NMR and TRLFS studies of Ln(III) and An(III) C5-BPP complexes†

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C5-BPP is a highly efficient N-donor ligand for the separation of trivalent actinides, An(III), from trivalent lanthanides, Ln(III). The molecular origin of the selectivity of C5-BPP and many other N-donor ligands of the BTP-type is still not entirely understood. We present here the first NMR studies on C5-BPP Ln(III) and An(III) complexes. C5-BPP is synthesized with 10% 15N labeling and characterized by NMR and LIFDI-MS methods. 15N NMR spectroscopy gives a detailed insight into the bonding of C5-BPP with lanthanides and Am(III) as a representative for trivalent actinide cations, revealing significant differences in 15N chemical shift for coordinating nitrogen atoms compared to Ln(III) complexes. The temperature dependence of NMR chemical shifts observed for the Am(III) complex indicates a weak paramagnetism. This as well as the observed large chemical shift for coordinating nitrogen atoms show that metal–ligand bonding in Am(C5-BPP)3 has a larger share of covalence than in lanthanide complexes, confirming earlier studies. The Am(C5-BPP)3 NMR sample is furthermore spiked with Cm(III) and characterized by time-resolved laser fluorescence spectroscopy (TRLFS), yielding important information on the speciation of trace amounts of minor complex species.

Introduction

In 2010 about 13% of the world’s electricity is supplied by nuclear power plants,1 producing 10 500 tons of spent nuclear fuel annually.2 Among the major challenges of used nuclear fuel are the long-term radiotoxicity and long-term thermal power that are dominated by plutonium and the minor actinides (MA = Np, Am, and Cm).

Both problems are addressed by the Partitioning and Transmutation strategy (P&T)3 that could have a beneficial impact on the design of a safe final repository.4–6 It involves separating plutonium and the minor actinides from the used fuel and converting them into shorter-lived fission products by neutron-induced nuclear reactions. In this context the separation of trivalent actinides An(III) from fission lanthanides Ln(III) is the key step, as some lanthanides have high neutron cross sections, consequently diminishing the efficiency of the transmutation step. Due to the similarity of Am(III) and Ln(III) both in chemical properties and ionic radii, highly selective extracting agents are needed to achieve a reasonable separation.5

It has been shown that extractants containing either soft sulfur or soft nitrogen donor atoms exhibit the required selectivity.6 Heterocyclic N-donor ligands derived from the terpyridine motif have shown higher complex strengths towards trivalent actinides than towards trivalent lanthanides.7 Among these, heteroaromatic nitrogen donor ligands 2,6-bis(1,2,4-triazine-3-yl)pyridines (BTPs) were the first extractants to achieve separation factors for Am(III) over Eu(III) higher than 100 from nitric acid solutions.7–9 They show good solubility in a range of organic diluents and form stable and isostructural 1 : 3 complexes with lanthanides and actinides.9–16 Furthermore, they co-extract nitrate anions from the aqueous phase and, unlike other similar extracting agents, do not need additional lipophilic anion sources such as 2-bromocarboxylic acid.17–19 In order to attain a fundamental understanding of the BTP-type ligands’ selectivity on a molecular level, the tridentate N-donor ligand C5-BPP was synthesized and tested for its extraction behavior.20 It was found that C5-BPP serves as a useful extracting agent with separation factors for Am(III) over Eu(III) over 100. However, it does not co-extract nitrate anions from the aqueous phase and is thus dependent on a lipophilic anion source. The ability to form stable 1 : 3 complexes and the different extraction behavior made C5-BPP an interesting target for investigations on the reason of the observed selectivity, especially in comparison to recent studies with nPrBTP.21

The molecular reason for the observed selectivity of some N-donor ligands is still largely unclear. A larger degree of covalence in the actinide–ligand bond, compared to lanthanide...
complexes, has been assumed to account for the observed extraction behavior. A more covalent bonding might result from a better overlap of the soft nitrogen lone pair with the diffuse 5f-orbitals of the actinide ions. In this case, the ratio of covalent to dative electrostatic bonding in actinide–N-donor complexes is expected to be larger than in isostuctural lanthanide compounds. Results from K-edge XAS spectroscopy on An(III) complexes with ligands containing sulfur, oxygen and chlorine seem to support this explanation.

Actinide compounds are a challenge for quantum chemistry due to various reasons, like for example the inclusion of relativistic effects. So far, prediction of bonding modes and NMR shifts is limited to simple systems and hardly implemented in commercial software packages. As an example for these problems, quantum chemical treatment of Am(III) and Eu(III) complexes with Cyanex 301 seemed to show a more covalent bonding in the actinide case, based on consideration of the bond length as a marker for covalence. Yet this produces misleading results, as calculated lanthanide–ligand bond lengths are too long in comparison to experimental data and bond lengths calculated by more sophisticated quantum-chemical methods.

Recently we were able to obtain the first NMR spectroscopic proof of a fundamentally different bonding mode in Am(III) complexes with N-donor ligands.

In general, NMR is an excellent spectroscopic method for the investigation of bonding interactions: the electrons of soft donor ligands can interact with positively charged cations either by sharing electron density in overlapping frontier orbitals or by electrostatic interactions. Both mechanisms lead to a rearrangement of electron density, which is monitored very precisely as the chemical shift in NMR spectroscopy. NMR focusing on the paramagnetism of the compounds allows the separation of the overall chemical shift into a part that is due to delocalization of electron spin density through covalent bonds (Fermi contact shift, FCS) and a distance- and angle-dependent part due to interaction of the anisotropic electron magnetic moment, assumed to be located at the metal ion, and the nuclear magnetic moment of ligand nuclei (pseudo contact shift, PCS). Currently, several methods for this separation of FCS and PCS are under investigation regarding their applicability to actinide complexes.

The scope of the work presented in this paper is to generate a reliable base of NMR spectra of diamagnetic and weakly paramagnetic C5-BPP lanthanide complexes and of the Am(III) complex. With these data, we aim to elucidate the bonding mode and potential bonding differences in lanthanide and actinide C5-BPP complexes, as this is expected to be the driving force of the ligand’s selectivity for actinide over lanthanide extraction.

Theoretical and NMR background

The chemical shift – and thus the electron distribution – of the coordinating nitrogen atoms are of particular interest for the investigation of bonding interactions. The effect of covalent bonding is especially pronounced here, since transferred electron density can normally only be detected over a few covalent bonds. Only in some cases nuclei more than three bonds away from coordinating atoms are influenced by FCS. Unfortunately, obtaining resonance signals in one-dimensional direct excitation spectra from 15N atoms at natural abundance is impossible in a time-effective manner. This is due to the fact that 15N has a low natural abundance of 0.364% and a low negative gyromagnetic ratio (γ = -0.28), resulting in low receptivity of the nucleus (about 1% of the 13C receptivity at natural abundance). Furthermore, a negative gyromagnetic ratio means that the Nuclear Overhauser Effect will decrease the signal intensity for 15N if 1H broadband decoupling is used.

In paramagnetic coordination compounds the overall experienced chemical shift δtot has several contributions:

\[ \delta_{\text{tot}} = \delta_{\text{dia}} + \delta_{\text{con}} + \delta_{\text{pc}} + \delta_{\text{anion}} \] (1)

δdia is the diamagnetic (or orbital) shift of the compound, δcon represents the Fermi contact shift, a through-bond effect, δpc is the pseudo contact shift that originates from coupling of the electron magnetic moment on the metal ion and the ligand nuclei spins and δanion is the influence of the counter-anion. All published methods for the separation of these terms have in common that they rely on an isostuctural diamagnetic analog to the paramagnetic complexes. The purely paramagnetic shift \( \delta_{\text{para}} = \delta_{\text{con}} + \delta_{\text{pc}} \) is calculated by simply subtracting the chemical shift values of the diamagnetic reference compound from the measured chemical shifts of the paramagnetic complexes (eqn (2)). If reference and paramagnetic complexes have the same counter-anion, δanion cancels out.

\[ \delta_{\text{para}} = \delta_{\text{tot}} - \delta_{\text{dia}} \] (2)

In the lanthanide series, La(III) and Lu(III) are diamagnetic ions and their complexes are generally used as diamagnetic reference compounds. Furthermore, Y(III) often forms complexes which are isostructural to the lanthanide compounds and can also be used as a reference.

In principle, paramagnetic compounds provide a detailed insight into the bonding mode via the separation of the observed paramagnetic chemical shift \( \delta_{\text{para}} \) into FCS and PCS. For this task, several methods have been proposed in the literature. Methods based on the chemical shift dependence on the temperature have been a matter of controversy and their application has to be evaluated very carefully. Currently, the standard procedure is the evaluation of the purely paramagnetic shift throughout the complete lanthanide series vs. tabulated lanthanide-depending constants, i.e. spin expectation values \( \langle S_i \rangle \) and Bleaney parameters \( C_{1n}, C_{2n}, \ldots \). Lanthanide shift reagents and lanthanide probes for protein structure determination have been widely used in NMR spectroscopy and thus lanthanide magnetic properties are quite well understood. This is not the case for magnetic properties of elements of the actinide series, thus Bleaney parameters \( C_{1n} \) and spin expectation values \( \langle S_i \rangle \) are unknown for these cations.

So far, only a small number of proton spectra and a few heteronuclear spectra of actinide containing compounds are observed...
available. These are largely limited to uranium in several oxidation states and hence there is a paucity of NMR studies on organic complexes with transuranium elements.

The magnetic properties of the free Am(\(\text{III}\)) ion are still a matter of debate in literature, as deviations from the expected diamagnetism arising from a predicted \(J = 0\) ground state have been found. Optical spectroscopy and DFT calculations show that the first non-diamagnetic excited states are some thousand wavenumbers higher in energy and thus thermally not populated and mixing of the states is not expected.\(^{49}\) This was also confirmed by experimental work on an [\(\text{Am(H}_{2}\text{O})_{6}\)](\(\text{CF}_{3}\text{SO}_{3}\))\(_{3}\) crystal in solid state, which exhibited a magnetic susceptibility curve that can be interpreted as non-magnetic behavior.\(^{49}\) On the other hand, surprisingly large magnetic susceptibilities and magnetic moments have been reported for different Am(\(\text{III}\)) compounds in the solid state, indicating that Am(\(\text{III}\)) is not purely diamagnetic.\(^{21-34}\) Recently, the magnetic susceptibility of Am(\(\text{III}\)) in perchloric acid solution was studied using the Evans NMR method.\(^{25}\) Results show a significant deviation from the expected magnetic behavior for Am(\(\text{III}\)) and Cf(\(\text{III}\)). Magnetic susceptibilities for both ions were found to be higher than expected.\(^{26}\)

In a recent publication, the effect of spin–orbit coupling on the alignment of spins in a magnetic field and the applicability of the Russel–Saunders coupling scheme was discussed.\(^{37}\) Spin expectation values for different configurations of Am(\(\text{III}\)) were calculated. The authors showed that the expected \(J = 0\) ground state has a spin expectation value \(\langle S_z \rangle = 0\) and contains the expected Hund’s multiplet to 63%. The energy difference to the \(J = 1\) state with \(\langle S_z \rangle = 0.5\) is only 0.24 eV, which is significantly lower than the expected value of 3.0 for six unpaired electrons. The authors conclude that there is a significant deviation from the multiplicites expected from Hund’s rules, but that pure \(j\)-\(j\) coupling cannot describe the electronic states as well.

In the case of the \(^{243}\text{Am(nPrBTP)}\)\(_{3}\)(\(\text{NO}_{3}\))\(_{3}\) complexes, we were able to acquire one- and two-dimensional \(\text{H}, \text{C}\) and \(\text{N}\) spectra in good quality.\(^{29}\) The observed linewidths in the spectra and the range of chemical shifts indicate that Am(\(\text{III}\)) has only a weak paramagnetism, with effects even smaller than observed for Sm(\(\text{III}\)). These results encouraged us to expand our NMR studies to the Am(\(\text{III}\)) complexes of C5-BPP.

### Synthesis

To compensate the unfavourable NMR spectroscopic properties of \(\text{\textsuperscript{15}N}\), we synthesized a \(\text{\textsuperscript{15}N}\) enriched C5-BPP ligand, \(\text{\textsuperscript{15}N}\)C5-BPP, in accordance to the already published \(\text{\textsuperscript{15}N}\) enriched nPrBTP ligand.\(^{23}\) The synthesis pathway is shown in Scheme 1. Successful labeling was confirmed by NMR spectroscopy and LIFDI-MS\(^{38,39}\) (cf. ESI†).

Using \(\text{\textsuperscript{15}N}\)C5-BPP, 1 : 3 complexes with lanthanides (La(\(\text{III}\)), Sm(\(\text{III}\)), Yb(\(\text{III}\)), Lu(\(\text{III}\))) and Y(\(\text{III}\)) were prepared. In order to compare these complexes to a 1 : 3 complex with a trivalent actinide we also prepared a \(\text{\textsuperscript{15}N}\)C5-BPP complex with \(^{244}\text{Am}\) (Fig. 1).

All complexes were prepared in deuterated methanol. Earlier studies on crystal structures of the Ln(\(\text{III}\)) complexes report that C5-BPP does not displace all nitrate anions from the inner coordination sphere of the central metal ions during crystallization.\(^{29}\) In our case \(\text{\textsuperscript{1}H}\) NMR spectroscopy on Ln(\(\text{III}\)) complexes showed that more than one complex species was formed in samples in which nitrate anions were present. Diffusion-ordered NMR spectroscopy (DOSY)\(^{40-42}\) proved that several complex species with varying diffusion coefficients were present. This is due to the fact that nitrate anions are strongly complexing ligands in pure organic solvents. The formation of numerous different complex species was overcome by using trilate salts (\(\text{OTf}^{-}, \text{CF}_{3}\text{SO}_{3}^{-}\)) for which the counter-anion has been shown to be non-coordinating. Indeed, in NMR spectra of C5-BPP lanthanide trilate complexes, only the desired 1 : 3 complex and, occasionally, small traces of a 1 : 2 complex, were found.

In order to perform NMR investigations using complexes with the same counter-anion, Am(\(\text{OTf}^{-}\)) was prepared from an Am(\(\text{NO}_{3}\))\(_{3}\) stock solution. Subsequently \(\text{\textsuperscript{15}N}\) labeled and unlabeled C5-BPP was used to synthesize [Am(C5-BPP)\(_{3}\)](\(\text{OTf}^{-}\)). To avoid potential magnetic impurities due to radioysis of the solvent and impurities from radioactive decay products we used the long-lived isotope \(^{243}\text{Am}\) (\(\text{I}_{1/2} = 7370\) a).

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Results and discussion

Diamagnetic Ln(m)–(C5-BPP)₃ complexes

As a first step in the investigation of bonding modes in C5-BPP complexes of lanthanide and actinide ions, we focused on diamagnetic or nearly diamagnetic compounds. A comparison of spectra of diamagnetic compounds is straightforward, as significant changes between isosctructural complexes can be attributed to a change in binding mode.

In our studies with nPrBTP we used the Lu(m) complex as a diamagnetic reference, since spectra of [La(nPrBTP)₃]NO₃ showed broadened spectral lines. This is due to a relatively weak coordination of nPrBTP to the large La(m) ion which decreases the complex symmetry and thus results in broad spectral lines. The bigger bite angle of the pyrazole nitrogen lone pairs in C5-BPP should enable this ligand to form structurally rigid complexes even with slightly larger cations. Indeed, we found that C5-BPP forms stable complexes with La(m), resulting in well-resolved NMR spectra with sharp lines.

Comparison of the three diamagnetic C5-BPP complexes (Y(m), La(m) and Lu(m)) shows that although all three metal ions are diamagnetic, there are significant differences in ¹H, ¹³C and ¹⁵N NMR chemical shifts. These differences are strongest in ¹H close proximity to the metal ion, and only very weak at the aliphatic side chain. Differences between proton spectra of Y(m) and Lu(m) complexes are small, with a maximum of 0.01 ppm at the H₄ triplet. The maximal discrepancy between proton signals of the La(m) and Lu(m) complexes is found for the signals of H₂/₅ with 0.04 ppm. The differences are more pronounced in ¹³C spectra. Again the spectra of Lu(m) and Y(m) complexes strongly resemble each other. Only for C₂/₆ (Δδ = 0.3 ppm) and C₁₀ (Δδ = 0.28 ppm) significant discrepancies are observed. Differences between the La(m) and Lu(m) complexes are stronger in particular for the quaternary carbon atoms C₅ (Δδ = 0.73 ppm), C₂/₆ (Δδ = 1.70 ppm), and C₁₀ (Δδ = 1.20 ppm).

As the influence of the central metal ion seems to be strongly dependent on the distance to the observed nucleus it should be even more pronounced on the nitrogen atoms. In ¹⁵N spectra we observe only weak shift differences for the non-coordinating N₉ (Y/Lu (Δδ = 0 ppm), La/Lu (Δδ = 1.2 ppm)). The coordinating nitrogen shifts show a stronger dependence on the central metal ion. The shift differences for N₁ (Y/Lu (Δδ = 1.0 ppm), La/Lu (Δδ = 4.0 ppm)) from ¹H, ¹⁵N-gHMOC spectra are smaller than for N₈ (Y/Lu (Δδ = 1.7 ppm), La/Lu (Δδ = 7.2 ppm)). These results coincide with the differences in ionic radii, which are quite similar for Y(m) (90.0 pm) and Lu(m) (86.1 pm), whereas La(m) is significantly larger (103.2 pm). Changes in the complex geometry and subsequently changed interaction between the metal ion and the ligand can explain the observed behavior.

These results clearly show that the diamagnetic reference compound needs to be chosen carefully, as the shift differences between La(m) and Lu(m) compounds are significant and several orders of magnitude larger than the spectral resolution. Based on our results we assume that La(m) is a good diamagnetic reference compound for the lighter part of the lanthanide series. The smaller metal ions Lu(m) and Y(m), which both have closed shells, are better suited as reference compounds for the heavier lanthanides. The error inferred from the reference compound on the determination of the purely diamagnetic chemical shift in strongly paramagnetic systems, where shifts of several hundred ppm can occur, are limited. However, the influence on weakly paramagnetic systems should not be underestimated.

It should be noted that in all diamagnetic lanthanide C5-BPP complexes and in the Y(m) C5-BPP complex, resonance signals for the coordinating nitrogen atoms N₈ and N₉ are generally found in a 12 ppm range around 266 ppm. For the non-coordinating nitrogen atom N₉, resonance signals are found in a narrow 2 ppm range around 206 ppm (cf. Table 1 and Fig. 2). In comparison to the free ligand, the coordination of C5-BPP to a M(III) cation hardly influences the chemical shift of N₈. N₉ on the other hand is shifted approximately 30 ppm upfield. This is due to the rearrangement of electron density upon complex formation. Unfortunately, in the free ligand no resonance signal is observed for N₉. Nevertheless, based on the diamagnetic lanthanide compound spectra, we would expect the resonance signal for N₈ in the same shift range as N₉. The same problems were encountered when we measured ¹⁵C spectra of the free ligand. We found that resonance signals for the quaternary carbon atoms C₂/₆, C₇, and C₁₀ are severely broadened and sometimes, as in the free ¹⁵N enriched ligand, unobservable in ¹D spectra. So far we do not have a clear explanation for this behavior.

Paramagnetic Ln(m)–(C5-BPP)₃ complexes

In the following we studied the influence of a weakly and a strongly paramagnetic central metal cation on the NMR spectra of the C5-BPP complexes. We used [Sm(C5-BPP)₃][OTf]₃ as a representative for a weakly paramagnetic ion (Sm³⁺; μeff = 0.85μ₀) and [Yb(C5-BPP)₃][OTf]₃ as a strongly paramagnetic ion (Yb³⁺; μeff = 4.54μ₀). With the ¹⁵N labeled ligand in hand, our focus was on the influence of paramagnetism on the resonance signals of the coordinating nitrogen atoms. In the Sm(m) complex, the N₉ resonance signal is observed at 205 ppm, i.e. without additional shift compared to the diamagnetic compounds. In contrast to the non-coordinating nitrogen, a

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* a Labeled values are taken from 2D ¹H,¹⁵N-gHMOC spectra of the 1:3 complexes with unlabeled C5-BPP.
larger shift is found for the coordinating nitrogen atoms. Compared to the La(m) complex, N₈ is shifted 45 ppm upfield and N₉ is shifted 48 ppm upfield. These values are in good agreement with observed shifts for nPrBTP complexes.²¹ Yb(m) complexes usually show the expected strong paramagnetic shifts, but paramagnetic relaxation enhancement for Yb(m) is still weak enough that spectral lines are not too broad to be observed and most multi-dimensional NMR experiments yield good results. Thus, unambiguous assignment of most signals is possible by heteronuclear correlation spectroscopy. However, due to the enhanced relaxation, the ¹⁵N signals for N₈ and N₉ in Yb(C₅-BPP)₃(OTf)₃ are only found after ¹⁵N-labeling. N₈ shows a notable shift of −10 ppm compared to the diamagnetic references, which can be attributed to the stronger PCS. The coordinating N₈ is shifted by approximately −200 ppm to 20 ppm.

A comparison of ¹⁵N direct excitation spectra of all investigated 1:3 M(m) C₅-BPP complexes is shown in Fig. 2. The N₉ signals (green circles) for the diamagnetic metal ions (Y(m), La(m), Lu(m)) show almost identical chemical shift values (green dotted line), while for N₈ (red circles) there is a notable difference (red dotted line). Furthermore, the chemical shift for the non-coordinating N₉ remains nearly constant even for the paramagnetic ions, while N₈ shows a strong dependency on the paramagnetism of the ion.

The larger shift in the Yb(m) and the Sm(m) cases can be attributed to a stronger PCS (especially for Yb, which predominantly exhibits PCS) but as well a non-negligible FCS. Heteronuclei directly bonded to paramagnetic cations have only scarcely been investigated with respect to the different contributions to the experienced paramagnetic shift. Most research is limited to protons in close proximity to the metal ion center. However, although the influence of FCS decreases rapidly along covalent bonds, it often cannot be neglected.²⁵ A strong impact of the FCS on directly coordinated nuclei can thus be expected, and, as in our case, might even contribute to a larger than expected share. Further research into this topic is necessary and currently under way in our group.

NMR-spectroscopy on Am(m)–(C₅-BPP)₃

The spectra of the Am(m)–C₅-BPP complexes with and without ¹⁵N labeling show that more than one complex species was formed. Upon addition of further ligand solution two of the complex species could be assigned to the free ligand and the
1 : 2 complex. Signals from the 1 : 3 complex, which forms the major species present in the sample, increase in intensity with increasing ligand-to-metal ratio. However, during titration another minor complex species that contains only one ligand molecule and a so far unknown contaminant not visible by NMR spectroscopy is formed. However, the NMR signals of the 1 : 3 complex as the major species could easily be identified and unambiguously assigned. To further elucidate the composition of the complex speciation we studied the samples by further NMR spectroscopic methods and time-resolved laser fluorescence spectroscopy (TRLFS, see below).

For a sample containing several different components, diffusion-ordered NMR spectroscopy (DOSY) is a versatile method. $^1$H DOSY spectra show three well separated complex species. $^{19}$F direct excitation spectra only show one signal at $-79.97$ ppm, which corresponds to the triflate anion. $^{19}$F DOSY spectra yield one diffusion coefficient for this peak which differs from the coefficients for the complex species calculated from $^1$H DOSY measurements. Thus a coordinated triflate anion or exchange between a bound and a free form can be excluded. All 1D spectra are well-resolved, and unambiguous assignment of the signals of the 1 : 3 Am(III) C5-BPP complex is possible. The complex is fully characterized by $^1$H, $^{13}$C and $^{15}$N direct excitation spectroscopy at different temperatures as well as a range of 2D heteronuclear correlation spectroscopy methods.

Information about magnetic properties and the bonding situation can be deduced by comparison of the Am(III) complex spectra and those of a diamagnetic reference compound. Unfortunately, the diamagnetic actinides Ac(m) and Lr(m) have short half-lives ($t_{1/2}^{232}$ Ac = 21.8 a, $t_{1/2}^{262}$ Lr = 3.6 h) and are not available in sufficient amounts. As we lack a diamagnetic actinide reference compound, we have to compare the Am(m) complex’s chemical shifts to those of Ln(m) complexes. This comparison is displayed in Fig. 3 for [Am($^{15}$N) C5-BPP]$_3$ OTf and [Sm($^{15}$N) C5-BPP]$_3$ OTf compared to [La($^{15}$N) C5-BPP]$_3$ OTf.

For most nuclei the effect of the Am(m) cation on the chemical shift is approximately ten times stronger than that of the Sm(m) cation. For the weakly paramagnetic Sm(m) a magnetic moment of $\mu_{\text{eff}} = 0.85\mu_B$ is known. Measurements using the Evans method yield a magnetic moment of $\mu_{\text{eff}} = 1.64\mu_B$ for Am(m). Recently, work on the influence of radioactive decay and radiolysis product formation on the accuracy of the Evans method has been published, suggesting a reduced magnetic moment of approximately $\mu_{\text{eff}} = 1.42\mu_B$. We therefore expect the paramagnetic influence of Am(m) to be stronger than the influence of Sm(m), but both should produce paramagnetic chemical shift effects in the same order of magnitude.

The large differences in the chemical shifts cannot be explained by the difference in the magnetic moments of the cations, but point to a fundamental change in the bonding mode. Fig. 4 shows a $^1$H,$^{15}$N gHMOC spectrum of [Am($^{15}$N) C5-BPP]$_3$ OTf. Indicated with the red boxes are the chemical shifts for the coordinating nitrogen atoms as expected from the free ligand and diamagnetic Ln(m) compounds. In the red circles are the measured values that differ vastly from the expectations. For the Am(m) complex, the immense shift differences of the coordinating nitrogen atoms ($N_1$: -256 ppm, $N_8$: -295 ppm) are noteworthy. Shifts of a comparable magnitude have only been found for a Yb(m) C5-BPP complex which has a considerably stronger effective magnetic moment ($\mu_{\text{eff}} = 4.54\mu_B$).

Furthermore, carbon atoms in both the Sm(m) and the Am(m) complexes show alternating positive and negative chemical shift differences along the carbon backbones of the ligands (see Fig. 4).
Fig. 3). This phenomenon is indicative of the simultaneous existence of spin polarization and spin delocalization at the ligand (polarized spin density delocalization). This spin delocalization would be due to a Fermi contact interaction between metal cation and N-donor ligands and thus to a share of covalence in the bonding. The pattern of the shift differences suggests that a part of the delocalized spin electron density resides in $p_\pi$ orbitals of the $sp^2$ hybridized carbon atoms. However, if this is true we would expect the signs of the chemical shift difference of the protons to be inverse to the attached carbons’ shift differences (two spins that are coupled electronically over one bond will have opposite signs). We find that for Sm(III) and Am(III) all pyridine proton shifts have the same signs. In both cases $H_4$ is shifted more towards deeper fields than the $H_{3/5}$ protons. This behavior suggests that two (or even more) different mechanisms take part in the delocalization of electron spin density, showing that the bonding between Am[III] and the soft N-donor ligand is a very complicated matter. An explanation for the downfield shift of $H_4$ could be that unpaired electron spin density is also transferred through $\pi$ bonds in the aromatic ring or the conjugated double bonds, respectively.

To gain insight into magnetic and bonding behavior, we acquired NMR spectra at different temperatures between 185 K and 335 K (cf. ESI†). In [La[$^{15}$N(C5-BPP)$_3$](OTf)$_3$, $N_8$ shows a temperature-dependent shift of 0.3 ppm. The $N_8$ signal shows strong line broadening (FWHM 18.54 ± 0.28 Hz) even at low temperatures. Thus, $N_8$ is only observable up to 315 K. In the monitored 130 K temperature range, the $N_8$ signal shows a temperature-dependent shift of −1 ppm.

In the case of the Y(III) C5-BPP complex, $N_8$ experiences a 0.5 ppm downfield shift between 185 K and 335 K, while $N_9$ (FWHM at 185 K: 19.10 ± 0.18 Hz) shows a −1.4 ppm upfield shift.

The temperature-dependent chemical shift of the Am[III] complex shows a different behavior (Fig. 5). The non-coordinating $N_8$ (FWHM at 285 K: 20.04 ± 0.19 Hz) shows an upfield shift of approximately 1 ppm at 275 K with increasing broadening of the resonance signal. This signal is not observable above 300 K, while a new doublet appears 0.8 ppm downfield of the last broad signal. Up to a temperature of 335 K this signal is again shifted upfield by 0.5 ppm. In total, the temperature-dependent shift of $N_8$ is 0.6 ppm. The coordinating $N_9$, however, shows a continuous 11.3 ppm downfield shift. This temperature-dependent shift is approximately ten times the shift measured for diamagnetic reference compounds, which is another distinct piece of evidence showing that Am[III] is not diamagnetic.

The observed chemical shift of the coordinating nitrogen atoms in the Am[III] complex cannot be explained by the different strength of the ions’ magnetic moments alone, as a comparison to the temperature dependent NMR spectra of the strongly paramagnetic [Yb[$^{15}$N(C5-BPP)$_3$](OTf)$_3$ shows. As the chemical shift of $N_8$ at room temperature is +20 ppm and thus close to the observed shift of the Am[III] complex, one could assume that the two ion’s paramagnetism were in the same order of magnitude. However, the complex with the more paramagnetic Yb[III] cation shows a larger chemical shift range upon temperature change: In the monitored temperature range, $N_8$ shows a 152 ppm shift, for the non-coordinating $N_9$ the shift is still −8 ppm. Furthermore, even in the weakly paramagnetic Sm[III] complex, $N_8$ shows a temperature-dependent shift of −162 ppm in the observed temperature range. Thus it is clear that the paramagnetism of the Am[III] ion is considerably weaker than at least in the Yb[III] ion and cannot satisfactorily explain the observed highfield shift of the $N_8$ signal in [Am[$^{15}$N(C5-BPP)$_3$](OTf)$_3$. The smaller temperature-dependent shift in the Am[III] complex, compared to Sm[III], could be due to a different ratio of covalent and dipolar bonding: FCS is transmitted through covalent bonds and has a linear temperature dependency. PCS, which can be associated to dipolar interactions, has a $T^{−2}$ dependency. However, as long as no clear separation of the chemical shift contribution can be performed, this has to be seen as indicative of a more covalent bond, but not yet as a proof.

The observed behavior of alternating chemical shift effects in the carbon backbone, but not on the protons in the ligand, points towards a combination of direct spin delocalization and polarized spin density delocalization. Both rely on a Fermi contact interaction arising from covalent bonding between the trivalent metal cation and the nitrogen atoms of the ligands. From comparison of the observed chemical shift differences in the slightly paramagnetic Sm[III] complex and the Am[III] complex, which cannot be explained by paramagnetism alone. We interpret this fact as indicative of an higher share of covalence in the actinide compound, which is consistent with recently reported XAS and EXAFS studies. Another effect that might explain the shift differences between Am[III] and the Ln[III] complexes is the existence of spin-orbit coupling effects on the metal ion which influences the shift of the nitrogen atom. Spin–orbit coupling is strongly dependent on the atomic number of the nucleus and is thus considerably stronger in the actinide series than for lanthanides. Spin polarization from spin–orbit coupling resembles spin-spin coupling effects in NMR spectroscopy that are mediated by s-type orbitals. This is another mechanism that could explain the substantial shifts on the nitrogen atoms and why the shift differences cannot be observed on neighboring atoms. As a consequence, both paramagnetic effects in the form of FCS and spin–orbit coupling seem to play an important role in the observed chemical shifts. Fermi-contact interactions and thus the existence of a certain covalence compared to the lanthanide compounds could thus explain the observed shifts on the nitrogen atoms of the americium complex.

**Cm[III] TRLFS studies to identify the minor complex species**

As shown above minor Am[III] complex species are formed in addition to the prevailing 1 : 3 Am[III] complex. Small amounts of impurities or minor complex species cannot be characterized using NMR spectroscopy. Hence we used a different spectroscopic method to elucidate the composition of the minor complex species. Addition of a trace amount (6.6 × 10$^{-8}$ mol L$^{-1}$) of Cm[III] to the [Am[$^{243}$C5-BPP)$_3$](OTf)$_3$. NMR sample enabled us to make use of the excellent fluorescence properties
of Cm(III). Furthermore, the chemical properties of Am(III) and Cm(III) are highly comparable which is for example reflected in very similar M(III)–N bond lengths in 1 : 3 complexes in solution4 (Am(nPr-BTP)₃ : Am(III)–N = 256 pm; Cm(nPr-BTP)₃ : Cm(III)–N = 257 pm).13,14

After addition of Cm(III) to the NMR sample the Cm(III) emission bands are recorded with increasing amounts of C5-BPP ligand. Upon addition of ligand solution to the Cm(III) spiked NMR sample the initial Am(III) and Cm(III) metal concentrations are diluted, and hence the ligand-to-metal ratio stepwise increases. The development of the Cm(III) fluorescence emission resulting from the ⁶D ⁷/₂ → ⁸S ⁷/₂ transition as function of the ligand-to-Am(III) ratio are shown in Fig. 6. The spectra are normalized to the same peak area for better comparison.

At an initial C5-BPP-to-Am(III) + Cm(III) ratio of 3 : 1 (corresponding to a ligand-to-Cm(III) ratio of 4.5 × 10⁻⁵ : 1) an emission band with a maximum at λₘₐₓ = 606.0 nm and two weak shoulders at λₘₐₓ = 603.2 nm and λₘₐₓ = 610.3 nm are observed.

With increasing ligand-to-metal ratio the intensity of the emission band at λₘₐₓ = 606.0 nm decreases significantly while both shoulders gain in intensity. At a final ligand-to-Am(III) + Cm(III) ratio of 6.8 : 1 two distinct emission bands with maxima at λₘₐₓ = 603.2 nm and λₘₐₓ = 610.6 nm are observed with an intensity ratio of approximately 1 : 1.

In earlier studies the emission bands of the Cm(III)–C5-BPP 1 : 1, 1 : 2 and 1 : 3 complex species in methanol were observed at λₘₐₓ = 603.7 nm, λₘₐₓ = 607.7 nm and λₘₐₓ = 611.6 nm, respectively.26 Hence, the observed emission bands at λₘₐₓ = 606.0 nm and λₘₐₓ = 610.6 nm are attributed to a Cm(III)–C5-BPP 1 : 2 and a Cm(III)–C5-BPP 1 : 3 complex species. The hypsochromic shi of 1.0 nm in comparison to the literature known 1 : 2 and 1 : 3 complex species are assigned to the use of a deuterated solvent and the high concentration of triflate anions.

With increasing ligand-to-metal ratio a decreasing ratio of the 1 : 2 complex species and an increasing ratio of the 1 : 3 complex species are observed, showing a stepwise complexation of Cm(III). The emission band at λₘₐₓ = 603.2 nm also gains in intensity upon increasing amount of C5-BPP, which confirms that does not result from the 1 : 1 Cm(III)–C5-BPP complex, and is attributed to a Cm(III) complex species with a minor impurity from the C5-BPP synthesis. At significantly higher metal
concentrations used for NMR studies these minor complex species do not play an important role and all signals of the 1:3 complex species can be assigned unambiguously (see above).

Conclusions

We present the first NMR study on a series of 1:3 complexes of Ln(III) and Am(III) with the tridentate N-donor ligand C5-BPP. A key step in our investigations was the synthesis of a C5-BPP molecule with $^{15}$N enrichment in the pyrazole moieties.

Using [$^{15}$N]C5-BPP we prepared 1:3 complexes with trivalent lanthanide ions (La, Sm, Yb, Lu and Y) and Am(III) as a representative of the trivalent actinides. In diamagnetic complexes, signals of the non-coordinating N$_8$ are observed in a small chemical shift range between 195 ppm and 206 ppm. At room temperature, the coordinating N$_8$ signals are found in a chemical shift range between 224 ppm and 275 ppm. Comparing the three diamagnetic complexes Y(III), La(III), and Lu(III), we found significant differences in $^1$H, $^{13}$C and $^{15}$N spectra. This shows that diamagnetic reference compounds for the extraction of purely paramagnetic signals of the non-coordinating N$_9$ are observed in a representative of the trivalent actinides. In diamagnetic complexes, signals of the non-coordinating N$_8$ are observed in a small chemical shift range between 195 ppm and 206 ppm. At room temperature, the coordinating N$_8$ signals are found in a chemical shift range between 224 ppm and 275 ppm. Comparing the three diamagnetic complexes Y(III), La(III), and Lu(III), we found significant differences in $^1$H, $^{13}$C and $^{15}$N spectra. This shows that diamagnetic reference compounds for the extraction of purely paramagnetic shifts have to be chosen with care. We conclude that La(III) serves as a good diamagnetic reference for the lighter part of the lanthanide series and Lu(III) and Y(III) are better suited for the heavier lanthanides.

We furthermore prepared the [Am($^{15}$N)(C5-BPP)$_3$](OTf)$_3$ complex and showed that NMR resonance signals for this complex have a stronger temperature dependence than signals of complexes with diamagnetic Ln(III), but weaker than for paramagnetic Yb(III) and Sm(III). This indicates a weak paramagnetism of the Am(III) complex, similar to earlier findings for BTP complexes.

In comparison to the diamagnetic lanthanide complexes, the coordinating N$_8$ experiences a significant upfield shift to $\sim$20 ppm, which is in excellent agreement with data from earlier studies with the Am(III)–nPrBTP complex. As comparison to the Sm(III) and Yb(III) complexes shows, this extraordinary upfield shift cannot be explained as paramagnetic effects known from studies of similar lanthanide complexes, since shifts of the coordinating N$_8$ in the same order of magnitude have only been found for the Yb(III) complex, which has a bigger magnetic moment. Explanations for this behavior are transfer of electron spin density to the nitrogen atoms by several possible mechanisms and spin–orbit coupling effects from Am(III). All transfer mechanisms rely on the existence of a Fermi contact interaction, which is mediated by covalent bonding through s-orbital containing binding orbitals.

Our results are an important contribution within current research efforts to identify the origin of selectivity of N-donor ligands in actinide–lanthanide separation. They show that NMR spectroscopy is a versatile and sensitive tool in the elucidation of fundamental bonding mechanisms especially for actinide compounds. Important insights into the metal–ligand bonding were obtained, which reveal valuable information for an optimized design of future extractants for the separation of actinides from lanthanides. Further temperature dependent NMR experiments with paramagnetic cations of the entire lanthanide series and further transuranium element cations are in progress. Moreover, we endeavor to investigate the contributions to the chemical shift using quantum chemical calculations. The obtained experimental chemical shift values for all nuclei in the complexes are important benchmarks for those calculations.

Experimental section

General

All NMR spectra were recorded at $T = 300$ K on a Bruker Avance III 400 spectrometer operating at 400.18 MHz for $^1$H, 100.63 MHz for $^{13}$C and 40.56 MHz for $^{15}$N. The spectrometer was equipped with a z-gradient broadband observe probe optimized for $x$-magnetization detection. Chemical shifts are referenced internally to TMS ($\delta$(TMS) = 0 ppm). $^{15}$N chemical shifts are referenced to $^{15}$NH$_4$Cl with $\delta$(NH$_4$Cl) = 0 ppm. For all direct excitation and correlation spectra, standard Bruker pulse sequences were used. DOSY spectra were acquired using one-shot sequences. All 1D spectra for diamagnetic complexes and Am(III) were recorded with 32k data points and are zero filled to 64k points. $^{15}$N spectra were recorded at lower spectral resolution if necessary, allowing fast pulsing and high repetition rates to compensate the paramagnetic relaxation enhancement. The reported chemical shifts are taken from 1D spectra unless stated otherwise. $^{15}$N data at natural abundances are obtained from $^1$H, $^{15}$N-HMQC spectra. Deuterated solvents were purchased from Euriso-Top GmbH. Chemicals for synthesis were purchased from VWR International and used as-is. $^{15}$N-labeled hydrazine hydrate (98% $^{15}$N) was purchased from Sigma-Aldrich and used as-is. Mass spectra using LIFDI and EI ionization methods were recorded using a JEOl JMS-700 magnetic sector instrument. Mass spectra using ESI ionization methods were recorded using a Bruker ApexQe FT-ICR instrument. All mass spectra were recorded at the mass spec facility of the Institute of Organic Chemistry at the University of Heidelberg. All mass spectra of $^{15}$N-labeled compounds were acquired using LIFDI-MS technology.$^{58,59}$ Melting points were measured using a Stuart SMP30 melting point apparatus.

TRLFS setup

All compounds for TRLFS experiments were used as received. Methanol (absolute) was purchased from Merck and stored over molecular sieves. The concentration of Cm(III) was set to $6.6 \times 10^{-8}$ mol $\cdot$ L$^{-1}$ by adding an aliquot of a stock solution [Cm(III)] = $6.7 \times 10^{-4}$ mol $\cdot$ L$^{-1}$ in HClO$_4$ ($1.0 \times 10^{-2}$ mol $\cdot$ L$^{-1}$) to the [Am($^{15}$N)(C5-BPP)$_3$](OTf)$_3$ NMR sample. The isotopic mass distribution of the Cm(III) solution was 89.7% $^{248}$Cm, 9.4% $^{246}$Cm, <0.5% $^{243}$Cm, $^{244}$Cm, $^{245}$Cm, and $^{247}$Cm, determined by alpha spectroscopy and ICP-MS. TRLFS measurements were performed using a Nd:YAG-pumped dye laser system [Surelite II laser (Continuum), NARROWscan D-R dye laser [Radiant Dyes Laser Accessories]]. For Cm(III) excitation a wavelength of 396.6 nm was used. The emission spectra were recorded at an angle of 90° to the exciting laser beam. A Shamrock 303i spectrograph
TRIFLS sample preparation

The \(^{149}\text{Am}(\text{C5-BPP})_3\)(OTf)_3 NMR sample in 600 \(\mu\)L MeOD-d_4 was transferred from a J. Young-type NMR tube into a quartz cuvette. 6 \(\mu\)L of an aqueous Cm(ClO_4)_2 stock solution (1.0 \(\times\) 10^{-5} mol L^{-1} HCIO_4, [Cm(III)] = 6.7 \times 10^{-6} mol L^{-1}) was added and carefully shaken. The change in volume was limited to 1.0% (vol). Titrations were performed by stepwise addition of a C5-BPP solution (3.0 \(\times\) 10^{-2} mol L^{-1}) in MeOD. After each addition of the ligand solution the sample was carefully shaken and a Cm(III) fluorescence spectrum was recorded.

Dimethyl 2,6-pyridinedicarboxylate

The preparation of dimethyl 2,6-pyridinedicarboxylate was carried out by a modification of a previously published method. 2,6-Dipicolinic acid (10.0 g, 59.8 mmol) and 2.0 mL sulphuric acid (conc.) were refluxed in 40 mL methanol for 3 h. After the solution was cooled to room temperature the solution was neutralized with 1.5 g (14.2 mmol) Na_2CO_3. The resulting white precipitate was separated by filtration and washed three times with 20 mL portions of cold water. The solid was dried for 2 h at 60 °C in high vacuum yielding the desired product (10.65 g, 54.6 mmol, 91%) as a white solid.

\(^{1}H\)-NMR (400.18 MHz, MeOD-d_4, \(T = 328 \text{ K}\)) keto form: \(\delta\) (ppm) = 8.19 (d, 2H, H_{3/5}), 8.08 (dd, 1H, H_4), 6.87 (s, 4H, H_8), 3.27 (s, 4H, H_{10}), 1.08 (s, 18H, H_{12}); enol form: 12% according to \(^{1}H\)-NMR \(\delta\) (ppm) = 8.27 (dd, 2H, H_{3/5}), 6.80 (s, 4H, H_8), 4.56 (s, OH), 2.59 (s, 4H, H_{10}), 1.03 (s, 18H, H_{12}).

\(^{13}C\)-NMR (100.63 MHz, MeOD-d_4, \(T = 328 \text{ K}\)) keto form: \(\delta\) (ppm) = 195.6 (C_q, C_4), 183.7 (C_q, C_2), 153.5 (C_q, C_{2/6}), 139.7 (C_s, C_8), 125.3 (C_s, C_{1/3}), 99.6 (C_s, C_6), 53.1 (C_p, C_10), 32.7 (C_p, C_{11}), 30.4 (C_p, C_12); enol form: \(\delta\) (ppm) = 195.5 (C_q, C_8), 183.2 (C_q, C_7), 136.3 (C_q, C_{2/6}), 139.9 (C_s, C_4), 126.6 (C_s, C_{3/5}), 99.6 (C_s, C_5), 56.6 (C_p, C_10), 30.7 (C_p, C_12).

HR-MS (EI) calculated for C_{21}H_{29}NO_4 [M]^+ 359.2097; found: 359.2114; calculated for C_{20}H_{28}NO_3 [M – C_H_2O]^+ 344.1862; found: 344.1841; calculated for C_{13}H_{12}NO_4 [M – C_H_4O]^+ 303.1471; found: 303.1492; calculated for C_{12}H_{11}NO_3 [M – C_H_3O]^+ 288.1236; found: 288.1275; calculated for C_{12}H_{12}O_3 [M – C_H_2O]^+ 275.1521; found: 275.1534; calculated for C_{12}H_{12}O_2 [M – C_H_2O]^+ 260.1287; found: 260.1283; calculated for C_{11}H_{11}O_3NO [M – C_H_3O]^+ 233.1416; found: 233.1416; calculated for C_{11}H_{11}NO_2 [M – C_H_3O]^+ 204.0657; calculated for C_{11}H_{11}NO 190.0868, found: 190.0488; calculated for C_{9}H_{10}O 99.0810, found: 99.0791.

2,6-Bis(5-(2,2-dimethylpropyl)-1H-pyrazol-3-yl-pyridine (C5-BPP)

8.0 mL (129 mmol) N\textsubscript{2}H\textsubscript{4}H\textsubscript{2}O (80% in H\textsubscript{2}O) were added to a solution of 540 mg (1.5 mmol) 1,1’-(pyridine-2,6-diyl)bis(4,4-dimethylhexane-1,3-dione) 40 mL methanol (abs.) and refluxed for 3 h. After the solution was cooled to room temperature the resulting white precipitate was collected and washed three times each with 30 mL water and 30 mL diethyl ether. The desired product was obtained by drying in high vacuum (0.415 g, 1.18 mmol, 79%) as a white, crystalline solid. mp: 266.5 °C.
Syntheses of lanthanide complexes

6 μmol of Ln(OTf)3 were weighed in a screw-cap glass. 18 μmol C5-BPP or [15N]C5-BPP, respectively, were dissolved in 600 μL MeOD-d4 with traces of TMS. The C5-BPP or [15N]C5-BPP ligand solution was added to the metal salt. After mixing the complex solution was transferred into an NMR tube. The sample was degassed by three freeze–pump–thaw cycles and subsequently flame-sealed. Complexes with the labeled and unlabeled ligand were prepared in the same way. The chemical shift values for the unlabeled complex are equal to those of the labeled complexes and are not stated here for brevity. However, N1 chemical shifts could only be determined from 1H,15N-gHSQC spectra and are labeled accordingly (f).

Synthesis of [243Am(C5-BPP)3]([OTf]3)

1.0 mL of a solution containing 4 mg mL−1 243Am in HNO3 (0.5 mol L−1) was transferred into a screw-cap glass. A total of 280 μL NaOH (2.0 mol L−1) was added in portions, resulting in precipitation of americium hydroxide. After 20 μL of additional NaOH (2.0 mol L−1) were added, the solution was centrifuged at 6000 rpm for 3 min. Additional 10 μL NaOH solution (2.0 mol L−1) were added, the solution was centrifuged again (6000 rpm, 2 min) and the supernatant was removed. Following this procedure, the precipitate was washed three times with 1.0 mL portions of NaOH (0.01 mol L−1) and once with 1.0 mL water. The americium hydroxide was dissolved in 1.0 mL H2O and 10 μL trifluoromethanesulfonic acid, forming Am(OTf)3. For complexation of Am(OTf)3 with C5-BPP or [15N]C5-BPP, respectively, 420 μL Am(OTf)3 solution were heated to dryness at about 100 °C on a heating plate. The obtained pale-pink solid was subsequently washed with 250 μL D2O and heated to dryness. The ligand solution (18 μmol in 600 μL MeOD-d4) was added to the Am(OTf)3, carefully mixed and transferred into a J. Young-type NMR tube.

[Y([15N]C5-BPP)3]([OTf]3)

1H-NMR (400.18 MHz, MeOD-d4): δ (ppm) = 8.16 (t, 1H, H4, J = 7.9 Hz), 7.96 (d, 2H, H3/5, J = 7.9 Hz), 6.77 (s, 2H, H11), 2.41 (s, 4H, H12), 0.70 (s, 18H, H14).

13C-NMR (100.63 MHz, MeOD-d4): δ (ppm) = 154.3 (Cq, C7), 151.6 (Cq, C2, C6), 148.3 (Cq, C10), 142.9 (Cq, C4), 123.7 (Cq, C3, C5), 106.4 (Cq, C11), 39.5 (Cq, C12), 32.2 (Cq, C13), 29.7 (Cp, C14).

15N-NMR (40.56 MHz, MeOD-d4): δ (ppm) = 287 (N4)∗, 206 (N4)∗. Value taken from an 1H,15N-HMQC spectrum.


Synthesis of [151Am(C5-BPP)3]([OTf]3)

1H-NMR (400.18 MHz, MeOD-d4): δ (ppm) = 8.16 (t, 1H, H4, J = 7.9 Hz), 7.96 (d, 2H, H3/5, J = 7.9 Hz), 6.77 (s, 2H, H11), 2.41 (s, 4H, H12), 0.70 (s, 18H, H14).

13C-NMR (100.63 MHz, MeOD-d4): δ (ppm) = 154.3 (Cq, C7), 151.6 (Cq, C2, C6), 148.3 (Cq, C10), 142.9 (Cq, C4), 123.7 (Cq, C3, C5), 106.4 (Cq, C11), 39.5 (Cq, C12), 32.2 (Cq, C13), 29.7 (Cp, C14).

15N-NMR (40.56 MHz, MeOD-d4): δ (ppm) = 287 (N4)∗, 206 (N4)∗. Value taken from an 1H,15N-HMQC spectrum.


ESI-MS (CH3OH) calculated for C63H85N15La [La(C5-BPP)3]([OTf]3): 1440.5367, found: 1440.5361; calculated for C64H88F6N15O6S2La [La(C5-BPP)3]([OTf]3) + C2H5OH + HF+ + Na+: 1441.5922, found: 1441.5880; calculated for C64H88F6N15O6S2La [La(C5-BPP)3]([OTf]3) + Na+: 1442.6404, found: 1442.6491; calculated for C64H88F6N15O6S2La [La(C5-BPP)3]([OTf]3): 1443.6404, found: 1443.6526.

[La([15N]C5-BPP)3]([OTf]3)

1H-NMR (400.18 MHz, MeOD-d4): δ (ppm) = 8.16 (t, 1H, H4, J = 7.9 Hz), 7.96 (d, 2H, H3/5, J = 7.9 Hz), 6.77 (s, 2H, H11), 2.41 (s, 4H, H12), 0.70 (s, 18H, H14).

13C-NMR (100.63 MHz, MeOD-d4): δ (ppm) = 154.3 (Cq, C7), 151.6 (Cq, C2, C6), 148.3 (Cq, C10), 142.9 (Cq, C4), 123.7 (Cq, C3, C5), 106.4 (Cq, C11), 39.5 (Cq, C12), 32.2 (Cq, C13), 29.7 (Cp, C14).

15N-NMR (40.56 MHz, MeOD-d4): δ (ppm) = 272 (d, N4, J = 9.9 Hz), 206 (m, N4)∗.
for C$_{42}$H$_{58}$Na$_{10}$Na [2C$_{5}$-BPP + Na$^{+}$]: 725.4744, found: 725.4803; calculated for C$_{62}$H$_{80}$F$_{3}$N$_{10}$O$_{3}$SNa [2C$_{5}$-BPP] + Na$^{+}$: 1274.6055, found: 1274.6056; calculated for C$_{42}$H$_{58}$Na$_{10}$Na [2C$_{5}$-BPP + Na$^{+}$]: 874.4260, found: 874.4262; calculated for C$_{42}$H$_{58}$Na$_{10}$Na [2C$_{5}$-BPP + Na$^{+}$]: 725.4744, found: 725.4754; calculated for C$_{62}$H$_{80}$F$_{3}$N$_{10}$O$_{3}$SNa [2C$_{5}$-BPP + Na$^{+}$]: 374.2320, found: 374.2321.

[Lu$^{3+}$C$_{5}$-BPP$_{3}$]$_{2}$[OTf]$_{3}$

1H-NMR (400.18 MHz, MeOD-d$_4$): $\delta$ (ppm) = 2.29 (d, $\frac{J}{C_{16}, C_{17}}$, 14.0 Hz), 2.55 (d, $\frac{J}{C_{16}, C_{17}}$, 14.0 Hz), 0.71 (s, 18H, H$_{18}$).

13C-NMR (100.63 MHz, MeOD-d$_4$): $\delta$ (ppm) = 153.6 (C$_q$, C$_7$), 149.9 (C$_q$, C$_{15}$), 149.5 (C$_q$, C$_{16}$), 142.9 (C$_q$, C$_4$), 123.2 (C$_q$, C$_3$), 123.5 (C$_q$, C$_6$), 106.2 (C$_q$, C$_{11}$), 39.6 (C$_d$, C$_{12}$), 32.2 (C$_d$, C$_{13}$), 29.7 (C$_p$, C$_{14}$).

15N-NMR (40.56 MHz, MeOD-d$_4$): $\delta$ (ppm) = 267 (N$_1$), 265 (N$_6$, N$_9$) = 9.4 Hz), 205 (m, N$_5$).

[Eu$^{3+}$C$_{5}$-BPP$_{3}$]$_{2}$[OTf]$_{3}$

1H-NMR (400.18 MHz, MeOD-d$_4$): $\delta$ (ppm) = 3.90 (m, H$_{13}$, 3J = 7.9 Hz), 3.15 (m, H$_{12}$, 2J = 7.9 Hz), 6.77 (s, 2H, H$_{14}$), 2.64 (d, 2H, H$_{13}$, 3J = 13.9 Hz), 0.81 (s, 18H, H$_{18}$).

13C-NMR (100.63 MHz, MeOD-d$_4$): $\delta$ (ppm) = 150.4 (C$_q$, C$_7$), 153.4 (C$_q$, C$_{15}$), 153.4 (C$_q$, C$_{16}$), 143.4 (C$_q$, C$_4$), 122.9 (C$_q$, C$_3$), 106.2 (C$_q$, C$_{11}$), 39.5 (C$_d$, C$_{12}$), 32.3 (C$_d$, C$_{13}$), 29.8 (C$_p$, C$_{14}$).

15N-NMR (40.56 MHz, MeOD-d$_4$): $\delta$ (ppm) = 221 (s, N$_1$), 224 (s, N$_6$, N$_9$), 205 (m, N$_5$).

[Sm$^{3+}$C$_{5}$-BPP$_{3}$]$_{2}$[OTf]$_{3}$

1H-NMR (400.18 MHz, MeOD-d$_4$): $\delta$ (ppm) = 7.69 (d, 2H, H$_{13}$, $\frac{J}{C_{16}, C_{17}}$ = 7.9 Hz), 7.47 (t, 1H, H$_4$, $\frac{J}{C_{16}, C_{17}}$ = 7.9 Hz), 6.58 (s, 2H, H$_{14}$), 2.91 (d, 2H, H$_{12}$, $\frac{J}{C_{16}, C_{17}}$ = 13.9 Hz), 2.55 (d, 2H, H$_{12}$, $\frac{J}{C_{16}, C_{17}}$ = 13.9 Hz), 0.64 (s, 18H, H$_{18}$).

13C-NMR (100.63 MHz, MeOD-d$_4$): $\delta$ (ppm) = 164.8 (C$_q$, C$_7$), 164.2 (C$_q$, C$_{15}$), 147.9 (C$_q$, C$_{16}$), 147.7 (C$_q$, C$_4$), 116.3 (C$_q$, C$_3$), 101.7 (C$_q$, C$_{11}$), 38.7 (C$_d$, C$_{12}$), 33.9 (C$_d$, C$_{13}$), 29.6 (C$_p$, C$_{14}$).

15N-NMR (40.56 MHz, MeOD-d$_4$): $\delta$ (ppm) = 216 (N$_1$), $\frac{1}{2}$ (N$_6$), $-22$ (N$_9$) * Value taken from an $^{1}H$,$^{15}N$-HMOC spectrum.

19F-NMR (376.54 MHz, MeOD-d$_4$): $\delta$ (ppm) = -80.00 (s, CF$_3$SO$_3^{-}$).

[Am$^{3+}$C$_{5}$-BPP$_{3}$]$_{2}$[OTf]$_{3}$

1H-NMR (400.18 MHz, MeOD-d$_4$): $\delta$ (ppm) = 7.64 (d, 2H, H$_{13}$, $\frac{J}{C_{16}, C_{17}}$ = 7.9 Hz), 7.47 (t, 1H, H$_4$, $\frac{J}{C_{16}, C_{17}}$ = 7.9 Hz), 6.58 (s, 2H, H$_{14}$), 2.91 (d, 2H, H$_{12}$, $\frac{J}{C_{16}, C_{17}}$ = 13.9 Hz), 2.55 (d, 2H, H$_{12}$, $\frac{J}{C_{16}, C_{17}}$ = 13.9 Hz), 0.64 (s, 18H, H$_{18}$).

13C-NMR (100.63 MHz, MeOD-d$_4$): $\delta$ (ppm) = 164.8 (C$_q$, C$_7$), 164.2 (C$_q$, C$_{15}$), 147.9 (C$_q$, C$_{16}$), 147.7 (C$_q$, C$_4$), 116.3 (C$_q$, C$_3$), 101.7 (C$_q$, C$_{11}$), 38.7 (C$_d$, C$_{12}$), 33.9 (C$_d$, C$_{13}$), 29.6 (C$_p$, C$_{14}$).

15N-NMR (40.56 MHz, MeOD-d$_4$): $\delta$ (ppm) = 217 (s, N$_1$), 21 (N$_6$), $-22$ (N$_9$)* Value taken from an $^{1}H$,$^{15}N$-HMOC spectrum.

19F-NMR (376.54 MHz, MeOD-d$_4$): $\delta$ (ppm) = -80.00 (s, CF$_3$SO$_3^{-}$).

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Notes and references