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Cite this: DOI: 10.1039/c0xx00000x

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ARTICLE TYPE

Remarkable enhancement in Am³⁺/Eu³⁺selectivity by an ionic liquid based solvent containing bis-1,2,4-triazinyl pyridine derivatives: DFT validation of experimental results

Arunasis Bhattacharyya*1, Seraj A. Ansari¹, Trilochan Gadly², Sunil K. Ghosh², Manoj Mohapatra¹ and P.K. Mohapatra*1

5 Received (in XXX, XXX) Xth XXXXXXXXX 20XX, Accepted Xth XXXXXXXXX 20XX DOI: 10.1039/b000000x

Mutual separation of trivalent actinide (An3+) and lanthanide (Ln³⁺) using several soft (N) donor ligands (bis(5,6-dialkyl-1,2,4-triazinyl)pyridine (R-BTP)) is attempted for the first 10 time in room temperature ionic liquid (RTIL) medium. The results indicate a spectacular enhancement in the selectivity as compared to that in molecular diluents with a separation factor (S.F.) of >3000 for Am³⁺ over Eu³⁺ using the methyl derivative (Me-BTP) in RTIL medium using [C_nmim].[NTf₂] 15 as the diluents (where n = 2, 3, 4, 6 or 8). Such a high S.F. value has never been reported before with any of the R-BTP derivatives in molecular diluents. An opposite trend in the distribution ratio values of both Am3+ and Eu3+ with increasing the size of alkyl (R) group is observed in RTIL 20 medium when compared with that in molecular diluents. The differences in the extraction behaviour of R-BTPs in RTILs vis-à-vis molecular diluents are explained on the basis of difference in the nature of complexes extracted in these two distinctly different media as supported by the time resolved 25 fluorescence (TRFS) study. Unusually high extractability and selectivity for Am³⁺ over Eu³⁺ with Me-BTP was attributed to the formation of 1:4 complex for Am³⁺, which was never reported earlier with any of the R-BTP derivatives in molecular diluents. DFT studies indicated higher metal 'd' 30 and 'f' orbital participation (covalence) in the bonding with R-BTP in case of Am3+ complexes as compared to that in case of Eu³⁺ complexes, which resulted in the selectivity of these classes of ligands. The observed results may have great significance in the radioactive waste management involving 35 the partitioning and transmutation strategy.

Introduction

One of the most challenging problems in the back end of the nuclear fuel cycle is the separation of trivalent actinides (An³⁺) from the lanthanides (Ln3+) as these two classes of 'f' block 40 elements, possess very similar complexation behaviour due to their comparable charge to radius ratios [1,2]. Soft (S,N) donor ligands show preference for An³⁺ ions over Ln³⁺ ions due to their ability to form stronger covalent bonds with An3+ ions. This has been attributed to the higher spatial distribution of the '5f' 45 orbitals of the actinides as compared to that of the '4f' orbitals of the lanthanides. A large number of literature is available on the Ln³⁺/An³⁺ separations using various 'S' and 'N' donor ligands [3-9]. Alkyl substituted bis-triazinyl pyridine (R-BTP), bis-triazinyl bipyridine (R-BTBP) and bis-triazinyl phenanthroline (R-50 BTPhen) (Figure 1) ligands are found to be quite promising in this regard showing high selectivity for An³⁺ over Ln³⁺ ions [6]. The major drawbacks of these 'N' donor heteropolycyclic ligands include their poor solubility in suitable organic long-chain

with n-dodecane [3,5,6]. Room temperature ionic liquid (RTIL) based solvent systems are being extensively investigated for the recovery of actinides and fission products from feeds relevant in the nuclear fuel cycle due 60 to many of their advantages over the molecular oraganic solvents which include (i) low volatility, (ii) non-flammability and (iii) high thermal stability, resulting in higher safety in the processes employing the RTILs, etc. [10-12]. RTILs display higher stability under α and γ radiation [13]. Moreover, it has been, recently, 65 reported that the R-BTP based extraction system is more radiation resistant in RTIL medium as compared to that in molecular solvents [14]. It is, therefore, important to explore the use of RTILs for process applications using radioactive elements with better recycling possibilities. Cocalia et al. [15], have 70 observed similar extraction and complexation behaviour of various actinide and lanthanide ions with dialkyl phosphoric and phosphinic acids in RTIL and molecular diluent. On the other hand, a number of literature reports are available where a differential complexation and extraction behaviour is noticed in 75 RTIL and molecular solvents [16]. Significant enhancement in the extraction of trivalent lanthanide ions using N,N,N',N'-tetra*n*-octyl diglycolamide (TODGA) was reported in [C₂mim][NTf₂] as compared to that in molecular solvent, viz. iso-octane medium and mutual selectivity among the lanthanide ions was also found 80 to be altered in the RTIL medium [17]. Different extraction behaviour in molecular solvent and RTIL media was explained

hydrocarbon solvents, viz. n-dodecane, kerosene etc. In all the

55 cases, a long chain alcohol, viz. n-octanol, was required along

on the basis of different extraction mechanism in these two kinds of solvents. In the RTIL media, the lanthanide ions are extracted

following the cation exchange mechanism, where the extraction

amount of the cationic part of the RTIL (C₂mim⁺) to the aqueous

phase. Sun et al. [18], reported an enhancement in the selective

extraction among the Ln³⁺ ions using a TALSPEAK type solvent

85 of metal ion resulted in the simultaneous transfer of an equivalent

have also studied the azide complexation of An³⁺ and Ln³⁺ ions in RTIL medium and observed slower complexation in case of the former [20]. From the observed differential behavior between An^{3+} and Ln^{3+} towards the N_3 complexation in RTIL, they 100 mentioned the possibility of using RTIL media for Ln³⁺/An³⁻

separation using the selective extractants for the An³⁺. However, to the best of our knowledge, only one literature report is available where the mutual separation of trivalent actinides and the lanthanides has been carried out in RTIL medium [21] where, 5 a task specific ionic liquid (TSIL) containing DTPA moiety was used to enhance the selectivity in a TALSPEAK type separation scheme. However, no literature report is available on the selective extraction of An3+ in RTIL media using any of the well known 'N' and 'S' donor extractants (Figure 1) mentioned above. The 10 evaluation of RTILs in the Ln³⁺/An³⁺ separation using BTP based 'N' donor ligands is of particular interest due to two main reasons, viz. (i) to avoid the use of toxic and hazardous long chain alcoholic solvents to solubilize the BTP derivatives in molecular solvents, viz. n-dodecane or kerosene. (ii) to look for 15 the possibility of improvements in the separation behaviour of An³⁺ and Ln³⁺ ions using the BTP - RTIL solvent systems as differential coordination behaviour of these two classes of 'f block elements is noticed in RTIL medium.

 $_{20}$ **Figure 1:** Promising soft (S, N) donor ligands for the selective extraction of $\mathrm{An^{3+}}$ over $\mathrm{Ln^{3+}}$

In the present paper, a systematic study on the separation of Am³⁺ and Eu³⁺ ions has been carried out using several substituted BTP 25 ligands viz., Me-BTP, Et-BTP and *n*-Pr-BTP in [C_nmim][NTf₂] (where n = 2,3,4,6,8). Time resolved fluorescence spectroscopy (TRFS) studies have also been carried out with the organic extract of Eu³⁺ ion in order to understand the difference in its extraction behaviour in the ionic liquid vis-à-vis molecular 30 diluent (*n*-dodecane) based solvent systems. Computational studies were also carried out on the Am³⁺ and Eu³⁺ complexes in order to find out the differences in the metal-ligand orbital overlap and bond distances in these two metal complexes.

Experimental

35 Reagents and Chemicals

The ligands (Me-BTP, Et-BTP and *n*-Pr-BTP) were synthesized following the procedure described elsewhere [22] and characterized by elemental analysis, melting point and NMR (¹H as well as ¹³C) spectral analysis. ²⁴¹Am was purified by a method reported earlier [23] and the purity was checked by alpha spectrometry. ^{152,154}Eu was procured from the Board of Radiation and Isotope Technology (BRIT), Mumbai, India and was used after ascertaining its radiochemical purity. Suprapur nitric acid (Merck) and MilliQ water (Millipore) were used for preparing the tracer solutions. High purity crystals (99.99%) of Eu(NO₃)₃.5H₂O (Sigma-Aldrich) were used to prepare the organic extract of Eu³⁺ ion for the luminescence studies.

Distribution Studies

Distribution studies were carried out with 241 Am and 152,154 Eu as 50 the tracers spiked in aqueous solutions containing varying nitric acid concentrations from 0.1-1.0 M. The strip solution from the

'actinide partitioning' step which is usually at ~0.2 M HNO₃ when the diglycolamides are used as the extractant for actinide partitioning [24] is proposed to be used as the feed for the 55 lanthanide – actinide separation. It was, therefore, thought of interest to limit HNO₃ concentration in the range of 0.1 M - 1.0 M in the present studies. The ionic liquid solvents were prepared using 0.01 M solutions of Me-BTP, Et-BTP or n-Pr-BTP in $[C_n mim][NTf_2]$ (where n = 2,3,4,6,8). Equal volumes (0.25 mL) 60 of the organic and the aqueous phases were taken in a leak-tight stoppered tube and agitated in a thermostated water bath at 25±0.1°C for 2 hours. The two phases were then centrifuged and assayed by taking suitable aliquots (0.1 mL) from both the phases followed by their radiometric assay by gamma ray counting using 65 a NaI(Tl) scintillation detector. In the cases of mixture of radiotracers, HPGe detector was used. The distribution ratio for a given metal ion (D_M) was calculated as the ratio of counts per minute of the corresponding radiotracer per unit volume in the organic phase to that in the aqueous phase. The separation factor $_{70}$ (S.F.) is defined as D_{Am} / D_{Eu} . Material balance was within the error limits of $\pm 5\%$.

Luminescence Study

The luminescence studies of the organic extracts of the Eu³⁺ ion by Me-BTP, Et-BTP and *n*-Pr-BTP in 2-bromooctanoic acid/*n*-75 dodecane and in [C₄mim].[NTf₂] were performed using an Edinburgh FLS 900 unit provided with CD-920 controller and micro sec (Xe) flash lap. The data acquisition and analysis were done by the F-900 software provided by Edinburgh Analytical Instruments, UK. The fluorescence decay curves for all the samples were recorded on 12 ms scale and fitted via an iterative method.

Computational Study

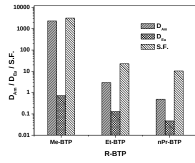
Gas phase geometries of the Am³⁺ and Eu³⁺ complexes of Me-BTP were optimized at the GGA level of density functional stheory (DFT) by using Becke's exchange functional [25] in conjunction with Perdew's correlation functional [26] (BP86) with generalized gradient approximation (GGA) where 28 and 60 electron core pseudopotentials (ECPs) along with the corresponding def-SV(P) basis set were selected for the Eu³⁺ and on Am³⁺ ions, respectively. All other lighter atoms were treated at the all electron (AE) level. All the calculations were performed using the TURBOMOLE program package [27-28]

Results and Discussion

Distribution Studies in $[C_n mim]$. $[NTf_2]$

⁹⁵ Distribution studies of Am^{3+} and Eu^{3+} were carried out using the R-BTP derivatives (R = Me, Et and *n*-Pr) in [C_nmim][NTf₂] media (n = 2,3,4,6 and 8) from 0.1, 0.5 and 1.0 M HNO₃ medium (Figure S1-S6 in supporting information).

Figure 2: Extraction and separation behaviour of Am^{3+} and Eu^{3+} from 0.1 M HNO₃ medium using 0.01 M R-BTP in $[C_8mim]$.[NTf₂]



Am3+ selectivity of Me-BTP over Eu3+ was found to be enormously high in [C_nmim][NTf₂] media, which is clearly shown in figure 2 and table 1.

Table 1: Comparison of two phase extraction and separation behaviour of Am³⁺ and Eu³⁺ using R-BTP in molecular solvents and RTIL media

Ligand (L)	Org. Phase	Aq. Phase	D_{Am}	S.F.	Ref
Me-BTP	0.004 M L + 1 M BDA/ TCE	0.013 M HNO ₃	213	24	4
	0.02 M L + 1 M BOA /n- dodecane	0.4 M HNO ₃	0.15	41	30
	0.01 M L in [C ₄ mim].[NTf ₂]	0.1 M HNO ₃	231	412	PW
	0.01 M L in [C ₈ mim].[NTf ₂]	0.1 M HNO ₃	>2000	>3000	PW
Et-BTP	0.004 M L + 1 M BDA in TCE	0.013 M HNO ₃	420	26	4
	0.02 M L + 1M BOA/ n- dodecane	0.4 M HNO ₃	7.31	66	30
	0.01 M L in [C ₄ mim].[NTf ₂]	0.1 M HNO ₃	2.9	14.5	PW
	0.01 M L in [C ₈ mim].[NTf ₂]	0.1 M HNO ₃	2.97	23	PW
nPr-BTP	0.0344 M L in TPH/EhOH (4/1)	0.3 M HNO ₃ + 1.6 M NH ₄ NO ₃	45.3	143	4
	0.01 M L in [C ₄ mim].[NTf ₂]	0.1 M HNO ₃	0.67	9	PW
	0.01 M L in [C ₈ mim].[NTf ₂]	0.1 M HNO ₃	0.49	10.4	PW

BDA: 2-bromo decanoic acid; BOA: 2-bromo octanoic acid; EhOH: 2ethyl-hexanol; TCE: 1,1,2,2-tetrachloroethane; PW: Present work; S.F. = $D_{\text{Am}}\!/D_{\text{Eu}}$

10 The S.F. values were found to be in the range of several thousands using Me-BTP even with 0.1 M HNO₃ and as shown from table 1, such a high S.F. value for Am³⁺ over Eu³⁺ is unprecedented with any of the BTP derivatives in molecular diluents. Carbamoylmethylenephosphine oxide (CMPO), an 15 extensively studied ligand for the actinide partitioning does not show any selectivity between trivalent actinides and lanthanides [1]. However, Naganawa et al. reported a S.F. value of 30 for Am³⁺ over Eu³⁺ with CMPO in the presence of the weakly complexing hydrophobic anion, tetrakis[3,5-20 bis(trifluoromethyl)phenyl] borate (TFPB-) [29]. In a previous study, we reported an enhancement in the selectivity (S.F. = ~500) for Am³⁺ over Eu³⁺ using Et-BTP in presence of a hydrophobic weak anion, chlorinated cobalt dicarbollide (HCCD). Et-BTP, otherwise, showed a S.F. value of <100 in 25 presence of 2-bromocarboxylic acid [30]. The results from the aqueous phase nitric concentration variation studies (Fig. 3) indicated a decrease in the D_M and a concomitant decrease in the S.F. values with increasing HNO₃ concentration. The decrease in D_M with increasing HNO₃ concentration agreed with our previous 30 results in molecular diluents with the difference that the metal ion extraction was much higher even with a significantly lower ligand concentration. Difference in complexation behaviour of An³⁺ and Ln³⁺ with azide ion was reported to be higher in $[C_4 mim][NTf_2]$ as compared to that in aqueous medium. [20]. 35 Furthermore, increase in metal ion extraction with increasing alkyl functionality in the ionic liquids in case of Me-BTP did not support the conventional ion-exchange mechanism and an ionpair mechanism may be proposed as given below.

$$M_{aq}^{3+} + n L_{IL} + 3 NTf_{2IL}^{-} == [M(L)_n]^{3+} [NTf_2]_{3,IL}^{-}$$

40 The lower extraction with ionic liquids having smaller alkyl groups may be attributed to lower solubility of the bulky metal ligand ion-pair complexes containing 3 extractant molecules in the ionic liquid phase. On the other hand, in most of the other cases involving Et-BTP or n-Pr-BTP for the extraction of either 45 Am³⁺ or Eu³⁺, the extraction with [C_nmim][NTf₂] was higher with n = 2 or 3 and lower extraction was seen with higher alkyl chains (n = 4, 6 or 8) suggesting the scope for cation-exchange mechanism of the type:

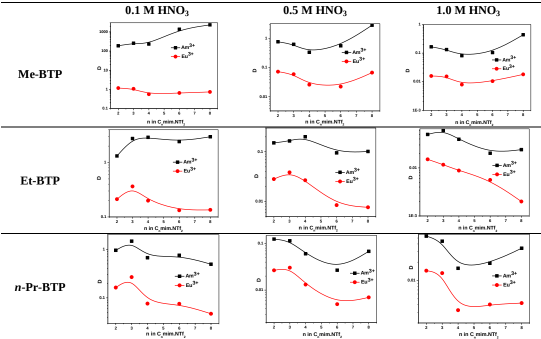
$$M^{3+}_{\ aq} + n \ L_{IL} + 3 \ C_n mim^+_{\ IL} = = \left[M(L)_n \right]^{3+}_{\ IL} + 3 \ C_n mim^+_{\ aq}$$

50 Though both extraction mechanism can be conveniently suggested based on the dependence of metal ion extraction with HNO₃ concentration in a particular RTIL, the higher concentration of nitric acid is likely to decrease metal ion extraction due to protonation and subsequent aqueous partitioning 55 of the extractant. Furthermore, as the acidity of the aqueous phase increases, the nitrate ion activity increases and due to the higher complexing nature of nitrate ion as compared to the weakly coordinating NTf2 ion of RTIL, metal ion interaction with R-BTP ligands decreases, which results in a concomitant decrease 60 in the selectivity.

An interesting difference in the trends of the D_M and S.F. values was observed when the R-BTP ligands with varying 'R' groups are compared in molecular diluents and RTIL (Table 1). An increase in the D_M values with increasing the size of the 'R' 65 group from methyl (Me) to ethyl (Et) in n-dodecane medium was reported in the presence of 2-bromo-octanoic acid [4,30]. In the present work, however, we observe a significant decrease in the D_M values with increasing size of the 'R' group from methyl to ethyl and from ethyl to n-propyl which is in sharp contrast to the 70 previously reported trend. This has been largely attributed to the differences in the extracted complex stoichiometry. It was interesting to note that while 1:1 (M:L) species were found to be extracted in molecular diluent systems, 1:3 (M:L) species were extracted in ionic liquids. In molecular diluents, the increasing 75 hydrophobicity of the ligand with increasing alkyl chain length favours the partitioning of the metal-ligand complex in the organic phase resulting in the enhancement in D_M values. This has been corroborated by luminescence studies which indicated 1:1 stoichiometry of the extracted complex (vide supra), where 80 the role of increasing steric hindrance with increasing size of the alkyl substituent is expected to be less pronounced. On the other hand, in the RTIL medium, the partitioning of the metal-R-BTP complex in the organic phase is facilitated by the presence of bulk concentration of the hydrophobic anion, NTf₂. Moreover, 85 the luminescence studies showed the formation of D₃ symmetric 1:3 Eu³⁺ complex and, therefore, the increasing alkyl chain length hinders the approach of three such bulky ligands resulting in sterically strained geometry which in turn is reflected in lower D_M values. In order to further understand the unusually high 90 selectivity of Me-BTP as compared to Et-BTP and n-Pr-BTP in RTIL medium, Am³⁺ and Eu³⁺ extraction studies were carried out varying the ligand (RBTP) concentration in [C₄mim][NTf₂] medium to identify the nature of the extracted species (Figure S7-S9 in supporting information). The results indicated that Eu³⁺ was 95 extracted as their respective 1:2 complexes with all the three ligands studied. Am^{3+*} was, however, extracted as 1:3 complex with Et-BTP and n-Pr-BTP. In case of Me-BTP, on the other hand, it is extracted as an unusual 1:4 complex, which resulted in unusually high selectivity of Me-BTP for Am3+ over Eu3+ in 100 RTIL medium due to the extraction of a significantly more hydrophobic complex of Am³⁺ (compared to the respective Eu³⁺

complex). Formation of 1:4 complex for the Am³⁺

Figure 3: Effect of alkyl chain length (n) in the C_n mim.NTf₂ on the extraction of Am^{3+} and Eu^{3+} using RBTP at different HNO₃ concentration



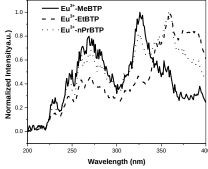
extraction with Me-BTP was, however, could not be explained on 5 the basis of well known facts as it required Am³+ coordination number of 12 for binding of all the four Me-BTP ligands in a tridentate manner. Am³+ extraction in the [Cnmim][NTf2] medium was, therefore, more favoured with respect to the Eu³+ extraction as the hydrophobic chain length (n) increased. The weakly coordinating NTf2 ion of RTIL cannot compete with the R-BTP ligands for the metal ion coordination and hence, due to more affinity of the R-BTP ligands for Am³+ over Eu³+ ion higher metal-ligand stoichiometry in case of Am³+ (1:4 for Me-BTP) as compared to Eu³+ (1:2 for Me-BTP) ion was observed which resulted in improved selectivity for Am³+ over Eu³+ in Cnmim.NTf2 medium.

Luminescence Study

In molecular diluents

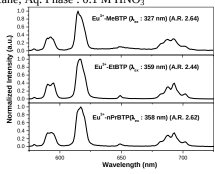
Luminescence studies were carried out with the $\mathrm{Eu^{3+}}$ extract in the organic phase containing 0.01 M R-BTP + 1 M 2-bromocotanoic acid in n-dodecane. Figure 4, shows the excitation spectra of the $\mathrm{Eu^{3+}}$ -R-BTP complexes.

Figure 4: Excitation spectra of Eu³⁺ complexes in the organic ²⁵ extract ($\lambda_{em} = 616$ nm): Org. Phase: 0.01 M RBTP + 1 M 2-bromooctanoic acid in *n*-dodecane; Aq. Phase : 0.1 M HNO₃



The relative intensities of various $f \rightarrow f$ transitions alter in these complexes. In case of Eu³⁺-Me-BTP complex, λ_{max} is at 326 nm whereas in the cases of Eu³⁺-Et-BTP and Eu³⁺-n-Pr-BTP complexes, the λ_{max} values are at 359 and 358 nm, respectively. The emission spectra of these complexes are shown in Figure 5, which indicates similar spectra for the Eu³⁺ complexes with all the three ligands. The asymmetry ratio (A.R.>2) values are quite high indicating the formation of asymmetric complexes. In order

Figure 5: Emission spectra of Eu^{3+} complexes in the organic extract: Org. Phase: 0.01 M RBTP + 1 M 2-bromooctanoic acid in *n*-dodecane; Aq. Phase: 0.1 M HNO₃



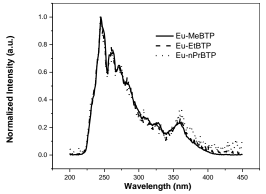
to get more insight into the stoichiometry of the complexes, their lifetimes were measured by recording the decay profiles (Figure S10) of the Eu³⁺-R-BTP complexes which are extracted in the *n*-45 dodecane phase in the presence of 2-bromooctanoic acid, which fitted to a single exponential decay pattern indicating formation the 1:1 complexes similar to those reported in the literature [31]. The 2-bromooctanoic acid, being a stronger anion, can compete with the R-BTP ligands. The presence of more than one unit of the R-BTP molecule in the inner sphere of the Eu³⁺ ion is, therefore, not observed in the present case.

In $[C_4mim][NTf_2]$

The results of the luminescence studies carried out using the Eu³⁺ extract in ionic liquid based solvents (R-BTP in [C₄mim].[NTf₂]) are entirely different from that observed in *n*-dodecane medium.

5 The excitation spectra (Figure 6) indicate that the ligand to metal charge transfer band at 247 nm is much stronger as compared to the $f \rightarrow f$ transition bands. This clearly indicates strong ligand sensitized luminescence of Eu³⁺ in its R-BTP complexes in the RTIL medium. Such a strong ligand sensitization was not 10 observed in molecular diluent (vide supra). The emission spectra (Figure 7) of all the three Eu³⁺-RBTP complexes are similar with A.R. values of ~1 where the peak at ~580 nm due to the ${}^5D_0 \rightarrow {}^7F_0$ transition is completely absent. This is indicative of D_n, D_{nd} or D_{nh} site symmetry around the Eu³⁺ ion [32-33]. The doublet at

Figure 6: Excitation spectra of Eu³⁺ complexes in the organic extract ($\lambda_{em} = 616$ nm): Org. Phase: 0.01 M RBTP in [C₄mim][NTf₂]; Aq. Phase: 0.1 M HNO₃



 $_{20}$ ~593 nm due to the $^5D_0 \rightarrow ^7F_0$ transition and singlet at ~616 nm due to the ${}^5D_0 \rightarrow {}^7F_0$ transition is the signature of D₃ symmetric 1:3 complexes [34-35]. The anion present in bulk (NTf₂) in the RTIL medium is a weak anion and, therefore, cannot compete with the R-BTP ligands. The metal coordination is, therefore, 25 saturated by three R-BTP ligands resulting in nine coordination numbers around the Eu $^{3+}$ ion. This is supported by very high τ values (~2 ms or higher). Similar high τ values suggesting complete dehydration of the inner coordination sphere has been previously reported [34-35]. This was, however, not consistent 30 with the results of distribution studies, which indicated the presence 1:2 complex of Eu³⁺.

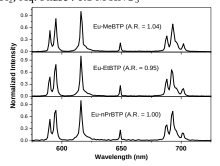
Table 2: Lifetime values (in µs) of Eu³⁺ complexes of R-BTP observed in the organic extract of molecular diluent and RTIL

	0.02 M Ligand +		
Licand	1 M 2-bromo-	0.02 M Ligand in	
Ligand	octanoic acid in	$[C_4mim].[NTf_2]$	
	n-dodecane		
Me-BTP	419 ± 2	2154 ± 2	
Et-BTP	692 ± 1	2576 ± 5	
nPr-BTP	625 ± 1	1947 ± 3	

The lifetime spectra of all the three Eu³⁺ complexes are given in Figure S11 and large lifetime values (Table 2) for all the three R-BTP complexes conform to such high stoichiometric (1:3) complexes [31] in the RTIL medium. This unusual behaviour 40 could be rationalized based on the assumption that due to the high

ligand sensitization, the fluorescence yield of 1:3 complex was much higher as compared to the 1:2 complex. The 1:3 complex, in spite of its negligible concentration was, therefore, dominated in the fluorescence spectra of the Eu³⁺ complexes of the R-BTP 45 ligands in its organic extract in RTIL medium.

Figure 7: Emission spectra of Eu³⁺ complexes in the organic extract ($\lambda_{ex} = 247$ nm): Org. Phase: 0.01 M RBTP in C₄mim.NTf₂; Aq. Phase: 0.1 M HNO₃

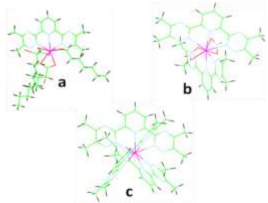


Computational Study

Geometry Optimization

The "M-N" bond distances calculated for Am(Me-BTP)₃³⁺ complex in the present study were found to be very close to those obtained for the $Am(n-Pr-BTP)_3^{3+}$ complex from the EXAFS study [31] and the $Cm(H-BTP)_3^{3+}$ complex obtained from DFT calculations [36]. The alkyl (R) groups, therefore, have very little effect on the structure and bonding of the Am3+ and Eu3+ complexes of R-BTP. Me-BTP can, therefore, be considered a 60 good model compound for the other R-BTP (Et-BTP and n-Pr-BTP) derivatives studied in the present work.

Figure 8: Optimized geometries of different Am³⁺ /Eu³⁺ complexes of MeBTP



65 In the present work, all the computational studies were, therefore, carried out on the Am³⁺ and Eu³⁺ complexes of Me-BTP. Optimized geometries of various Am³⁺ and Eu³⁺ complexes of Me-BTP with different stoichiometries (1:1 as obtained in molecular solvent (n-dodecane and 2-bromo-octanoic acid), 1:2 70 and 1:3 as obtained in RTIL medium) were shown in figure 8. The bond distances between Am³⁺/Eu³⁺ ion and coordinating 'N' atoms (central pyridinyl 'N' atom (N_c) and lateral triazinyl 'N' atoms (N₁)) were listed in table 3. It is to be noted here that 'M-N_c' distances were found to be decreasing as we go from 1:1 to 75 1:3 complex. This was because of the fact that 1:1 complex contained three anionic (2-bromo-octanoic acid) ligands, which lowered the charge on the central metal ion, thereby lowering its interaction with the electron donor 'Nc' atom of the ligand.

Table 3: 'M-N' bond distances in the optimized geometries of different Am³⁺ and Eu³⁺ complexes of MeBTP

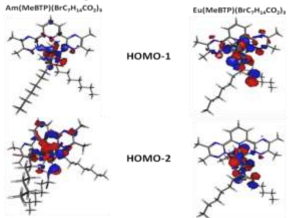
M ³⁺	M(L)(BOA) ₃	$[M(L)_2(NO_3)_2]^+$	$[M(L)_3]^{3+}$
Am ³⁺	$Am-N_c = 2.676$	$Am-N_c = 2.624(0)$	$Am-N_c = 2.595(3)$
	$Am-N_l = 2.59(3)$	$Am-N_l = 2.575(4)$	$Am-N_1 = 2.60(1)$
Eu ³⁺	Eu- $N_c = 2.737$	Eu- $N_c = 2.667(2)$	Eu- $N_c = 2.629(8)$
	Eu- $N_l = 2.629(5)$	Eu- $N_l = 2.629(5)$	Eu- $N_l = 2.63(1)$

L: Me-BTP: BOA: 2-bromo octanoic acid

In the 1:2 complex, two anionic (NO₃-) ligands are present and 5 therefore metal ion interacts with 'N_c' more strongly as compared to 1:1 complex. In the 1:3 complex, no anionic ligand is present in the inner coordination sphere of the metal ion. The interaction of metal ion with the ligand's electron donor (N_c) atom is, therefore, strongest in the tri-positive 1:3 complex with shortest 10 'M-N_c' bond distance amongst all the three possible complexes studied in spite of the presence of the three bulky Me-BTP ligands. Shorter 'Am-N' bonds as compared to the 'Eu-N' bonds were observed in all the cases.

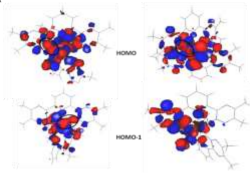
Molecular Orbitals Analysis

- 15 Shorter 'Am-N' bonds as compared to the 'Eu-N' bonds in all the three kinds (1:1, 1:2 and 1:3) of complexes could be understood by analyzing of the frontier orbitals of those Am3+ and Eu3+ complexes (Figure 9-11). In the M(Me-BTP)(2-BrC₇H₁₄CO₂)₃ complex, the bonding interactions of metal ions with Me-BTP
- 20 Figure 9: Frontier orbitals of Am3+ and Eu3+ complexes of MeBTP and 2-bromo-octanoic acid with 1:1 stoichiometry as obtained in n-dodecane medium showing the difference in metalligand orbital overlap



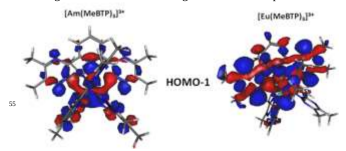
25 was mainly observed in the frontier molecular orbitals (HOMO-1 and HOMO-2) as shown in figure 9. These orbitals showed higher metal-ligand (Me-BTP) overlap in Am3+ complex as compared to that in Eu³⁺ complex. This was also reflected in the contributions of metal 'd' and 'f' orbitals in those frontier 30 molecular orbitals (Table 4), which showed higher 'd' and 'f' orbitals contributions for Am³⁺ as compared to that for Eu³⁺. Zaiter et al., have also reported higher orbital interaction in the Am³⁺ complex of pyrazine as compared to those in the Eu³⁺ complex showing higher Am³⁺-ligand back bonding [37]. Similar 35 features were also observed for the frontier molecular orbitals of $M(Me-BTP)_2(NO_3)_2^+$ (HOMO, HOMO-1) and $M(Me-BTP)_3^{3+}$ (HOMO-1), which mainly participated in the bonding with Me-

Figure 10: Frontier orbitals of Am³⁺ and Eu³⁺ complexes of 40 MeBTP with 1:2 stoichiometry (M(MeBTP)₂(NO₃)₂⁺) as obtained in RTIL medium showing the difference in metal-ligand orbital



In the $Eu(Me-BTP)_3^{3+}$ complex, both the 'd' and 'f' orbitals 45 contributions of Eu³⁺ were found to be lower as compared to that of Am³⁺ in the Am(Me-BTP)₃³⁺ complex.

Figure 11: Frontier orbitals of Am³⁺ and Eu³⁺ complexes of MeBTP with 1:3 stoichiometry as obtained in RTIL medium 50 showing the difference in metal-ligand orbital overlap



In order to have a quantitative idea about the comparative 'M-N' bond strength for Am3+ and Eu3+ complexes the molecular 60 orbitals were analysed to calculate the two center Mayer's bond order (Table S12 in the electronic supporting information), which showed higher bond orders in 'Am-N' bonds as compared to those in 'Eu-N' bonds in the cases of all the all the complexes studied.

65 Table 4: Total contribution of metal 'd' and 'f' orbitals in the frontier molecular orbitals of different Am3+ and Eu3+ complexes of MeBTP calculated using c² population analysis (SCPA) method [38]

Complex	Frontier MO	d%(Am/Eu)	f%(Am/Eu)
M(L)(DOA)	HOMO-1	1.1/0.5	27.3/15.0
$M(L)(BOA)_3$	HOMO-2	0.8/0.4	35.7/29.7
$[M(L)_2(NO_3)_2]^+$	НОМО	2.5/0.7	23.2/16.0
$[M(\mathbf{L})_2(N\mathbf{O}_3)_2]$	HOMO-1	2.4/0.8	35.6/24.4
$[M(L)_3]^{3+}$	HOMO-1	2.4/0.2	25.2/1.1

The molecular orbitals analysis, therefore, clearly showed the 70 preference of this class of ligands towards An³⁺ over Ln³⁺. This could be the reason for the tendency of Am3+ to form higher stoichiometric complexes with R-BTP as compared to Eu³⁺ [34]. This has been reflected in the present work, where Am³⁺ is extracted as 1:3 complex for Et-BTP and n-Pr-BTP, whereas Eu³⁺ 75 is extracted as 1:2 complex only. Me-BTP, moreover, forms extra

ordinarily high stoichiometric and unusual (1:4) complex with Am^{3+} . Further studies are required to investigate the structure of this Am^{3+} complex in detail.

Conclusions

5 The role of RTIL as a diluent for N-donor ligands (R-BTPs) for the mutual separation of An3+ and Ln3+ ions is explored from the nitric acid medium for the first time. Significant enhancement in the selectivity is observed as compared to that in molecular diluents with a S.F. value as high as >3000 using Me-BTP as the 10 extractant. The enhanced selectivity in RTIL medium is explained on the basis of the formation of extractable species with higher metal-ligand stoichiometry in case of Am³⁺ as compared to that in case of Eu³⁺. The nature of the complexes extracted in RTIL vis-à-vis n-dodecane media is characterized by 15 the TRFS technique. Computational studies showed higher 'd' and 'f' orbital participation with higher metal-ligand orbital overlap for Am³⁺ complex as compared to that for Eu³⁺ complex in the bonding with Me-BTP. The results of the present work show the possibility of application of RTIL as the promising 20 diluent for the Ln3+/An3+ separation using 'N' donor heteropolycyclic ligands, which are otherwise not soluble in hydrocarbon based suitable molecular solvent, viz. n-dodecane. It will, therefore, be of interest to explore the selective extraction of An³⁺ over Ln³⁺ using soft donor ligands (R-BTBP, R-BTPhen 25 and dithiophosphinic acids) which are known to be more selective than R-BTP derivatives studied in the present work.

Acknowledgement

The authors (A.B., M.M. and P.K.M.) would like to thank Dr. A. Goswami, Head, Radiochemistry Division for his continuous support and encouragement during the course of this work. The authors also thank the supercomputing facility, BARC for carrying out the computational studies.

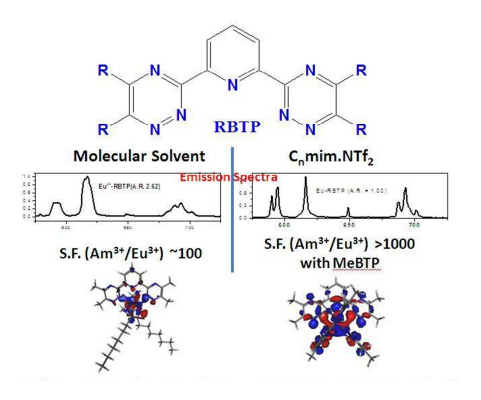
Notes and references

- 35 Authors for Correspondence: arunasis12@yahoo.co.in (A.B.) and mpatra@barc.gov.in (P.K.M.)
 - ^{1.} Radiochemistry Division; ^{2.} Bioorganic Division; Bhabha Atomic Research Centre, Trombay, Mumbai-400085, India
- † Electronic Supplementary Information (ESI) available: **Distribution**⁴⁰ **studies: Figure S1:** Effect of aqueous phase acidity on the extraction of Am³⁺ and Eu³⁺ by 0.01 M MeBTP in different C_nmim.NTf₂; **Figure S2:** Effect of aqueous phase acidity on the selectivity of Am³⁺ over Eu³⁺ by 0.01 M MeBTP in different C_nmim.NTf₂; **Figure S3:** Effect of aqueous phase acidity on the extraction of Am³⁺ and Eu³⁺ by 0.01M EtBTP in
- 45 different C_nmim.NTf₂; Figure S4: Effect of aqueous phase acidity on the selectivity of Am³⁺ over Eu³⁺ by 0.01M EtBTP in different C_nmim.NTf₂; Figure S5: Effect of aqueous phase acidity on the extraction of Am³⁺ and Eu³⁺ by 0.01M *n*-PrBTP in different C_nmim.NTf₂; Figure S6: Effect of aqueous phase acidity on the selectivity of Am³⁺ over Eu³⁺ by 0.01M *n*-
- 50 PrBTP in different C_nmim.NTf₂; Figure S7: Effect of MeBTP concentration on the Am³⁺ and Eu³⁺ extraction; Org. phase: 0.004-0.02M MeBTP in C4mim.NTf₂; Aq. Phase: 0.1M HNO₃; Figure S8: Effect of EtBTP concentration on the Am³⁺ and Eu³⁺ extraction; Org. phase: 0.004-0.02M EtBTP in C4mim.NTf₂; Aq. Phase: 0.1M HNO₃; Figure S9:
- 55 Effect of nPrBTP concentration on the Am³+ and Eu³+ extraction; Org. phase: 0.004-0.02M nPrBTP in C4mim.NTf₂; Aq. Phase: 0.1M HNO₃; **Luminescence studies: Figure S10:** Decay of Eu³+ complexes in the organic extract ($\lambda_{ex} = 327$ nm (for Eu-MeBTP), 359 nm (for Eu-EtBTP) and 358 nm (for Eu-nPrBTP) and $\lambda_{em} = 616$ nm):
- 60 Org. Phase: 0.01M RBTP + 1M 2-bromooctanoic acid in n-dodecane; Aq. Phase: 0.1M HNO₃; Figure S11: Decay of Eu³⁺

complexes in the organic extract ($\lambda_{ex} = 247$ nm and $\lambda_{em} = 616$ nm): Org. Phase: 0.01M RBTP in C₄mim.NTf₂; Aq. Phase: 0.1M HNO₃; **Table S12:** Two center Mayer's bond order in the 'M-N' 65 bonds in different Am³⁺ and Eu³⁺ complexes of Me-BTP

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Different Complexation behaviour in RTIL medium resulted in enhancement in selectivity upto few thousands with MeBTP $196 \times 141 \text{mm}$ (96 x 96 DPI)