Northumbria Research Link

Citation: Lewis, Frank, Harwood, Laurence, Hudson, Michael, Drew, Michael, Desreux, Jean, Vidick, Geoffrey, Bouslimani, Nouri, Modolo, Giuseppe, Wilden, Andreas, Sypula, Michal, Vu, Trong and Simonin, Jean (2011) Highly efficient separation of actinides from lanthanides by a phenanthroline-derived bis-triazine ligand. Journal of the American Chemical Society, 133 (33). pp. 13093-13102. ISSN 0002-7863

Published by: American Chemical Society

URL: http://dx.doi.org/10.1021/ja203378m < http://dx.doi.org/10.1021/ja203378m >

This version was downloaded from Northumbria Research Link: https://nrl.northumbria.ac.uk/id/eprint/10881/

Northumbria University has developed Northumbria Research Link (NRL) to enable users to access the University's research output. Copyright © and moral rights for items on NRL are retained by the individual author(s) and/or other copyright owners. Single copies of full items can be reproduced, displayed or performed, and given to third parties in any format or medium for personal research or study, educational, or not-for-profit purposes without prior permission or charge, provided the authors, title and full bibliographic details are given, as well as a hyperlink and/or URL to the original metadata page. The content must not be changed in any way. Full items must not be sold commercially in any format or medium without formal permission of the copyright holder. The full policy is available online: http://nrl.northumbria.ac.uk/policies.html

This document may differ from the final, published version of the research and has been made available online in accordance with publisher policies. To read and/or cite from the published version of the research, please visit the publisher's website (a subscription may be required.)





Highly Efficient Separation of Actinides from Lanthanides by a Phenanthroline-Derived Bis-Triazine Ligand

Frank W. Lewis,[†] Laurence M. Harwood, *[†] Michael J. Hudson,[†] Michael G. B. Drew,[†] Jean F. Desreux,[‡] Geoffrey Vidick,[‡] Nouri Bouslimani,[‡] Giuseppe Modolo,[§] Andreas Wilden,[§] Michal Sypula,[§] Trong-Hung Vu,^{||} and Jean-Pierre Simonin^{||}

Department of Chemistry, University of Reading, Whiteknights, Reading RG6 6AD, UK, Coordination and Radiochemistry, University of Liège, Sart Tilman B16, B-4000 Liège, Belgium, Forschungszentrum Jülich GmbH, Sicherheitsforschung und Reaktortechnik, D-52425 Jülich, Germany, and Université Pierre et Marie Curie, Laboratoire PECSA (UMR CNRS 7195), Case 51, 4 Place Jussieu, 75252 Paris Cedex 05, France

E-mail: lim.harwood@reading.ac.uk

RECEIVED DATE

Highly Efficient Separation of Actinides from Lanthanides

[†] University of Reading, [‡] University of Liège, [§] Forschungszentrum Jülich GmbH, ^{||} Université Pierre et Marie Curie.

Abstract. The synthesis, lanthanide complexation and solvent extraction of actinide(III) and lanthanide(III) radiotracers from nitric acid solutions by a phenanthroline-derived quadridentate bis-

triazine ligand is described. The ligand separates Am(III) and Cm(III) from the lanthanides with remarkably high efficiency, high selectivity and fast extraction kinetics compared to its 2,2'-bipyridine counterpart. Structures of the 1:2 bis complexes of the ligand with Eu(III) and Yb(III) were elucidated by X-ray crystallography and force field calculations, respectively. The Eu(III) bis-complex is the first 1:2 bis-complex of a quadridentate bis-triazine ligand to be characterized by crystallography. The faster rates of extraction were verified by kinetics measurements using the rotating membrane cell technique in several diluents. The improved kinetics of metal ion extraction are related to the higher surface activity of the ligand at the phase interface. The improvement in the ligands properties on replacing the bipyridine unit with a phenanthroline unit far exceeds what was anticipated based on ligand design alone.

Keywords: actinides, extraction, ligand design, phenanthroline, lanthanides, sustainable chemistry.

Introduction

During the production of electricity from nuclear power, a typical 1000 MW(e) light water reactor produces 20-30 t of spent nuclear fuel per annum.¹ This spent fuel consists mainly (>98.5 %) of uranium and short-lived fission products which do not pose a long-term hazard. However, approximately 1 wt % of the spent fuel is composed of plutonium and the minor actinides (Am, Cm, Np), which are highly radiotoxic. Plutonium is the main contributor to the long-term radiotoxicity of spent nuclear fuel. Its separation through the PUREX (Plutonium and URanium EXtraction) process and reuse as fuel (MOX fuel) in nuclear reactors is current industrial practice.² The remaining waste still contains the minor actinides necessitating the containment and separation of the waste from the biosphere for many thousands of years, a situation that represents a serious environmental concern.

If the minor actinides could be removed, the mandatory storage time of the remaining waste decreases from several thousand years to a few hundred years. One strategy for reducing the radiotoxicity of the waste involves partitioning and transmutation of the long-lived minor actinides into shorter-lived or stable elements by neutron fission.³ However, it is first necessary to separate the actinides from the bulk

of the lanthanides and other fission and corrosion products that are also present in the acidic PUREX waste streams because the lanthanides have high neutron capture cross-sections.⁴ The separation of the radioactive minor actinides from the lanthanides is therefore one of the principal current challenges in nuclear waste reprocessing. This separation is made all the more difficult, given the chemical similarities between the two groups of elements.⁵ However, it is thought that the greater availability of the valence orbitals in the actinides mean that there is a more covalent contribution to metal-ligand bonding than with the lanthanides. The origins of this covalency are still not fully understood, and remain the subject of ongoing debate and study.⁶

It has been shown that ligands with soft *N*-donor atoms can exploit this small but significant difference between the actinides and lanthanides, and many *N*-heterocyclic ligands have been investigated for their ability to carry out this separation by solvent extraction.⁷ Two ligand classes have emerged as the most promising; the terdentate 2,6-bis(1,2,4-triazin-3-yl)pyridine ligands (BTPs)⁸ and the quadridentate 6,6'-bis(1,2,4-triazin-3-yl)-2,2'-bipyridine ligands (BTBPs). The BTBP ligands are currently the most suitable ligands for these separations.⁹ The annulated BTBP **1** has extraction and back-extraction properties which could be suitable for a continuous separation process (Figure 1).^{10,11} Unlike the BTBPs bearing alkyl side chains on the triazine rings, **1** is stable to hydrolysis and resistant towards radiolysis and degradation by free radicals because the labile benzylic hydrogens have been replaced by methyl groups.¹² Recently, some unusual uranyl (UO₂²⁺) complexes of **1** have been isolated and characterized.¹³



Figure 1. Conformations of the annulated BTBP ligand 1 and structure of 2,9-di(2-pyridyl)-1,10-phenanthroline **2**.

However, the rates of extraction of the actinides by 1 was rather slow and a phase-transfer agent (N,N'-dimethyl-N,N'-dioctyl-2-(2-hexoxyethyl) malondiamide) was required to improve the kinetics of extraction.¹⁰ This represents a disadvantage for the use of 1 in the proposed Generation (IV) fast reactors. Since a conformational change by 1 from the *trans*-conformation to its less-favoured *cis*-conformation is required prior to metal ligation, we hypothesized that this need for molecular reorientation was responsible for the slow rates of extraction by 1.

The *cis*-locked 1,10-phenanthrolines are versatile ligands in a wide range of applications. Their rigidity and the juxtaposition of their donor atoms means that complex formation is both more rapid and more thermodynamically favoured than their 2,2'-bipyridine analogues.¹⁴ For example, it has been reported that the pre-organized, phenanthroline-based ligand 2^{15} exhibits higher levels of complex stability and selectivity towards Cd(II) than related ligands such as quaterpyridine.¹⁶ In addition, phenanthrolines have larger dipole moments than the 2,2'-bipyridines,¹⁷ and often coordinate strongly to water via hydrogen bonding.¹⁸ These properties could lead to improved extraction kinetics in a separation process since the ligand may interact more favourably with the organic/water interface. Based on this, we predicted that a *cis*-locked quadridentate bis-triazine ligand containing a 1,10-phenanthroline

moiety would show improved extraction properties compared to its 2,2'-bipyridine counterpart **1**. Herein, we report the synthesis, lanthanide complexation and solvent extraction chemistry of the first of a new class of bis-triazine ligand in which the 2,2'-bipyridine moiety of **1** has been replaced by a 1,10phenanthroline moiety.

Results and Discussion

Synthesis and X-ray Crystallography. The new ligand **9** was synthesized as shown in Scheme 1. The dinitrile **6** was synthesized from 2,9-dimethyl-1,10-phenanthroline **3** following known procedures.¹⁹ The conversion of dioxime **5** to dinitrile **6** gave a low yield (37 %) of product in our hands. Consequently, we developed a one-pot method for the synthesis of the dinitrile **6** from the dialdehyde **4**. The reaction of the dinitrile **6** with hydrazine hydrate in EtOH gave the novel diamide dihydrazide **7** in near quantitative yield. The condensation of **7** with diketone **8** in THF at reflux afforded 2,9-bis(1,2,4-triazin-3-yl)-1,10-phenanthroline (BTPhen) **9** in 59 % yield (see Supporting Information). The ligand **9** was obtained as a stable hydrate as indicated by its ¹H NMR spectrum, the water remaining bound to the ligand *in vacuo* (0.1 mm Hg) and after heating at 120° for 24 hours.



Scheme 1. Synthesis of the BTPhen ligand 9.

The known diketone 8^{20} was synthesized by a modified procedure (Scheme 2). The diester intermediate 11 proved very difficult to synthesize on a large scale using the literature method (oxidative coupling of pivalic acid using H₂O₂/FeSO₄, followed by esterification of the resulting diacid)²¹ but we have since developed a new synthesis of 11 involving the alkylation of ethyl isobutyrate 10 with ethane-1,2-disulfonate esters.²² This procedure gave the diester 11 in 69–70 % yield. This contrasts to when 1,2-dihaloethanes were used in attempted alkylation reactions, when none of the desired diester 11 was formed and the major product was the ethyl α -haloisobutyrate.²³ The diester 11 was converted to the diketone 8 as previously described (see Supporting Information).^{20a} Using this modified procedure, diketone 8 was readily synthesized on a multi-gram scale in much improved overall yield (44 % overall compared to <5 % using the previous method).



Scheme 2. Modified and improved synthesis of the diketone 8.

The X-ray crystal structure of the ligand **9** (Figure 2)²⁴ shows one of the bound water molecules located in the coordination cavity of the ligand. The two outer triazine rings of BTPhen **9** are twisted around away from the conformation required for metal ligation such that the N(11)-C-C-N(21) and N(32)-C-C-N(41) torsion angles are -159.0(2), -159.0(2) respectively. The central N(21)-C-C-N(32) torsion angle is $-8.6(3)^{\circ}$. The Eu(III) complex of BTPhen **9** was synthesized (see Supporting Information) and structurally characterized by X-ray diffraction.²⁵ We were surprised to find that the biscomplex of stoichiometry [Eu(**9**)₂NO₃]²⁺ had been formed (Figure 3), even though a single equivalent of **9** was used in the crystal growing experiment.



Figure 2. Molecular structure in the crystal of the BTPhen ligand **9**. Thermal ellipsoids are at 30 % probability. Hydrogen bonds shown as dotted lines. Distances are O(1)-N(21) 3.003(2), O(1)-N(45) 3.125(2)Å. Other solvent molecules are omitted for clarity.

This is the first time that a 1:2 lanthanide complex with any quadridentate bis-triazine ligands has been isolated and structurally characterized. Although it is known that the BTBP ligands can form both 1:1 and 1:2 complexes in solution, only 1:1 complexes have been previously characterized by X-ray diffraction.²⁶ The metal is ten-co-ordinate being fully enclosed by two molecules of **9** in addition to a single bidentate nitrate ion. The geometry can best be considered as a capped square antiprism with the bidentate nitrate group occupying the capping site. The bond lengths from the metal to the outer triazine nitrogen atoms are equivalent with a range of 2.546(6)-2.616(5)Å to those to the inner phenanthroline nitrogen atoms with a range of 2.561(6)-2.598(5)Å. Indeed the mean bond lengths are 2.587, 2.582Å respectively. These metal-nitrogen bond lengths are very similar to those found in the 1:1 complexes formed by the BTBP ligands.²⁶ By contrast to the ligand conformation, now the outer torsion angles N(11)-C-C-N(21) and N(32)-C-C-N(41) are 6.0(10), $-6.0(10)^{\circ}$ with the central torsion angle $-0.6(8)^{\circ}$. The corresponding angles in the second ligand are 4.6(11), 10.6(10), $-4.9(11)^{\circ}$ respectively. The four donor nitrogen atoms in each ligand are approximately planar (r.m.s. deviations 0.01, 0.07Å) with the metal 0.729(3) and 0.585(4)Å respectively from each N4 plane. These two N4 planes intersect at 69.8(2)^o The bond lengths to the two nitrate oxygen atoms are 2.540(6) and 2.593(5)Å.



Figure 3. Molecular structure in the crystal of the $[Eu(9)_2NO_3]^{2+}$ cation. Thermal ellipsoids are at 30 % probability. The $[Eu(NO_3)_5]^{2-}$ counterions and solvent molecules have been omitted for clarity. Selected bond lengths [Å]: Eu(1)-N(51) 2.546(6), Eu(1)-N(72) 2.561(6), Eu(1)-N(32) 2.571(7), Eu(1)-N(21) 2.598(5), Eu(1)-N(81) 2.572(6), Eu(1)-N(61) 2.598(5), Eu(1)-N(11) 2.615(7), Eu(1)-N(41) 2.616(5), Eu(1)-O(1) 2.540(6), Eu(1)-O(3) 2.593(5).

pKa Determination and NMR Studies of Lanthanide Complexes. It is rather paradoxical that metal ions are so well extracted from concentrated acid aqueous solutions by ligands such as **1** and **9** despite their easily protonatable central bipyridine or phenanthroline units. The basicity of these ligands must be strongly decreased by the electron withdrawing triazine substituents but little or no information on the pKa of these ligands is yet available to substantiate this hypothesis. The pKa of the ligand **9** was thus determined by ¹H NMR spectroscopy.²⁷ The dependence of the NMR shifts of BTPhen **9** upon the DCl concentration in deuterated methanol $(10^{-7}-3 \text{ M}, > 99 \%$ methanol) were recorded (see Supporting Information). The NMR titration curves display a well-defined shift jump between $-\log[DCl] = 4$ and 2 followed by a smooth shift increase at higher acid concentrations. The first protonation pKa of **9** (3.1 ± 0.1) is assigned to the protonation of the phenanthroline ring; it is close to the value reported for the

BTP ligands in 76 % methanol (pKa = 3.3)^{7c} and is much lower than the pKa of phenanthroline itself in water (4.92). The electron withdrawing effect of the triazine rings of **9** is also responsible for the low value of the second pKa (0.3 ± 0.1 , 0.2 ± 0.1 and 0.03 ± 0.03 for phenanthroline protons 3, 4 and 5 respectively), which is tentatively assigned to the second protonation of phenanthroline.

Using lanthanides as actinide surrogates, the complexation behavior of BTPhen **9** towards the lanthanides was studied by NMR spectroscopy to gain an understanding of the species likely to be involved in the extraction of the trivalent actinides.

As shown earlier,^{11a} the maximum stoichiometry of lanthanide complexes in an organic solvent can be deduced from a nuclear magnetic relaxation dispersion (NMRD) titration in which the relaxation rate of a Gd^{3+} solution is measured for different metal:ligand ratios. The progressive formation of a metal complex is accompanied by the removal of solvent molecules. Consequently, the ¹H relaxation time of these molecules increases as the latter are no longer in the immediate vicinity of the paramagnetic centers. The relaxation rate thus decreases and a plateau is reached when the metal complex is fully formed. NMRD titrations are presented in Figure 4 for Gd^{3+} perchlorate and nitrate salts to which BTPhen 9 is progressively added in anhydrous acetonitrile. The clear breaks in the titration curves for a 1:2 ratio indicate that stable bis-complexes are formed in both cases. However, the curvature in the NMRD titration plot of the nitrate salt indicates that the nitrate ion is better able to compete with the BTPhen ligand 9 than the perchlorate ion. In keeping with these data, NMR peaks due to the free ligand are observed as soon as 1:2 metal:ligand ratios are exceeded. The number of resonances and their shifts in all the NMR spectra discussed below remain identical whether this ratio is 1:1 or 1:2 and the biscomplex is thus the major solution component in these conditions.



Figure 4. Nuclear magnetic relaxation dispersion (NMRD) titration curve of an anhydrous CD₃CN solution of $Gd(ClO_4)_3$ (\bigcirc) and $Gd(NO_3)_3$ (\boxdot) by BTPhen 9.

The NMR spectrum of the diamagnetic bis-complex formed between anhydrous Lu(ClO₄)₃ and BTPhen **9** in acetonitrile displays two methyl peaks, three aromatic peaks and a multiplet between 1.45 and 1.65 ppm that originates from the rapidly inverting cyclohexenyl groups (see Supporting Information). These peaks broaden somewhat at lower temperatures but the changes are insufficient for a quantitative analysis that would lead to an estimation of the activation energy. Attempts in chloroform-acetonitrile mixtures at 223 K were also unsuccessful and were not pursued as the rigidification of cyclohexene itself is only reached at 109 K.²⁸ A NOESY spectrum was recorded to verify the assignments (see Supporting Information). Cross-peaks were observed between the cyclohexenyl protons and between the aromatic protons but not between these two groups. No cross peaks were observed between two different protons on separate ligands. The spectrum of Lu(BTPhen)₂³⁺ was used as a reference for deducing the paramagnetic shifts induced by the Yb³⁺ ion. The latter is known to induce essentially pure dipolar shifts from which solution structures can be inferred.²⁹

The ¹H NMR spectrum of a 1:2 mixture of Yb(ClO₄)₃ and BTPhen **9** in anhydrous acetonitrile is shown in Figure 5. All resonances are easily assigned from their respective areas and from COSY spectra. Three aromatic protons resonate at high and low fields and the cyclohexenyl substituents give

rise to two methyl peaks and two methylene peaks. The aliphatic substituents are thus rapidly inverting their conformation on the NMR time scale (over a 17 ppm shift range). This remains true between 298 and 230 K.



Figure 5. ¹H NMR spectrum of a 1:2 Yb(ClO₄)₃:BTPhen **9** mixture in anhydrous CD₃CN at 263 K. Assignments: (+) methyl protons, (o) methylene protons, (**x**) aromatic protons (broad peak at -3.7 ppm: traces of water).

A molecular modelling calculation was made following the procedure used for a related bis-triazine ligand derived from 2,2':6',2''-terpyridine (force-field approach with parameters published for lanthanide complexes).^{11b} The optimized structure of the 1:2 bis-complex of **9** with Yb³⁺ is shown in Figure 6. As expected, the most stable conformation is an arrangement in which the two ligands are perpendicular to each other. All cyclohexenyl groups in the modelled structure are in the half-chair conformation in agreement with molecular calculations performed for cyclohexene.²⁸ The red arrows in Figure 6 indicate the smaller interatomic distances found between two ligands in the bis-complex. All these distances are larger than the sum of the Van der Waals radii. The structure is thus not sterically crowded and there is ample room for additional coordinating anions or solvent molecules. The full dipolar equation 1 was used to unravel the solution structure of the Yb³⁺ BTPhen **9** bis-complex.

$$\delta_{i} = \frac{1}{12\pi} \left[\left(\chi_{zz} - \frac{1}{2} \left(\chi_{xx} + \chi_{yy} \right) \right) \frac{\left\langle 3\cos^{2}\theta_{i} - 1 \right\rangle}{r_{i}^{3}} + \left(\chi_{xx} - \chi_{yy} \right) \frac{\left\langle \sin^{2}\theta_{i}\cos 2\psi_{i} \right\rangle}{r_{i}^{3}} \right]$$
(1)



Figure 6. Force field simulation of the optimized structure of the 1:2 bis-complex of BTPhen **9** with Yb^{3+} . The arrows show the shortest interatomic distances in the metal complexes. These distances are larger than the sum of the van der Walls radii. See Supporting Information for a list of structural parameters.

In this equation,²⁹ the paramagnetic shifts depend on magnetic susceptibility terms that are identical for all nuclei and on geometric terms that are different for each nucleus and are easily computed from a molecular model. The factors θ_i , ψ_i and r_i are the polar coordinates of proton i in the set of axes of the magnetic susceptibility tensor with the metal ion at its origin. This equation has been used successfully for obtaining the solution structure of lanthanide chelates²⁹ or of proteins modified by grafting a paramagnetic metal complex.³⁰ In the latter case, the spectra display so many shifted resonances that the orientation of the susceptibility axes can be included in a fit between the experimental and calculated paramagnetic shifts. This is not the case for the BTPhen complex that exhibits only seven NMR resonances but this complex features symmetry elements with which the axes of the magnetic susceptibility tensor must coincide. The z axis was thus oriented with the C₂ axis that bisects the two phenanthroline groups. In addition to this axis, there are two symmetry planes, either through the

phenanthroline rings or between these rings (ie: at 45°). The x and y axes can be located in either plane. For symmetry reasons, the same correlation between experimental and calculated paramagnetic shifts will be obtained but the sign of the magnetic susceptibility terms will change. This is also true if the x (or y) axis is exchanged with the z axis (see Supporting Information). As shown in Figure 7, there is a very good correlation between the calculated and experimental paramagnetic shifts based on the dipolar equation 1. The correlation is especially good if one takes into account that the cyclohexenyl rings are rapidly inverting. The geometric factors of each exchanging group were obtained by computing the mean of the factors for two proton sets exchanging their positions during the ring inversion process. No account was taken of the transition states nor of the fact that cyclohexene adopts various less stable geometries in addition to the half-chair conformation.²⁸ Not surprisingly, the datum point that is the most removed from the correlation line in Figure 7 originates from one of the methyl proton sets undergoing rapid inversion. The geometric factors of the exchanging protons are very different in this particular case and larger errors are thus expected.



Figure 7. Correlation between the calculated and experimental paramagnetic shifts of the bis-complexes of BTPhen **9** with Yb^{3+} perchlorate (\bigcirc) or nitrate (\boxdot). Y axis on left: perchlorate, y axis on right: nitrate.

There is one feature worthy of note in these calculations that has not been mentioned in the literature so far. As the two ligands are planar in an environment of two-fold symmetry, an agreement indistinguishable from the one presented in Figure 7 is obtained if the angle between the two ligands is increased or decreased. The geometric factors are of course modified but the magnetic susceptibility terms are also changing and the correlation between calculated and experimental paramagnetic shifts remains unaltered (For instance, if the angle between the two BTPhen ligands are 90, 75, 60 and 45°, the magnetic susceptibility factors in equation 1 become 1392 and -4038, 1804 and -3600, 1617 and -3788, 1543 and -3863 Å⁻³ respectively). We are confronted here with a limit of the dipolar shift analysis method that does not arise with non-planar ligands such as one based on 2,2':6',2''-terpyridine^{11b} or other complexes investigated so far.²⁹ A consequence of this is that a good correlation will still be obtained if the structure of the complex becomes flatter to allow for the entrance of an anion or a solvent molecule in the first coordination sphere. A good correlation between experimental and calculated shifts is thus still expected for the bis-BTPhen nitrate complex depicted in Figure 3 as it features a pseudo-twofold symmetry axis and a larger opening for accommodating an anion.

A similar analysis was carried out for the 1:2 bis-complex of BTPhen **9** with Eu shown in Figure 3. The ¹H NMR spectrum of a 1:2 mixture of **9** and Eu(ClO₄)₃ was recorded and the analysis of the induced paramagnetic shifts was performed as for the Yb(III) bis-complex (see Supporting Information). However, unlike Yb(III), the contact paramagnetic shifts for Eu(III) are larger than the dipolar paramagnetic shifts and thus the dipolar equation (1) cannot be used as reliably for solution structure determinations.^{29d,31} Only a qualitative agreement was obtained from which the solution structure of the complex could not be reliably established.

In the ¹H NMR spectrum of a 1:2 mixture of Yb(NO₃)₃ and BTPhen **9** in acetonitrile (see Supporting Information), the entrance of a nitrate ion into the first coordination sphere is clearly indicated by paramagnetic shifts that are totally different from those observed in the perchlorate case. The magnetic susceptibility terms should indeed be totally different because of the presence of a charged species directly coordinated to the metal ion. However, the molecular model remains valid as shown by the good

correlation between experimental and calculated shifts presented in Figure 7. Nevertheless, the degree of opening of the bis-complex to accommodate the nitrate ion remains unknown and it appears that NMR spectroscopy is of no help in solving this particular problem.

Solvent Extraction Studies with Am(III), Cm(III) and Eu(III) Radionuclides. Preliminary solvent extraction experiments were then carried out to determine the ability of 9 to selectively extract An(III) over Ln(III). Solutions of the ligand 9 in octanol (0.01 M) were contacted with nitric acid solutions containing ²⁴¹Am and ¹⁵²Eu radiotracers. The distribution ratios (D) for Am(III) and Eu(III), and the separation factors ($SF_{Am/Eu}$) at different nitric acid concentrations are shown in Figure 8. Very high D values for Am(III) were observed ($D_{Am} > 1000$), indicating that the extraction of Am(III) by 9 is very efficient. The D values for Eu(III) were approximately two orders of magnitude lower than those of Am(III) ($D_{Eu} < 10$), and the resulting separation factors were in the range 68–400. These results show that 9 is able to extract Am(III) in preference to Eu(III) with very high selectivity. The formation of stable 1:2 bis-complexes by 9 certainly contribute to the higher D values observed for Am(III) compared to previous N-donor ligands.^{7c} The distribution ratios for Am(III) for 9 are about two orders of magnitude higher that those for the related BTBP ligand 1.^{10a} The considerable improvement in the extraction of Am(III) by 9 compared to 1 far exceeds what was anticipated on the basis of molecular design. The high D values observed for 9 make it especially suitable even for low-level waste decontamination where only trace levels of radionuclides are present.

In common with other bis-triazine ligands, BTPhen **9** is able to extract Am(III) from nitric acid solutions of high acidity (1–4 M HNO₃). Thus, although the ligand is likely to be protonated in contact with HNO₃ solutions (pKa = 3.1-3.2), proton competition does not prevent metal ion extraction. This is in contrast with other *N*-donor ligands based on 2,2':6',2''-terpyridine which can only extract Am(III) from weakly acidic (< 0.1 M HNO₃) solutions.³² The precise reasons for this extraction ability are unclear at this point. It should be noted that, although the competition between protonation and extraction leads to a decrease in *D* values, an increase in acidity is also an increase in the nitrate ion concentration that favours the extraction. The formation of highly thermodynamically stable 1:2 bis-

complexes by BTPhen **9**, coupled with the high hydrophobicity of the ligand could also explain the observed extraction results. In addition, a very low pKa value of -1.77 has been estimated for 1,2,4-triazine from a correlation between pKa's and ionization energies³³ and the triazine rings of **9** are thus believed to remain unprotonated even at very high acidities. The extraction of lanthanide and actinide salts from concentrated HNO₃ solutions by **9** is thus less thwarted by protonation than for other *N*-donor ligands.



Figure 8. Extraction of Am(III) and Eu(III) by BTPhen 9 in octanol (0.01 M) as a function of initial nitric acid concentration (D = distribution ratio, SF = separation factor, $\blacktriangle = D_{Am}$, $\bullet = D_{Eu}$, $\blacksquare = SF_{Am/Eu}$, mixing time: 60 min., temperature: 22°C ± 1°C).

Figure 9 shows the kinetics of extraction of Am(III) and Eu(III) by BTPhen 9 from 1 M HNO₃ into octanol. For Am(III), relatively fast extraction kinetics are observed even in the absence of a phase-transfer agent, and distribution ratios close to the equilibrium value are reached within 15 minutes of phase contact time. Thus the kinetics of extraction by the BTPhen ligand 9 are significantly faster than the BTBP ligand 1 which requires up to 60 minutes of contact time to reach its equilibrium D_{Am} value of approx. 4.5.^{10a} The faster rates of extraction by BTPhen 9 mean that the use of a phase-transfer agent such as N,N'-dimethyl-N,N'-dioctyl-2-(2-hexoxyethyl) malondiamide (DMDOHEMA),³⁴ which is needed to improve the extraction kinetics of BTBP 1,¹⁰ is not necessary. Furthermore, as D_{Am} is greater

than 10 even after 5 minutes of contact, an efficient extraction can still take place using shorter contact times if desired.



Figure 9. Extraction of Am(III) and Eu(III) from 1 M HNO₃ by BTPhen **9** in octanol (0.01 M) as a function of time ($\blacksquare = D_{Am}$, $\bullet = D_{Eu}$, temperature: 22°C ± 1°C).

In a separation process, it is desirable to back-extract (strip) the actinides from the loaded organic phase after the extraction has taken place so that the ligand can be re-used in further extraction cycles. Ideally this requires D_{Am} values < 1 for efficient stripping. Figure 8 shows that for **9**, the distribution ratios for Am(III) are lowest at low acidity but still exceed 1. The effect of the less polar diluents toluene and hydrogenated tetrapropene (TPH, a dodecane-like diluent) as co-diluents was then studied in order to reduce the *D* values for Am(III) at low acidities (see Supporting Information). With toluene as the codiluent, the distribution ratios for Am(III) and Eu(III) both decreased as the volume fraction of toluene increased. This effect was most pronounced at low acidities. With 0.001 M HNO₃ as the aqueous phase, D_{Am} decreased from 1.57 in octanol/toluene (80:20) to 0.11 in octanol/toluene (20:80). When toluene was the sole diluent, a D_{Am} value of 0.02 was observed (aqueous phase: 0.01 M HNO₃). With TPH as the co-diluent, D_{Am} values of 0.48 and 0.45 were observed in the extraction from 0.001 M HNO₃ into octanol/TPH mixtures (60:40 and 50:50), respectively.

Extraction experiments were then carried out using ²⁴¹Am and ²⁴⁴Cm radiotracers in order to probe the co-extraction of both Am(III) and Cm(III) by BTPhen **9** (see Supporting Information). Both Am(III) and

Cm(III) were efficiently co-extracted into octanol/toluene (40:60) solutions by **9** with virtually no selectivity between the two actinides ($SF_{Am/Cm} = 0.7-4$). In the extraction of both Am(III) and Cm(III) from 0.001 M HNO₃ into octanol/toluene mixtures, the *D* values for both metals decreased as the volume fraction of toluene increased, and *D* values below 1 were observed for both metals in 40:60, 20:80 and 0:100 octanol/toluene mixtures. These results indicate that the stripping of both Am(III) and Cm(III) from a loaded organic phase is feasible at lower acidities.

The waste streams produced in the PUREX process contain high concentrations of Y(III) and all the trivalent lanthanides in addition to the minor actinides. Information is therefore needed on the extraction of the lanthanides as well as the actinides. The extraction of the lanthanides by octanol solutions of BTPhen **9** at several different acidities showed a profile across the lanthanide series of first increasing, then decreasing *D* values, with the maximum *D* value being observed for Sm(III) (see Supporting Information). A similar trend was reported for the BTBP ligand **1**.^{10a} Thus BTPhen **9** efficiently separates both Am(III) and Cm(III) from the entire lanthanide series.

Kinetics and Surface Tension Measurements. In order to uncover the reasons for the faster extraction kinetics observed with **9** compared to its non-rigidified analogue **1**, a comparison of the extraction kinetics of ¹⁵²Eu by BTBP **1** and BTPhen **9** in different diluents was carried out using the rotating membrane cell technique.³⁵ With octanol, 2-methylcyclohexanone, 3-methylcyclohexanone and 4-methylcyclohexanone as the diluents, the calculated extraction rate constants (k_{ext}) for BTPhen **9** are substantially larger than those for BTBP **1** in each of these diluents (Figure 10). Conversely, the back-extraction (stripping) rate constants (k_{str}) for BTPhen **9** are larger than for BTPhen **9** in these diluents (Figure 11). These results suggest that BTPhen **9** will have a faster rate of extraction, but a slower rate of stripping compared to BTBP **1**. The fastest extraction kinetics for both ligands were observed in cyclohexanone ($k_{ext} = 121 \times 10^{-6}$ cm s⁻¹ for 0.01 M BTBP **1** from 2 M HNO₃; $k_{ext} = 74.8 \times 10^{-6}$ cm s⁻¹ for 0.01 M BTPhen **9** from 1 M HNO₃), which is known to improve significantly the kinetics of extraction by the BTBPs.³⁶ Using *N*,*N*,*N*,*N*-tetraoctyl diglycolamide (TODGA)^{34,37} as additive also

accelerated the extraction kinetics of BTBP **1** in octanol, but was less effective in accelerating the extraction kinetics of BTPhen **9** in octanol (see Supporting Information).



Figure 10. Extraction rate constants (k_{ext}) in different diluents for 0.01 M solutions of BTBP 1 and BTPhen 9 (aqueous phase: 2 M HNO₃).

The interfacial tensions between aqueous phases of 1 M HNO₃ and organic phases of BTBP 1 and **BTPhen** 9 diluted octanol, 2-methylcyclohexanone, 3-methylcyclohexanone in and 4methylcyclohexanone were then measured using the du Noüy ring method.³⁸ In the case of BTBP 1, surface activity was only observed when the ligand was diluted in 3-methylcyclohexanone, and was not observed in the other diluents. In the case of BTPhen 9, surface activity was observed in all four diluents (see Supporting Information). The sharpest decrease in surface tension was observed in octanol. The faster extraction kinetics observed with BTPhen 9 compared to BTBP 1 thus appears to be related to the greater surface concentration of the BTPhen ligand 9 at the interface. Hence BTPhen 9, which exhibits surface activity in all diluents, gives extraction kinetics that are faster than BTBP 1, which is surface active in only one diluent (3-methylcyclohexanone). The greater surface activity observed for BTPhen 9 compared to 1 is in keeping with what was predicted based on ligand design. The higher polarity of 9 and its ability to strongly coordinate to water is probably responsible for its ability to interact favourably with the interface.



Figure 11. Back-extraction rate constants (k_{str}) in different diluents for 0.01 M solutions of BTBP 1 and BTPhen 9 (aqueous phase: 2 M HNO₃).

Conclusions

In summary, we have reported the first example of a promising new class of highly-selective solvent extraction reagent for the partitioning of trivalent actinides from trivalent lanthanides in nuclear waste streams, and have demonstrated that pre-organization of the donor atoms for metal ligation with a rigid *cis*-locked 1,10-phenanthroline motif leads to a rapid and highly efficient separation of actinides from lanthanides. Interfacial tension measurements suggest that the improved extraction kinetics of the ligand relative to its 2,2'-bipyridine counterpart are related to higher concentrations of the ligand at the interface. The first X-ray crystal structure of a 1:2 bis-complex of a quadridentate bis-triazine ligand with Eu(III) has been determined. Lanthanide NMR complexation studies allowed the solution structure of the 1:2 bis-complex with Yb(III) to be deduced from the induced paramagnetic shifts. The aliphatic diketone precursor to the ligand has been synthesized by an improved procedure which, for the first time, allows the straightforward synthesis of BTPhen **9** and related ligands on at least a multi-gram scale. Further applications of the ligand in coordination chemistry are anticipated.

Acknowledgment. We thank the Nuclear Fission Safety Program of the European Union for support under the ACSEPT (FP7-CP-2007-211 267) contract. We also thank the EPSRC and the University of Reading for funds for the X-Calibur system. Use of the Chemical Analysis Facility at the University of Reading is gratefully acknowledged.

Supporting Information Available: Procedures and characterization data for all compounds. Tables and graphs of solvent extraction data. Figures of optimized structures and NMR spectra of lanthanide complexes. Procedure for pKa determination. X-ray crystallographic cif files for ligand 9 and its Eu complex. Tables and graphs of kinetic data and interfacial tension measurements. This material is available free of charge via the Internet at http://pubs.acs.org.

References

- OECD/NEA, Nuclear Fuel Cycle Transition Scenario Studies Status Report, NEA No. 6194, OECD Publications, Paris, 2009.
- (a) Baumgärtner, F. Kerntechnik 1978, 20, 74. (b) Baumgärtner, F.; Ertel, D. J. Rad. Chem.
 1980, 58, 11. (c) McKibben, J. M. Radiochim. Acta 1984, 36, 3. (d) Sood, D. D.; Patil, S. K. J. Rad. Nucl. Chem. 1996, 203, 547. (e) Madic, C. Actualite Chimique 1999, 16. (f) Musikas, C.; Schultz, W.; Liljenzin, J.-O. Solvent Extraction Principles and Practice, 2nd ed; Rydberg, J.; Cox, M.; Musikas, C.; Choppin, G., Eds.; Marcel Dekker, Inc.: New York, 2004, pp 507–557.
- Magill, J.; Berthou, V.; Haas, D.; Galy, J.; Schenkel, R.; Wiese, H.-W.; Heusener, G.; Tommasi, J.; Youinou, G. *Nuclear Energy* 2003, *42*, 263.
- 4. Mathur, J. N.; Murali, M. S.; Nash, K. L. Solvent Extr. Ion Exch. 2001, 19, 357.
- (a) Cotton, S. Comprehensive Coordination Chemistry II, Vol. 3; McCleverty, J. A.; Meyer, T. J., Eds.; Elsevier: Oxford, 2004, pp 93–188. (b) Burns, C. J.; Neu, M. P.; Boukhalfa, H.; Gutowski, K. E.; Bridges, N. J.; Rogers, R. D. Comprehensive Coordination Chemistry II, Vol. 3; McCleverty, J. A.; Meyer, T. J., Eds.; Elsevier: Oxford, 2004, pp 189–332. (c) Katz, J. J.; Morss, L. R.; Edelstein, N. M.; Fuger, J. The Chemistry of the Actinide and Transactinide

Elements, Vol. 1; Katz, J. J.; Morss, L. R.; Edelstein, N. M.; Fuger, J., Eds.; Springer: Dordrecht, 2006, pp 1–17.

- 6. (a) Choppin, G. R. J. Alloys Compd. 1995, 223, 174. (b) Alexander, V. Chem. Rev. 1995, 95, 273. (c) Choppin, G. R. J. Alloys Compd. 2002, 344, 55. (d) Gaunt, A. J.; Neu, M. P. C. R. Chimie 2010, 13, 821.
- (a) Dam, H. H.; Reinhoudt, D. N.; Verboom, W. *Chem. Soc. Rev.* 2007, *36*, 367. (b) Ekberg, C.;
 Fermvik, A.; Retegan, T.; Skarnemark, G.; Foreman, M. R. S.; Hudson, M. J.; Englund, S.;
 Nilsson, M. *Radiochim. Acta* 2008, *96*, 225. (c) Kolarik, *Z. Chem. Rev.* 2008, *108*, 4208.
- 8. For leading references, see: (a) Drew, M. G. B.; Guillaneux, D.; Hudson, M. J.; Iveson, P. B.; Russell, M. L.; Madic, C. Inorg. Chem. Commun. 2001, 4, 12. (b) Iveson, P. B.; Riviére, C.; Guillaneux, D.; Nierlich, M.; Thuéry, P.; Ephritikhine, M.; Madic, C. Chem. Commun. 2001, 1512. (c) Boucher, C.; Drew, M. G. B.; Giddings, P.; Harwood, L. M.; Hudson, M. J.; Iveson, P. B.; Madic, C. Inorg. Chem. Commun. 2002, 5, 596. (d) Ionova, G.; Rabbe, C.; Guillaumont, R.; Ionov, S.; Madic, C.; Krupa, J.-C.; Guillaneux, D. New J. Chem. 2002, 26, 234. (e) Berthet, J.-C.; Miquel, Y.; Iveson, P.B.; Nierlich, M.; Thuéry, P.; Madic, C.; Ephritikhine, M. J. Chem. Soc. Dalton Trans. 2002, 3265. (f) Colette, S.; Amekraz, B.; Madic, C.; Berthon, L.; Cote, G.; Moulin, C. Inorg. Chem. 2002, 41, 7031. (g) Colette, S.; Amekraz, B.; Madic, C.; Berthon, L.; Cote, G.; Moulin, C. Inorg. Chem. 2003, 42, 2215. (h) Colette, S.; Amekraz, B.; Madic, C.; Berthon, L.; Cote, G.; Moulin, C. Inorg. Chem. 2004, 43, 6745. (i) Denecke, M. A.; Rossberg, A.; Panak, P. J.; Weigl, M.; Schimmelpfennig, B.; Geist, A. Inorg. Chem. 2005, 44, 8418. (j) Drew, M. G. B.; Foreman, M. R. St. J.; Geist, A.; Hudson, M. J.; Marken, F.; Norman, V.; Weigl, M. Polyhedron 2006, 25, 888. (k) Steppert, M.; Walther, C.; Geist, A.; Fanghänel, T. New J. Chem. 2009, 33, 2437. (1) Banik, N. L.; Schimmelpfennig, B.; Marquardt, C. M.; Brendebach, B.; Geist, A.; Denecke, M. A. Dalton Trans. 2010, 39, 5117. (m) Benay, G.; Schurhammer, R.; Wipff, G. Phys. Chem. Chem. Phys. 2010, 12, 11089.

- For leading references, see: (a) Drew, M. G. B.; Foreman, M. R. S. J.; Hill, C.; Hudson, M. J.; Madic, C. *Inorg. Chem. Commun.* 2005, *8*, 239. (b) Nilsson, M.; Ekberg, C.; Foreman, M.; Hudson, M.; Liljenzin, J.-O.; Modolo, G.; Skarnemark, G. *Solvent Extr. Ion Exch.* 2006, *24*, 823.
 (c) Trumm, S.; Lieser, G.; Foreman, M. R. S. J.; Panak, P. J.; Geist, A.; Fanghänel, T. *Dalton Trans.* 2010, *39*, 923. (d) Hubscher-Bruder, V.; Haddaoui, J.; Bouhroum, S.; Arnaud-Neu, F. *Inorg. Chem.* 2010, *49*, 1363. (e) Ekberg, C.; Aneheim, E.; Fermvik, A.; Foreman, M.; Löfström-Engdahl, E.; Retegan, T.; Spendlikova, I. *J. Chem. Eng. Data* 2010, *55*, 5133. (f) Berthet, J.-C.; Maynadié, J.; Thuéry, P.; Ephritikhine, M. *Dalton Trans.* 2010, *39*, 6801. (g) Benay, G.; Schurhammer, R.; Wipff, G. *Phys. Chem. Chem. Phys.* 2011, *13*, 2922.
- (a) Geist, A.; Hill, C.; Modolo, G.; Foreman, M. R. St. J.; Weigl, M.; Gompper, K.; Hudson, M. J. Solvent Extr. Ion Exch. 2006, 24, 463. (b) Magnusson, D.; Christiansen, B.; Foreman, M. R. S.; A. Geist, A.; Glatz, J.-P.; Malmbeck, R.; Modolo, G.; Serrano-Purroy, D.; Sorel, C. Solvent Extr. Ion Exch. 2009, 27, 97.
- (a) For a related ligand derived from pyridine, see: Hudson, M. J.; Boucher, C. E.; Braekers, D.; Desreux, J. F.; Drew, M. G. B.; Foreman, M. R. St. J.; Harwood, L. M.; Hill, C.; Madic, C.; Marken, F.; Youngs, T. G. A. *New. J. Chem.* 2006, *30*, 1171. (b) For a related ligand derived from 2,2':6',2''-terpyridine, see: Lewis, F. W.; Harwood, L. M.; Hudson, M. J.; Drew, M. G. B.; Modolo, G.; Sypula, M.; Desreux, J. F.; Bouslimani, N.; Vidick, G. *Dalton Trans.* 2010, *39*, 5172.
- For an example of a radiolytically unstable BTBP, see: Fermvik, A.; Berthon, L.; Ekberg, C.;
 Englund, S.; Retegan, T.; Zorz, N. *Dalton Trans.* 2009, 6421.
- 13. (a) Berthet, J.-C.; Thuéry, P.; Foreman, M. R. S.; Ephritikhine, M. *Radiochim. Acta* 2008, 96, 189. (b) Berthet, J.-C.; Thuéry, P.; Dognon, J.-P.; Guillaneux, D.; Ephritikhine, M. *Inorg. Chem.*

2008, *47*, 6850. (c) Berthet, J.-C.; Siffredi, G.; Thuéry, P.; Ephritikhine, M. Dalton Trans. **2009**, 3478.

- (a) Sammes, P. G.; Yahioglu, G. Chem. Soc. Rev. 1994, 23, 327. (b) Luman, C. R.; Castellano, F. N. Comprehensive Coordination Chemistry II, Vol. 1; McCleverty, J. A.; Meyer, T. J., Eds.; Elsevier: Oxford, 2004, pp 25–39. (c) Accorsi, G.; Listorti, A.; Yoosaf, K.; Armaroli, N. Chem. Soc. Rev. 2009, 38, 1690. (d) Bencini, A.; Lippolis, V. Coord. Chem. Rev. 2010, 254, 2096.
- 15. Zong, R.; Thummel, R. P. J. Am. Chem. Soc. 2004, 126, 10800.
- Cockrell, G. M.; Zhang, G.; VanDerveer, D. G.; Thummel, R. P.; Hancock, R. D. J. Am. Chem. Soc. 2008, 130, 1420.
- 17. See for example: Cumper, C. W. N.; Ginman, R. F. A.; Vogel, A. I. J. Chem. Soc. 1962, 1188.
- 18. See for example: Donnay, G.; Donnay, J. D. H.; Harding, M. J. C. Acta. Cryst. 1965, 19, 688.
- 19. Chandler, C. J.; Deady, L. W.; Reiss, J. A. J. Het. Chem. 1981, 18, 599.
- 20. (a) Jones, P.; Villeneuve, G. B.; Fei, C.; DeMarte, J.; Haggarty, A. J.; Nwe, K. T.; Martin, D. A.; Lebuis, A-M.; Finkelstein, J. M.; Gour-Salin, B. J.; Chan, T. H.; Leyland-Jones, B. R. J. *Med. Chem.* 1998, 41, 3062. (b) Kikuchi, K.; Hibi, S.; Yoshimura, H.; Tokuhara, N.; Tai, K.; Hida, T.; Yamauchi, T.; Nagai, M. J. Med. Chem. 2000, 43, 409.
- 21. (a) Coffman, D. D.; Jenner, E. L.; Lipscomb, R. D. J. Am. Chem. Soc. 1958, 80, 2864. (b)
 Bremner, D. H.; Mitchell, S. R.; Staines, H. Ultrasonics Sonochemistry 1996, 3, 47. (c) Bremner,
 D. H.; Burgess, A. E.; Li, F-B. Green Chem. 2001, 3, 126.
- 22. Lewis, F. W; Harwood, L. M.; Hudson, M. J. WO2011077081 A1, (2011).
- This type of reactivity has been reported with 1,2-dibromoethane, see: Ibarzo, J.; Ortuña, R. M. *Tetrahedron* 1994, *50*, 9825.

- 24. Crystal data of ligand **9**: **9**, 0.5MeOH, 1.5H₂O, C_{34.5}H₄₃N₈O₂, M = 601.77, monoclinic, spacegroup $P2_1/n$, Z = 4, a = 15.5530(8), b = 10.2067(6), c = 20.833(2)Å, $\beta = 103.417(7)^{\circ}$, U = 3216.9(4)Å³, $d_{calc} = 1.243$ g cm⁻³, 8714 independent reflections, 4145 observed reflections, $R_1 = 0.0568$, $wR_2 = 0.1203$, CCDC 789753.
- 25. Crystal data of Eu(III) bis-complex of ligand **9**: [Eu(**9**)₂(NO₃)] [Eu(NO₃)₅], NCMe, H₂O, C₇₀H₈₁Eu₂N₂₃O₁₉, M = 1852.50, triclinic, spacegroup *P*-1, Z = 2, a = 13.5768(8), b = 15.3420(10), c = 20.5094(13)Å, a = 106.114(6), $\beta = 100.566(5)$, $\gamma = 98.556(5)^{\circ}$, U = 3943.4(4)Å³, $d_{calc} = 1.560$ g cm⁻³, 21133 independent reflections, 7523 observed reflections, *R*₁ = 0.0595, $wR_2 = 0.1088$, CCDC 789754.
- Foreman, M. R. S.; Hudson, M. J.; Drew, M. G. B.; Hill, C.; Madic, C. Dalton Trans. 2006, 1645.
- 27. (a) Ackerman, J. J. H.; Soto, G. E.; Spees, W. M.; Zhu, Z.; Evelhoch, J. L. *Mag. Res. Med.* 1996, 36, 674. (b) Grycova, L.; Dommisse, R.; Pieters, L.; Marek, R. *Mag. Res. Chem.* 2009, 47, 977.
- 28. Anet, F. A. L.; Haq, M. Z. J. Am. Chem. Soc. 1965, 87, 3147.
- (a) Desreux, J. F. *Inorg. Chem.* 1980, 19, 1319. (b) Spirlet, M. R.; Rebizant, J.; Desreux, J. F.; Loncin, M. F. *Inorg. Chem.* 1984, 23, 359. (c) Desreux, J. F.; Loncin, M. F. *Inorg. Chem.* 1986, 25, 69. (d) Bertini, I.; Luchinat, C.; Parigi, G. *Solution NMR of Paramagnetic Molecules*, Elsevier: Amsterdam, 2001, pp 62–67. (e) Ranganathan, R. S.; Raju, N.; Fan, H.; Zhang, X.; Tweedle, M. F.; Desreux, J. F.; Jacques, V. *Inorg. Chem.* 2002, 41, 6856.
- 30. (a) Keizers, P. H. J.; Desreux, J. F.; Overhand, M.; Ubbink, M. J. Am. Chem. Soc. 2007, 129, 9292. (b) Clore, G. M.; Iwahara, J. Chem. Rev. 2009, 109, 4108.
- 31. Inagaki, F.; Miyazawa, T. Prog. NMR Spectr. 1981, 14, 7.

- 32. (a) Cordier, P. Y.; Hill, C.; Baron, P.; Madic, C.; Hudson, M. J.; Liljenzin, J.-O. *J. Alloys Compd.* 1998, 271–273, 738. (b) Hagström, I.; Spjuth, L.; Enarsson, Å.; Liljenzin, J.-O.; Skålberg, M.; Hudson, M. J.; Iveson, P. B.; Madic, C.; Cordier, P. Y.; Hill, C.; Francois, N. *Solvent Extr. Ion Exch.* 1999, 17, 221. (c) Drew, M. G. B.; Iveson, P. B.; Hudson, M. J.; Liljenzin, J.-O.; Spjuth, L.; Cordier, P. Y.; Enarsson, Å.; Hill, C.; Madic, C. *J. Chem. Soc. Dalton Trans.* 2000, 821.
- 33. Brinck, T.; Murray, J. S.; Politzer, P. J. Org. Chem. 1991, 56, 2934.
- Serrano-Purroy, D.; Baron, P.; Christiansen, B.; Malmbeck, R.; Sorel, C.; Glatz, J.-P. Radiochim. Acta 2005, 93, 351.
- 35. (a) Simonin, J.-P.; Weill, J. Solvent Extr. Ion Exch. 1998, 16, 1493. (b) Simonin, J.-P.;
 Hendrawan, H. J. Phys. Chem. B 2000, 104, 7163. (c) Simonin, J.-P.; Hendrawan, H.; Dardoize,
 F.; Clodic, G. Hydrometallurgy 2003, 69, 23.
- 36. (a) Nilsson, M.; Andersson, S.; Drouet, F.; Ekberg, C.; Foreman, M.; Hudson, M.; Liljenzin, J.-O.; Magnusson, D.; Skarnemark, G. Solvent Extr. Ion Exch. 2006, 24, 299. (b) Nilsson, M.; Ekberg, C.; Foreman, M.; Hudson, M.; Liljenzin, J.-O.; Modolo, G.; Skarnemark, G. Solvent Extr. Ion Exch. 2006, 24, 823.
- Magnusson, D.; Christiansen, B.; Glatz, J.-P.; Malmbeck, R.; Modolo, G.; Serrano-Purroy, D.;
 Sorel, C. *Solvent Extr. Ion Exch.* 2009, *27*, 26.
- (a) Lecomte du Noüy, P. J. Gen. Physiol. 1925, 7, 625. (b) Yildirim Erbil, H. Surface Chemistry of Solid and Liquid Interfaces, Blackwell Publishing Ltd: Oxford, 2006.

Table of Contents Graphic

