service and the
Microanalysis Laboratory at the University of Manchester are
also acknowledged.

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