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## How can f-Block Mono-Cations Behave as Mono-Cations of d-Block Transition **Metals?**

# Alexandre Quemet,<sup>[a]</sup> Rene Brennetot,<sup>[a]</sup> Jean-Yves Salpin,<sup>[b,c]</sup> Alvaro Cimas,<sup>[d]</sup> Colin Marsden,<sup>[e,f]</sup> Jeanine Tortajada,<sup>[b,c]</sup> and Pierre Vitorge<sup>[c,g]</sup>

Keywords: Density functional calculations / Ion-molecule reactions / Lanthanides / Actinides / Multiple bonds

The electronic structures of LnNH<sup>+</sup> is studied by DFT (B3LYP) quantum calculation for the Ln = La, Eu and Gd 4f-block elements (lanthanides).  $Ln \equiv N$  triple bonds of essentially d-character are formed for La and Gd explaining why La<sup>+</sup> and Gd<sup>+</sup> behave like dblock elements as experimentally evidenced by mass spectrometry, and why the Ln<sup>+</sup> reactivity is correlated with its electron-promotion energy: the present theoretical study is a support to such correlation and qualitative knowledge. The  $Ln^+ + NH_3 \rightarrow LnNH_3^+ \rightarrow transition$ state  $\rightarrow$  HLn=NH<sub>2</sub><sup>+</sup>  $\rightarrow$  transition state  $\rightarrow$  Ln  $\equiv$  NH<sup>+</sup> + H<sub>2</sub> reaction pathway is calculated.

The formation of HLn=NH<sub>2</sub><sup>+</sup> corresponds to the formation of new covalent bonds associated with more electron pairing, and corresponding lowering of the spin multiplicity -spin crossing reaction. It is in this step that low electron-promotion energy is required to promote an Ln 4f electron onto an Ln 5d orbital as typically for La<sup>+</sup> and Gd<sup>+</sup>. Similar geometry, bonding and electronic cofiguration are calculated for NpNH<sup>+</sup> -an actinide complex observed by mass spectrometry- with higher participation of 5f-valence orbitals (20% and 25% for the  $\sigma$  and  $\pi$  bonds) as compared to the 4f-valence orbitals (3% and 8%) of GdNH<sup>+</sup>: Gd<sup>+</sup> and Np<sup>+</sup> are the only lanthanide and actinide cations with two non-f-valence electrons -one s and one d- in their ground states.

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#### Introduction

Very different chemical behaviours have been evidenced in mass spectrometry analysis across the lanthanide (Ln) series of f-block elements:<sup>[1]</sup> they can even behave as the Sc and Y d-block transition elements.<sup>[2]</sup> It has been proposed that the reactivity of Ln<sup>+</sup> with gas in collision-reaction cells of mass spectrometers is correlated with its electron-promotion energy, the energy to obtain two non-f-valence electrons: the  $4f^{n}5d^{1}6s^{1}$  electronic configuration. This is in contrast with the well established picture for the chemistry of the f-block elements in condensed phases.

Most of the rare earths are lanthanides. La, the first lanthanide (Ln) in the periodic table is usually stable in the form of the  $La^{3+}$ hard cation, where La(III) is in its group oxidation state. The heavier Ln elements are f-block elements corresponding to the progressive filling of the 4f-shell, with a few exceptions. The 4fvalence orbitals of Ln<sup>3+</sup> do not easily participate to covalent bonding, consistently with the hard character of Ln<sup>3+.[3]</sup> This accounts for the chemical analogy among the  ${\rm Ln}^{^{3+}}$  series -despite they have different electronic configurations, and with other hard

trications of similar sizes as typically the (An<sup>3+</sup>) tri-cations of the actinides, the series of the 5f-block elements.

This behaviour is in contrast with those of d-block transition elements, which ions often are rather soft -giving covalent bonds: their chemical behaviours are driven by their electronic configurations, which vary across the d-transition metal series. Nevertheless, another rare earth, Y, the transition element above La in the third column of the periodic table is a chemical analogue of La.  $Y^{3+}$  is of the size of a heavy  $Ln^{3+}$  as a result of the Ln contraction: it behaves as a heavy  $Ln^{3+}$ , while  $Sc^{3+}$  -Sc is the transition element above Y- is clearly smaller.<sup>[4]</sup>

Sc<sup>+</sup>, Y<sup>+</sup> or Ln<sup>+</sup> can be viewed as reduced Sc<sup>3+</sup>, Y<sup>3+</sup> or Ln<sup>3+</sup> stable hard ions, on which two electrons have been added:  $M^{3+}$  +  $2e^- \rightarrow M^+$ . These two added electrons are not expected to be firmly retained in the monocations, since rare earths are usually more stable in the chemical forms of trications: meaning that these monocations are not especially expected to be of hard character in contrast with the stable trications. Namely, Ln<sup>+</sup> chemistry is not especially expected to be similar to that of the  $Sc^+$  or  $Y^+$  d-cation, when the two Ln<sup>+</sup> extra electrons are in f-orbitals. Conversely, some analogy is not unexpected for those Ln<sup>+</sup> ions of similar electronic configurations as Sc<sup>+</sup> and Y<sup>+</sup>, namely when the promotion energy the 4f<sup>n</sup>5d<sup>1</sup>6s<sup>1</sup> electronic configuration is small. Note that d-valence orbitals can more easily be stabilized than the f-ones on covalent bonding, since the d-orbitals usually have greater spatial expansions, which facilitates covalency: electronic configuration can change on covalent bonding.

Experimental mass spectrometry showed the formation of  $MNH^+$ , the key product for the reactivities of d-elements ( $M^+ = Sc^+$ and Y<sup>+</sup>), and for Ln<sup>+</sup> ions of small (La<sup>+</sup>) or zero (Gd<sup>+</sup>) electronpromotion energies (see supporting information):[1a, 5]

> $Ln^+ + NH_3 \rightarrow LnNH^+ + H_2$ (1)

where the  $Ln^+$  strong reducer produces  $H_2$  by inserting into N-H bonds. The LnNH<sup>+</sup> product can be described with usual formal atomic oxidation states: Ln(III), N(-III) and H(I) corresponding to their column in the periodic table; however, this is not enough to explain the stability of LnNH<sup>+</sup>, since Reaction 1

was experimentally observed only for those f-block mono-cations with low electron-promotion energies.

Herein, we report computational studies of the electronic structure of LnNH<sup>+</sup> for Ln = La, Eu and Gd. Eu<sup>+</sup> and Gd<sup>+</sup> have different reactivities: <sup>[1a, 5]</sup> the electronic structure of LnNH<sup>+</sup> turned out to be that of a d-element only for those lanthanides (La and Gd) for which Reaction 1 was experimentally observed, while such an electronic structure is not possible for Eu<sup>+</sup>, a mono-cation of low reactivity. <sup>[1a, 5]</sup> Actinide mono-cations (An<sup>+</sup>) are also compared. More details on mass spectrometry results, reaction mechanism (Figure.1) and potential energy surfaces will be given elsewhere together with higher level of calculations: MP2, CCSD(T), CASPT2 confirming that DFT/B3LYP is enough for the calculations presented here.<sup>[5]</sup>

#### **Results and Discussion**

LnNH<sup>+</sup> is linear with Ln-N bond lengths of 1.88 Å (La) and 1.83 Å (Gd) (Table 1). These bond lengths are not unexpected as compared with published Sc-N (1.71 Å) and Y-N (1.85 Å) distances<sup>[6]</sup>. Ma et al.<sup>[7]</sup> found a Ce-N bond length (1.92 Å) longer than the La-N one by 0.04 Å: virtually the same distance is indeed expected, since Ce is just after La in the Ln series. The Ln-N bond length is the shortest in LnNH<sup>+</sup> as compared in all the intermediate species we calculated.<sup>[5]</sup> We calculated that the LaNH<sup>+</sup> geometry is more stable by 400 kJ.mol<sup>-1</sup> than the HLaN<sup>+</sup> one.

The linear geometry of LnNH+ suggests the sp hybridization for N, thus associated with a Ln $\equiv$ N triple bond. Indeed, the NBO5.9 population analysis finds the (La $\equiv$ N-H)<sup>+</sup> Lewis structure; namely four covalent bonds (Table 2).

The participation of the Ln 4f orbitals is small: 16% (La) or 8% (Gd). As expected, the  $\sigma$ -valence orbitals are lower in energy than the  $\pi$ -ones -which are the HOMOs, and the less nodal planes the lower is the energy among the  $\sigma$ -valence orbitals (Figure.2 and Table 2). Hui-Zhen et al.<sup>[6b]</sup> also reported a linear structure for YNH<sup>+</sup> with a Y=N triple bond formed by Y s- and d-orbitals, and N s- and p-orbitals. According to the octet rule, the maximum possible number of covalent bonds is here formed in Lewis interpretations, since the NH fragment cannot form more than 3 covalent bonds corresponding to the sp hybridization.

To check the influence of f-orbitals, we also suppressed the ffunctions from the basis set, and we re-optimized the geometry of LaNH<sup>+</sup>. NBO5.9 still found the (La=N-H)<sup>+</sup> Lewis structure, despite the La-N distance is larger by 0.10 Å, and the complex is destabilized by 115 kJ.mol<sup>-1</sup>. This compares well with similar calculations across the ThO<sub>2</sub>, PaO<sub>2</sub><sup>+</sup>, UO<sub>2</sub><sup>2+</sup> seemingly isoelectronic series:<sup>[8]</sup> adding f-functions decreased the Th=O bond length by 0.12 Å, and stabilized the complex by 376 kJ.mol<sup>-1</sup> (138 kJ.mol<sup>-1</sup> per Th≡O bond), while the influence is more important for Pa and U. After others, this was an argument to indicate that Th(IV) is rather a d-element in ThO<sub>2</sub>, while the isoelectronic transthorium elements are true f-block elements in the corresponding molecular ions. The effects of the f-orbitals appear to be of the same order of magnitude: slightly (20%) smaller for both energy and geometry for the La=N triple bond than for the Th=O one confirming that the importance of f-type functions is moderate for LaNH<sup>+</sup>. In LaNH<sup>+</sup> and GdNH<sup>+</sup>, the N 2s 2p orbitals, and the Ln 5d orbitals participate to the triple bonds. Ln-N bonding has a predominant d-character. In that respect, these (La<sup>+</sup> and Gd<sup>+</sup>) so called 4f-block monocations, actually have chemical behaviours of d-block elements: they form covalent bonds, which all have a dominating d-character. Furthermore, the M atomic charges (M = Y, La and Gd) are

comparable in MNH<sup>+</sup> for La (1.97), Gd (1.92) (Table 1), and as published for Y  $(1.94)^{[6b]}$  consistently with similar participation of the M electrons in the M-N bond. Note that Y is a d-block element.

 $Np^+$  is the only actinide mono-cation with zero electron-promotion energy as the Gd^+ lanthanide mono-cation:  $^{[1h]}$  Gd^+ and  $Np^+$  both have one s- and one d-valence electrons.  $Np^+$  have the same reactivity as Gd^+ both forming MNH^+ (M = Gd and Np, see supporting information). We calculated that the Np-N distance (1.88 Å) in NpNH<sup>+</sup> is a little longer than the Gd-N one in GdNH^+ (1.83 Å), in line with the small expansion usually observed from Ln to An ions.<sup>[9]</sup> The participation of Np 5f-orbitals (40% in the  $\sigma$  Np-N bond, and 20% in the two  $\pi$  ones) is greater as compared to that of Gd 4f orbitals (3% in the  $\sigma$  bond and 8% in the  $\pi$  ones), in line with the less compact geometry of the An 5f-orbitals as compared to the 4f-ones.<sup>[10]</sup>

Several intermediate species make the connection between the  $(La^+ + NH_{3,})$  reactants and the  $(LaNH^+)$  final product.<sup>[5]</sup> The first intermediate species is LaNH<sub>3</sub><sup>+</sup>. Its optimized geometry is  $C_{3v}$  tetrahedral. The La-N distance is larger in LaNH<sub>3</sub><sup>+</sup> (2.61 Å) as compared in LaNH<sup>+</sup> (1.88 Å). This corresponds to weaker La-N interaction in LaNH<sub>3</sub><sup>+</sup> as also reflected by the La-N  $\sigma$ -bond vibrational frequency (283cm<sup>-1</sup> for LaNH<sub>3</sub><sup>+</sup> as compared to 838 cm<sup>-1</sup> for LaNH<sup>+</sup>). This suggests purely electrostatic La-N interaction, or at most single La-N covalent bond. In both cases it corresponds to the sp<sup>3</sup> hybridization for N in LaNH<sub>3</sub><sup>+</sup>.

This is in line with the charge of La (0.92): virtually no change in the charge of La as compared to initial La<sup>+</sup>. The two electrons initially described as the N 2p<sub>z</sub> lone pair in  $|NH_3|$  are now (in La-NH<sub>3</sub><sup>+</sup>) shared with La on its 5d<sub>z<sup>2</sup></sub> orbital, formally corresponding to donation from one of the 2p<sub>z</sub> N electrons to the 5d<sub>z<sup>2</sup></sub> La orbital (see the scheme for Reaction  $La^+ + NH_3 \rightarrow LaNH_3^+$  in Figure.1 and Scheme 1).

The rearrangement of LaNH<sub>3</sub><sup>+</sup> into the HLaNH<sub>2</sub><sup>+</sup> intermediate species is an H transfer from NH<sub>3</sub> to La breaking an (N-H) covalent bond of LaNH<sub>3</sub><sup>+</sup>, and making new covalent bonds in HLaNH<sub>2</sub><sup>+</sup> (La=N and La-H). This results in more covalent bonds, and more electron pairing, which decreases the spin multiplicity. La must have the 4f<sup>0</sup>5d<sup>1</sup>6s<sup>1</sup> electronic configuration to make the new La=N covalent bond in HLaNH<sub>2</sub><sup>+</sup> (see the scheme for Reaction LaNH<sub>3</sub><sup>+</sup>  $\rightarrow$  HLaNH<sub>2</sub><sup>+</sup> in Figure.1 and Scheme 2). It is in this step that low promotion energy is needed.

HLaNH<sub>2</sub><sup>+</sup> have a planar structure (in line with sp<sup>2</sup> hybridization for N, and La=N double bond) and five bonds: one  $\sigma$  Ln-H bond, two  $\sigma$  N-H bonds (one for each of the two H atoms bonded to N), one  $\sigma$  Ln-N bond (La 6s 5p 5d / N 2s 2p) and one  $\pi$  Ln-N bond (La 5d / N 2p). The 4f orbitals slightly participate (16% for La and 4% for Gd) to the  $\sigma$  Ln-N bonds.

LaNH<sup>+</sup> can further be formed -from HLaNH<sub>2</sub><sup>+</sup>- via La(H<sub>2</sub>)NH<sup>+</sup>, a four-center transition state (TS2), where H<sub>2</sub> is weakly bonded to LaNH<sup>+</sup> by both its La and N atoms (Figure.1). The charges of H<sub>N</sub> and H<sub>La</sub> are +0.5 and -0.5 respectively. They further attract each other to finally form H<sub>2</sub>: LaNH<sup>+</sup> is formed with H<sub>2</sub> loss.

Eu is just before Gd in the periodic table; for this reason, differences in their chemical behaviours cannot essentially be originated in the sizes of their ions when at the same oxidation state: it should rather be originated in differences in their electronic configurations. Eu<sup>+</sup> has the highest electron-promotion energy (388 kJ.mol<sup>-1</sup>) among the Ln<sup>+</sup> cations from the 4f<sup>7</sup> 6s<sup>1</sup> to the 4f<sup>6</sup> 5d<sup>1</sup> 6s<sup>1</sup> electronic configuration. This high value is not unexpected: in the ground state of Eu<sup>+</sup> the 4f sub-layer is half occupied conferring a great stability to this fundamental electronic configuration (4f<sup>7</sup> 6s<sup>1</sup>). In the EuNH<sub>3</sub><sup>+</sup> initial complex Eu has only one non-f-electron, a 6s electron, which would pair with an H 1s electron giving the Eu-H

covalent bond if  $\text{HEuNH}_2^+$  would form, but one electron would remain unpaired on N, since the 4f to 5d promotion energy is too high to allow the formation of a second Eu-N bond (see the scheme for Reaction  $\text{Eu}NH_3^+ \rightarrow HEuNH_2^+$  in Figure.1 and Scheme 3) consistently with the NBO population analysis that gives one  $\sigma$  Eu-N bond in  $\text{HEuNH}_2^+$ , as also found for Eu-NH<sub>3</sub><sup>+</sup>.

 $\rm HEuNH_2^+$  is calculated to have a non planar structure indicating that N is not in the sp<sup>2</sup> hybridization: Ln-N cannot be a double bond. Finally, all this suggests that the  $\rm HEuNH_2^+$  geometry is not much stable because the electronic configuration of Eu does not enough facilitate covalency for the Eu-N bond in  $\rm HEuNH_2^+$  It is further confirmed in the EuNH<sup>+</sup> species where the Eu=N bond is only a double bond. This confirms that the formation of  $\rm HLnNH_2^+$ is the step, where the electron-promotion energy plays a key role.

Am is the An with the same electronic configuration as Eu. They each have high electron-promotion energy, and the f-sublayer is half occupied conferring a great stability to this fundamental electronic configuration. We calculated that the Am-N distance (2.16 Å) in AmNH<sup>+</sup> is the same for Eu-N in EuNH<sup>+</sup> (2.16 Å). They have similar electronic structures with double An=N bonds, in line with their similar reactivities: AnNH<sup>+</sup> is not formed for An = Eu and Am species (see supporting information).

Scheme 1



Scheme 2



Scheme 3



Figure 1. Scheme of reaction  $Ln^+ + NH_3 \rightarrow LnNH^+ + H_2$  for  $La^+/Gd^+$  (a) and  $Eu^+$  (b). Transition states are in []. The dot (•) represents a valence electron. The (actually f-unpaired) electrons that do not participate to the bonding are not represented. The limiting step for the reaction of Eu is represented by the red cross: the spin crossing (SC) reaction is the key steps that requires low electron-promotion energy. Dotted arrows are for reactions that are not observed





Figure.2. Molecular valence orbitals of LaNH<sup>+</sup>

Table 1. Electronic configuration and electron-promotion energy  $(PE)^{[1h]}$  in kJ.mol^1 of M<sup>+</sup>. M-N bond length (in Å), M NBO5.9 charge and  $\nu_{\sigma}$  M–N harmonic vibrational frequency (in cm<sup>-1</sup>) in MNH<sup>+</sup>

$\mathbf{M}^+$	Elec. Conf.	PE	d(M-N)	M charge	$\nu_{\sigma}$ M-N.	[2
$La^+$	$4f^05d^2$	19	1.88	1.98	838	
$\mathrm{Eu}^+$	$4f^76s^1$	388	2.16	1.83	575	
$\mathbf{Gd}^+$	$4f^75d^16s^1$	0	1.83	1.92	850	
$Np^+$	$5f^46d^17s^1$	0	1.88	1.93	862	
$Am^+$	$5f^77s^1$	245	2.16	1,70	574	[3

Table 2. Molecular orbitals of of LaNH $^{\scriptscriptstyle +},$  energy (E) is in eV. LaNH $^{\scriptscriptstyle +}$  is linear along the z axis.

Е	Main character	Sym.	La	N	Н	[5]
-23.0	$\sigmaN\text{-}H$	A1	6s 5d <sub>z<sup>2</sup></sub>	2s 2pz	1s	
-20.4	σ La-N	A1	$5dz^2 4fz(5z^2-3r^2)$	2s 2pz	1s	
-11.5	$\pi$ La-N	E1	5dxy 4fx(5z <sup>2</sup> -3r <sup>2</sup> )	2px		[6]
-11.5	π La-N	E1	5dyz 4fy(5z <sup>2</sup> -r <sup>2</sup> )	2py		

#### Conclusions

In conclusion, the original MOs used for the quantum calculations evidence covalent bonding with dominant d-character for the contributions of Ln = La and Gd in compounds along their reaction pathway. This is confirmed by NBO5.9 population analysis. Ln<sup>3+</sup> hard trications can be viewed as La<sup>3+</sup> -which does not have any valence electron- where the extra electrons are usually added onto 4f-orbitals that do not participate to the bonding. Double reduction of some Ln<sup>3+</sup> ions into Ln<sup>+</sup> -typically La<sup>+</sup> and Gd<sup>+</sup>- can bring one electron in an s-orbital, and another one in a d-orbital participating to covalent bonds as for the Sc<sup>+</sup> and Y<sup>+</sup> d-block ions of transition elements: Sc<sup>+</sup> and Y<sup>+</sup> are isoelectronic to La<sup>+</sup> in M=NH<sup>+</sup>. The same is here calculated for Np<sup>+</sup>, an An<sup>+</sup> 5-f block cation with two non-f electrons in its ground state (5f<sup>4</sup>6d<sup>1</sup>7s<sup>1</sup>).

#### **Experimental Section**

Computational and experimental details are given in the Supporting Information

Supporting Information (see footnote on the first page of this article)

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 $M \equiv N$  triple bonds of essentially dcharacter are formed for lanthanide (M = La and Gd) and actinide (M = Np) fblock elements explaining why La<sup>+</sup>, Gd<sup>+</sup> and Np<sup>+</sup> behave like d-mono-cations as experimentally evidenced by mass spectrometry, and why their reactivities are correlated with their electronpromotion energies. Triple M≡N bond

Quemet A., Brennetot R., Salpin J.Y., Cimas A., Tortajada J., Marsden C, and Vitorge P...... Page No. – PageNo.

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### Alexandre Quemet, Rene Brennetot, Jean-Yves Salpin, Alvaro Cimas, Colin Marsden, Jeanine Tortajada, and Pierre Vitorge\*

**Computational methods**: The quantum calculations were performed with the Gaussian03 suite of programs.<sup>[1]</sup> The geometries and the natural population analysis of the stationary points on the potential energy profiles have been obtained with DFT/B3LYP calculations, this level of calculation compares well with MP2, CCSD(T) and CASPT2 (as described in the companion paper: ref.5 of the communication) for the data presented here.

The Stuttgart's small core relativistic effective core potentials ECP28MWB with 28 core electrons were used for the Ln atoms.<sup>[2]</sup> We call STUTf, the ECP28MWB\_SEG associated basis where the g-functions have been suppressed. They were used for the (B3LYP) geometry optimizations, and to extract the electronic structures. 2 g-functions were added for the MP2 and CCSD(T) calculations (see ref.5 of the communication). The 6-31+G(d,p) basis set was used for N and H. Similarly, we note STUT no f no g the STUTf basis set from which the f-functions are suppressed.

Electronic configuration, spin contamination, energy difference between the highest occupied (HUMO) and the lowest unoccupied (LUMO) molecular orbitals, vibrational frequencies, and other usual calculation results were examined to check the quantum calculations. Natural population analysis (NPA) was performed with software NBO5.9,<sup>[3]</sup> which also provides atomic charges. However, we are essentially discussing the mono- (or bi- for close shell calculations) electronic Molecular Orbitals (MO) used and produced by the B3LYP/STUTf quantum calculations called MOs when nothing else is stated. The molecular graphic package Visual Molecular Dynamics<sup>[4]</sup> was used for visualizing geometries, and molecular orbitals with an isosurface value of 0.01 atomic unit from Gaussian03 cube files.

**Experimental methods:** Mass spectrometry measurements were carried out with a single quadrupole ICP-MS X Series I and Serie II (Thermo-Fisher Scientific, Bremen, Germany) equipped with the Collision Cell Technology (CCT). Main operating parameters are resumed in Table S1.The ICP-MS X serie I has been modified in order to work with radioactive materials. A glove box (JACOMEX, Dagneux, France) has been installed for this purpose. The nebulizer, the peristaltic pump, the plasma ion source, and the interface are located inside the glove box. An autosampler CETAX ASX 260 also situated inside the glove box, and controlled by the PlasmaLab software (Thermo Electron) minimizes human irradiation. Air tightness is ensured by a modified interface, and the pumping system was also adapted. A new pump was added (EDWARDS 65m3 h–1) from the standard configuration to ensure a better vacuum in the expansion, and to compensate the head losses due to the addition of filters.

Table S1. Typical ICP-MS operating conditions

	Q ICP-MS X serie I (Thermo Fischer)	Q ICP-MS X serie II (Thermo Fischer)	
RF power	1400 W	1250 W	
Plasma gas flow rate	14 L.min <sup>-1</sup>	14 L.min <sup>-1</sup>	
Auxiliary gas flow rate	0.8 L.min <sup>-1</sup>	0.75 L.min <sup>-1</sup>	
Nebulizer gas flow rate	0.85 L.min <sup>-1</sup>	0.84 L.min <sup>-1</sup>	
Sample uptake rate	0.4 mL.min <sup>-1</sup>	1 mL.min <sup>-1</sup>	
Nebulizer type	Quartz concentric	Quartz concentric	
Spray Chamber type	Conical with impact bead	Quartz chamber with impact bead	

Many elements can be analyzed in standard mode, i.e. without gas in the cell. Nevertheless, interferences can complicate the analysis, especially for nuclear fuels, where elements are not with their natural isotopic compositions. Collision/reaction devices are currently used to get rid of such interferences. This system is based on addition of collision and/or reaction gas after cation formation. The gas impacts or reacts with ion beam eliminating isobaric or poly-atomic interference. The ion lens setting, the nebulizer gas-flow rate and the torch position of the instrument were optimized every day with a multi element tuning solution (SPEX) in standard mode. This solution contains  $1 \mu g.L^{-1}$  of uranium and indium. It is used to maximize sensitivity, and to perform short term stability tests. Typical sensitivities were better than 150,000 counts.s<sup>-1</sup>.ug<sup>-1</sup>.L (X serie II) and 300,000 counts.s<sup>-1</sup>.ug<sup>-1</sup>.L (X serie

I) with a residual standard deviation better than 1 % for 10 measurements of 1 minute. In gas mode, a 1  $\mu$ g.L<sup>-1</sup> indium solution -which does not react with gas- was used to maximize sensitivity for each gas flow rate.

 $10 \ \mu g.L^{-1}$  Ln mono-elemental solutions were prepared from 1000 mg.L<sup>-1</sup> La and Eu standard SPEX, and <sup>160</sup>Gd, <sup>237</sup>Np and <sup>241</sup>Am mono-isotopic solutions diluted in a 0.5 N nitric acid solution. This 0.5 N nitric acid solution was prepared by diluting a sub-boiling 14.3N nitric acid solution with deionized water (resistivity 18.2 MΩ cm). The sub-boiling nitric acid solution was obtained by distillation of a 65% HNO<sub>3</sub> Normatom solution using an EVAPOCLEAN system (Analab, France). Indium has been added in each sample to follow the scattering losses consecutive to the introduction of a gas in the Q ICP-MS reaction cell. Ammonia (minimum purity of 99.98%) was obtained from MESSER (Puteaux, France). High purity was used to avoid secondary reactions with impurities. Nevertheless, deuterated ammonia (ND<sub>3</sub>) – D isotopic content of 99.75% - was used to check results obtained with NH<sub>3</sub>. It was purchased from Euriso-top (Gif-sur -Yvette, France). The gas flow rate was measured with a Mass Flow Controller (MFC), calibrated for He. The flow rate of NH<sub>3</sub>

$$\varphi_{\rm NH_3} = f_{\rm NH_3} \ \varphi_{\rm He} \tag{2}$$

was calculated by using  $f_{NH_3} = 0.551$  in our conditions.



Figure S1. Q ICP-MS mass spectra for 10 ppb of <sup>139</sup>La (a) and Eu (b) standard solutions, and <sup>160</sup>Gd (c) <sup>237</sup>Np (d) and <sup>241</sup>Am (e) monoisotopic solutions, when using NH<sub>3</sub> as reacting gas (0.05 mL.min<sup>1</sup>). LaO<sup>\*</sup> and GdO<sup>\*</sup> arise from secondary reactions with gas impurities as confirmed by using ND<sub>3</sub> instead of NH<sub>3</sub> (see ref.5 of the communication).

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