Cite this: Chem. Sci., 2022, 13, 681

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 6th October 2021
Accepted 15th December 2021
DOI: 10.1039/d1sc05517h
rsc.li/chemical-science

# Structure, reactivity and luminescence studies of triphenylsiloxide complexes of tetravalent lanthanides $\dagger$ † 

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#### Abstract

Among the 14 lanthanide elements ( $\mathrm{Ce}-\mathrm{Lu}$ ), until recently, the tetravalent oxidation state was readily accessible in solution only for cerium while $\operatorname{Pr}((\mathrm{v}), \mathrm{Nd}(\mathrm{iv}), \mathrm{Dy}(\mathrm{iv})$ and $\mathrm{Tb}(\mathrm{iv})$ had only been detected in the solid state. The triphenylsiloxide ligand recently allowed the isolation of molecular complexes of $\mathrm{Tb}(\mathrm{iv})$ and $\operatorname{Pr}(\mathrm{iv})$ providing an unique opportunity of investigating the luminescent properties of $\operatorname{Ln}(\mathrm{iv})$ ions. Here we have expanded the coordination studies of the triphenylsiloxide ligand with $\operatorname{Ln}(i I)$ and $\operatorname{Ln}(i v)$ ions and we report the first observed luminescence emission spectra of $\operatorname{Pr}(\mathrm{iv})$ complexes which are assigned to a ligand-based emission on the basis of the measured lifetime and computational studies. Binding of the ligand to the $\operatorname{Pr}(i v)$ ion leads to an unprecedented large shift of the ligand triplet state which is relevant for future applications in materials science.


## Introduction

Among the 14 lanthanide elements (Ce-Lu), until recently, the tetravalent oxidation state was readily accessible only for cerium while $\operatorname{Pr}(\mathrm{Iv}), \mathrm{Nd}(\mathrm{Iv}), \mathrm{Dy}(\mathrm{Iv})$ and $\mathrm{Tb}(\mathrm{iv})$ had only been detected in solid inorganic compounds, double fluorides, oxides, perovskites, ${ }^{1}$ or in concentrated aqueous carbonate solutions. ${ }^{2}$ Seminal attempts by various groups to isolate molecular complexes of $\operatorname{Tb}(\mathrm{rv})$ and $\operatorname{Pr}(\mathrm{rv})$ were unsuccessful, ${ }^{3}$ and only recently $\operatorname{Pr}(\mathrm{Iv})^{4}$ and $\operatorname{Tb}(\mathrm{Iv})^{5}$ complexes were finally successfully isolated in organic solvents by taking advantage of the tuning of the +III/+IV oxidation potential by ligand field effects and by a careful choice of the chemical oxidizing agents. These landmark results open a whole new field in lanthanide coordination chemistry and related photonic and magnetic applications.

The first molecular complexes of terbium(iv) were isolated using the bulky $\sigma$ and $\pi$ donor supporting ligands tris(tertbutoxy)siloxide ${ }^{5 a}$ and tris(amidyl)imidophosphorane. ${ }^{5 b}$ Both complexes display a coordination sphere of the Tb ion fully saturated by the supporting ligand. Despite the very different

[^0]redox potentials reported for the two $\mathrm{Tb}(\mathrm{iv})$ complexes (the imidophosphorane complex has a 1.13 V more negative reduction potential than the tris(tertbutoxy)siloxide complex), no other $\operatorname{Ln}(\mathrm{Iv})$ complexes, besides $\mathrm{Ce}(\mathrm{Iv}),{ }^{6}$ were so far isolated using these ligands.

Tris(triphenylsiloxide) complexes of $\operatorname{Ln}$ (III) were extensively studied after being first reported 30 years ago, ${ }^{7-10}$ and the structure of several $\left[\mathrm{M}^{\mathrm{III}}\left(\mathrm{OSiPh}_{3}\right)_{3}(\mathrm{THF})_{3}\right]$ complexes $\left(\mathrm{M}=\mathrm{La},{ }^{7 b}\right.$ $\mathrm{Ce},{ }^{8} \mathrm{Sm},{ }^{9}$ Dy, ${ }^{10}$ and $\mathrm{Er}^{11}$ ) is known. The triphenylsiloxide ligand was also reported to stabilize the $\mathrm{Ce}(\mathrm{Iv})$ complex $\left[\mathrm{Ce}\left(\mathrm{OSiPh}_{3}\right)_{4}(-\right.$ $\mathrm{DME})]^{12}$ more than thirty years ago but no attempts were made until recently ${ }^{4,5 c}$ to use the same ligand for stabilizing other $\operatorname{Ln}(\mathrm{Iv})$ ions.

Only in 2020 our group isolated the third example of a $\mathrm{Tb}(\mathrm{iv})$ molecular complex using the triphenylsiloxide ligand, $\left[\mathrm{Tb}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{MeCN})_{2}\right], \quad 3-\mathbf{T b}^{\mathrm{MeCN}} .{ }^{5 c} \quad$ The triphenylsiloxide ligand also allowed the isolation and characterization of the first molecular complex of $\operatorname{Pr}(\mathrm{iv})$ that proved to be isomorphous to the $\mathrm{Tb}(\mathrm{iv})$ one $\left[\operatorname{Pr}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{MeCN})_{2}\right]$, $3-\mathrm{Pr}^{\mathrm{MeCN}}{ }^{4}$ These complexes provide the opportunity of investigating for the first time the luminescent properties of molecular complexes of lanthanides in the +IV oxidation state.

For most $\operatorname{Ln}$ (iii) complexes luminescence emission arises from parity-forbidden $4 \mathrm{f} \rightarrow 4 \mathrm{f}$ transitions, ${ }^{13}$ but $\mathrm{Ce}($ (iI) complexes show broad absorption and emission bands in the UV/visible range which are due to electric dipole allowed 4f $\leftrightarrow$ $5 d$ transitions. ${ }^{14} \mathrm{Ce}($ III) emission has shorter lifetimes compared to other lanthanides but high emission intensities. These features have led to the application of $\mathrm{Ce}($ (ii) in light emitting diodes and in photocatalysis. ${ }^{15}$ Similar applications were
identified for $4 \mathrm{f} \leftrightarrow 5 \mathrm{~d}$ transitions of divalent lanthanides ions, such as $\mathrm{Sm}^{2+}$ and $\mathrm{Eu}^{2+} .{ }^{16}$ Considering that $\operatorname{Pr}(\mathrm{Iv})$ has formally the electronic structure of $\mathrm{Ce}(\mathrm{III}), 4 \mathrm{f}^{1}$, but with significantly higher energy and possibly non accessible 5d orbitals, we set out to investigate if it could display luminescence emission. It should be noted that there are no previous reports of luminescence emission from $\operatorname{Pr}(\mathrm{Iv})$ ions in solid state or in solution. Complexes of $\mathrm{Ce}(\mathrm{Iv})$ can display intense ligand-to-metal charge transfer transition (LMCT) and highly reactive LMCT excited states that can be tuned by ligand environment ${ }^{17,15 b, 15 c, 15 g}$ and can lead to photocatalytic application. ${ }^{18}$ However, examples of emissive $\mathrm{Ce}(\mathrm{Iv})$ complexes are extremely rare and were interpreted in term of an emission of Ce (III) from a metal-centred fd excited state, which is generated by electron transfer from the ligand to $\mathrm{Ce}(\mathrm{Iv}) .{ }^{19}$ Here we have expanded the coordination studies of the triphenylsiloxide ligand with $\operatorname{Ln}($ III $)$ and $\operatorname{Ln}($ iv $)$ ions and we report the first observed luminescence emission spectra of $\operatorname{Pr}(\mathrm{Iv})$ complexes which are assigned to a ligand-based emission. Binding of the ligand to $\operatorname{Pr}(\mathrm{iv})$ leads to an unprecedented large shift of the ligand triplet state. These findings are in line with theoretical prediction.

## Results and discussion

## Syntheses of lanthanide(iii) complexes

It was previously shown that tetrakis $\operatorname{Ln(iII)~complexes~with~}$ siloxide ligands are easily prepared using the salt metathesis route ${ }^{20 a, 6 a, 5 a, 20 b}$ and we recently reported that the potassium-tetrakis-(triphenylsiloxide) $\quad \mathrm{Tb}($ III $) \quad$ complex $\quad\left[\mathrm{KTb}\left(\mathrm{OSiPh}_{3}\right)_{4}(-\right.$ THF)], $\mathbf{1 - T b}$, is obtained in $76 \%$ yield by direct reaction of $\mathrm{TbI}_{3}$ with 4 equivalents $\operatorname{KOSiPh}_{3}\left(\mathbf{K L}^{\mathbf{P h}}\right)($ Scheme 1$) .{ }^{5 c}$ However, when we attempted to synthesize the analogous $\operatorname{Pr}(\mathrm{III})$ complex using the same salt metathesis route, the reaction between $\mathrm{PrCl}_{3}$ and 4 equivalents $\mathbf{K L}^{\mathbf{P h}}$ resulted in a mixture of the tris- and tetrakis(triphenylsiloxide) complexes of $\operatorname{Pr}(\mathrm{III}) .{ }^{4}$ In contrast, we recently showed that the previously reported tris(triphenylsiloxide) $\operatorname{Ln}($ III $)$ complexes $\left[\operatorname{Ln}^{\text {III }}\left(\mathrm{OSiPh}_{3}\right)_{3}(\mathrm{THF})_{3}\right]\left(\mathrm{Ln}=\mathrm{Ce},{ }^{8} \mathrm{Pr},{ }^{8}\right)$ could be used as precursors to access analytically pure tetrakis analogues $\left[\mathrm{KLn}^{\mathrm{III}}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{THF})_{2+x}\right], \mathbf{1 - L n},(\mathrm{Ln}=\mathrm{Ce}, x=0$; $\mathrm{Pr}, x$ $=1$ ) upon addition of 1 equivalent $\mathbf{K L}^{\mathbf{P h}}$ (Scheme 2). ${ }^{4}$ Using this route, tetrakis(triphenylsiloxide) complexes of trivalent Nd and Gd were also prepared.

The $\left[\mathrm{KNd}^{\mathrm{III}}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{THF})_{2}\right]$, 1-Nd, and $\left[\mathrm{KGd}^{\mathrm{III}}\left(\mathrm{OSiPh}_{3}\right)_{4}(-\right.$ THF $)_{3}$ ], 1-Gd, complexes were obtained in $75 \%$ and $82 \%$ yield respectively, by addition of 1 equivalent $\mathbf{K L}^{\mathbf{P h}}$ to their $\left[\operatorname{Ln}^{\text {III }}\left(\mathrm{OSiPh}_{3}\right)_{3}(\mathrm{THF})_{3}\right]$ precursors in THF (Scheme 2).


Scheme 1 Synthesis of 1-Tb. ${ }^{5 c}$


Scheme 2 Synthesis of 1- $\mathrm{Ln}\left(\mathrm{Ln}=\mathrm{Ce},{ }^{4} \mathrm{Pr},{ }^{4} \mathrm{Nd}, \mathrm{Gd}\right)$.

Upon addition of 1 equivalent 2.2 .2 -cryptand to THF-d ${ }_{8}$ solutions of 1-Ln $\left(\mathrm{Ln}=\mathrm{Ce}, \operatorname{Pr},{ }^{4} \mathrm{Nd}\right.$ and $\left.\mathrm{Tb}^{5 c}\right)$ a shift in the signals assigned to the siloxide protons can be identified in the ${ }^{1} \mathrm{H}$ NMR spectra of the reaction mixtures (Fig. S9 and S10 $\ddagger$ ). The shift is induced by the removal of the potassium cation from the coordination sphere of the complexes to form an outer-sphere $[\mathrm{K}(2.2 .2 \text {-cryptand })]^{+}$counterion and confirms that in absence of 2.2.2-cryptand, the potassium cation in 1- Ln remains bound to the complexes in solution.

Further addition of 1 equivalent $\mathbf{K L}{ }^{\mathbf{P h}}$ to the tetrakis complex 1-Nd in toluene resulted in the formation of the pentakis [ $\mathrm{K}_{2}-$ $\left.\mathrm{Nd}^{\mathrm{III}}\left(\mathrm{OSiPh}_{3}\right)_{5}\right], 2-\mathrm{Nd}$, in $78 \%$ yield (Scheme 3 and Fig. S11\$). Single crystals of 2-Nd suitable for X-ray diffraction analysis were isolated from a saturated toluene solution stored at $-40^{\circ} \mathrm{C}$ overnight. Complex 2-Nd provides the first example of a $\operatorname{Ln}($ III $)$ complex containing five siloxide ligands ${ }^{21}$ demonstrating that electron-rich triphenylsiloxide ligands allow the synthesis of pentakis $\operatorname{Ln}($ III $)$ complexes. In contrast, performing the ligand addition in coordinating solvent such as THF, or redissolving 2Nd in THF, only resulted in the presence in solution of the tetrakis $\mathbf{1 - N d}$ and 1 equivalent $\mathbf{K L}^{\mathbf{P h}}$ as shown by ${ }^{1} \mathrm{H}$ NMR spectroscopy (Fig. S12 $\ddagger$ ). These results show that the existence, in solution, of the pentakis $\mathbf{2 - N d}$ complex is limited to noncoordinating solvent and its decomposition in THF can be explained by a competitive binding between the THF and the fifth anionic siloxide ligand.

## Syntheses of lanthanide(iv) complexes

With the electron-rich tetrakis-siloxide complexes on hand we pursued their oxidation to yield the tetravalent analogues. The complex 1-Ce can be oxidized with 1.1 equivalent of $\mathrm{AgBPh}_{4}$ in THF resulting in the formation of the cerium(iv) complex


Scheme 3 Synthesis of 2-Nd.



Scheme 4 Synthesis of $3-\mathrm{Ce}^{\text {THF }}$
$\left[\mathrm{Ce}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{THF})_{2}\right], 3-\mathrm{Ce}^{\mathrm{THF}}$ (Scheme 4, Fig. S13 and S14 $\ddagger$ ). Slow diffusion at $-40{ }^{\circ} \mathrm{C}$ of $n$-hexane into the concentrated toluene filtrate of the reaction mixture resulted in the isolation of colorless crystals of $3-\mathrm{Ce}^{\mathrm{THF}}$ in $61 \%$ yield. Interestingly, once isolated, the colorless crystals of $3-\mathrm{Ce}^{\mathbf{T H F}}$ do not redissolve in toluene. This differs from the rather high solubility of its precursor 1-Ce in toluene. Single crystal of 3-Ce ${ }^{\text {THF }}$ suitable for X-ray diffraction analysis were grown from storage of a concentrated THF solution of $3-\mathrm{Ce}^{\mathrm{THF}}$ at $-40{ }^{\circ} \mathrm{C}$ overnight. The synthesis and the crystal structure of the DME analogue $\left[\mathrm{Ce}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{DME})\right]$ were reported more than 30 years ago., ${ }^{8,12}$ The $\left[\mathrm{Ce}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{DME})\right]$ was previously prepared either by salt metathesis, from $\left[\left(\mathrm{NH}_{4}\right)_{2} \mathrm{Ce}\left(\mathrm{NO}_{3}\right)_{6}\right]$, ${ }^{8}$ or by protonolysis from $\left[\mathrm{Ce}\left(\mathrm{O}^{\mathrm{i}} \mathrm{Pr}\right)_{4}\right] .^{12}$ The present results show that we can also access the tetrakis(triphenylsiloxide) $\mathrm{Ce}(\mathrm{rv})$ complex, 3-Ce ${ }^{\mathrm{THF}}$, in good yield, by direct oxidation of the potassium-tetrakis(triphenylsiloxide) Ce (iii) complex, 1-Ce.

Differently from 1-Ce, the oxidation of 1-Pr and 1-Tb has proven impossible with a mild oxidizing agent such as silver salts. ${ }^{4,5 c}$ However, we recently showed that the use of a stronger oxidizing agent such as tris(4-bromophenyl)aminium salts ( $E^{0}=$ $0.67 \mathrm{~V} v s$. Fc compared to $E^{0}\left(\mathrm{Ag}^{+}\right)=0.04 \mathrm{~V} v$ s. Fc in MeCN$)^{22}$ leads to the oxidation of the 1-Pr and 1-Tb complex, ${ }^{4,5 c}$ yielding the third $\mathrm{Tb}(\mathrm{iv})$ molecular complex and the first example of a molecular complex of praseodymium in the +IV oxidation state, $\left[\mathrm{Ln}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{MeCN})_{2}\right], 3-\mathrm{Ln}{ }^{\mathrm{MeCN}}(\mathrm{Ln}=\mathrm{Pr}, \mathrm{Tb})$ (Scheme 5)..$^{4,5 c}$ Complex $3-\mathbf{T b}^{\mathbf{M e C N}}$ is insoluble in $n$-hexane, has a very


Scheme 5 Synthesis of 3- $\mathrm{Ln}^{\mathrm{MeCN}}(\mathrm{Ln}=\mathrm{Pr}, \mathrm{Tb})^{4.5 \mathrm{c}}$ and isolation of 3$\mathrm{Tb}^{\text {THF }}$.
limited solubility in toluene and acetonitrile, but is completely soluble in THF. Complex $3-\mathbf{P r}^{\mathbf{M e C N}}$ shows similar solubility behaviour as $\mathbf{3 - T b}{ }^{\mathbf{M e C N}}$ except that it is also completely soluble in toluene. Single crystals of the $\mathrm{Tb}(\mathrm{Iv})$ THF adduct, $\left[\mathrm{Tb}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{THF})_{2}\right], 3-\mathrm{Tb}^{\mathrm{THF}}$ (Scheme 5), could be grown in this work from a concentrated THF solution of $\mathbf{3 - T b}{ }^{\mathbf{M e C N}}$ stored at $-40{ }^{\circ} \mathrm{C}$ overnight. The molecular structures of the MeCN adduct $\mathbf{3 - T b}{ }^{\mathbf{M e C N}}$ and the THF adduct $\mathbf{3 - T b}{ }^{\text {THF }}$ have the same coordination environment except for the nature of the bound solvent molecules (vide infra). It is noticeable that the bound solvent molecules can be easily exchanged without significant changes in the structure.

The addition of $\left[\mathrm{N}\left(\mathrm{C}_{6} \mathrm{H}_{4} \mathrm{Br}\right)_{3}\right]\left[\mathrm{SbCl}_{6}\right]$ to a MeCN suspension of 1-Nd did not result in the isolation of any $\operatorname{Nd}(\mathrm{Iv})$ oxidation products. It should be noted that 1-Nd is not soluble in MeCN. Furthermore, cyclic voltammetry experiments performed on 2 mM THF solutions of $\mathbf{1 - N d}$ with $\left[\mathrm{NBu}_{4}\right]\left[\mathrm{B}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{4}\right]$ as the supporting electrolyte remained silent. This also differs from 1-Tb and 1-Pr which displayed very well-defined oxidation events at $E_{\mathrm{pa}}=0.41 \mathrm{~V} v s . \mathrm{Fc}$ and $E_{\mathrm{pa}}=0.65 \mathrm{~V} v s . \mathrm{Fc}$, respectively. ${ }^{4,5 c}$ This can be explained by the large difference in the measured aqueous $\operatorname{Ln}($ Iv $) / \operatorname{Ln}($ III $)$ reduction potentials between the free Nd and the $\mathrm{Tb} / \mathrm{Pr}$ ions $(\mathrm{Nd}($ Iv $) / \mathrm{Nd}($ III $)-\mathrm{Tb}($ Iv $) / \mathrm{Tb}($ III $)=1.9 \mathrm{~V} ; \mathrm{Nd}(\mathrm{Iv}) /$ $\mathrm{Nd}(\mathrm{III})-\operatorname{Pr}(\mathrm{Iv}) / \operatorname{Pr}(\mathrm{III})=1.8 \mathrm{~V}) .{ }^{23}$

The addition of 1 equivalent $\mathrm{OPR}_{3}(\mathrm{R}=\mathrm{Et}, \mathrm{Ph})$ to $3-\mathrm{Tb}^{\mathbf{M e C N}}$ in THF was previously shown to yield the $\mathrm{Tb}(\mathrm{iv})$ complexes $\left[\mathrm{Tb}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}\left(\mathrm{OPPh}_{3}\right)\right], 4-\mathbf{T b}^{\mathrm{Ph}}$ and $\left[\mathrm{Tb}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}\left(\mathrm{OPEt}_{3}\right)\right]$, 4$\mathbf{T b}^{\text {Et }}$ (Scheme 6). ${ }^{5 \boldsymbol{c}}$ These complexes demonstrated the relevance of the two open coordination sites in $3-\mathbf{T b}^{\mathbf{M e C N}}$ for reactivity studies and the possibility of replacement of the labile solvent molecules. Interestingly, the phosphinoxide adducts $\mathbf{4 - T b}{ }^{\mathbf{R}}$ displayed a much higher solubility in toluene compared to the acetonitrile adduct $3-\mathbf{T b}^{\mathbf{M e C N}}$.

Here we have explored the reactivity of the previously reported ${ }^{4} \operatorname{Pr}(\mathrm{Iv})$ complex $3-\mathbf{P r}^{\text {MeCN }}$ with phosphinoxide and we found that the complex $\left[\mathrm{Pr}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}\left(\mathrm{OPPh}_{3}\right)(\mathrm{MeCN})\right], 4-\mathrm{Pr}^{\mathrm{Ph}}$, can be synthetized by treating $3-\mathbf{P r}^{\mathbf{M e C N}}$ with 1 equivalent triphenylphosphine oxide in toluene at room temperature (Scheme 7). The slow diffusion of $n$-hexane into the toluene reaction mixture at $-40^{\circ} \mathrm{C}$ allowed the isolation of dark brown crystals of $\mathbf{4}-\mathbf{P r}^{\mathbf{P h}}$ in $90 \%$ yield. The ${ }^{1} \mathrm{H}$ NMR spectrum of $\mathbf{4}-\mathbf{P r}^{\mathbf{P h}}$ in toluene-d ${ }_{8}$ shows four signals at $\delta=7.2 \mathrm{ppm}, 7.1 \mathrm{ppm}$, 6.9 ppm , and 6.7 ppm corresponding to the 75 protons of the siloxide and the phosphinoxide ligands (Fig. S15 $\ddagger$ ). The binding of the $\mathrm{OPPh}_{3}$ moiety is reversible in coordinating solvent such as THF, resulting in the formation of $3-\mathbf{P r}^{\mathbf{T H F}}$ and free $\mathrm{OPPh}_{3}$


Scheme 6 Synthesis of $\left.4-\mathrm{Tb}^{\mathrm{R}}(\mathrm{R}=\mathrm{Et}, \mathrm{Ph})\right)^{5 \mathrm{c}}$


Scheme 7 Synthesis of $4-\mathrm{Pr}^{\mathrm{Ph}}$.
immediately after the dissolution of $\mathbf{4 - P r}{ }^{\mathbf{P h}}$ in THF-d ${ }_{8}$ (Fig. S17\$).

Interestingly, the phosphinoxide adduct $4-\mathbf{P r}^{\mathbf{P h}}$ shows a higher solution stability than $3-\mathbf{P r}^{\mathbf{M e C N}}$. Notably, UV-visible stability studies showed that after storing a 3 mM solution of $4-\mathbf{P r}^{\mathbf{P h}}$ in toluene at room temperature for 24 hours, $80 \%$ of 4 $\mathbf{P r}^{\mathbf{P h}}$ is still present in solution (Fig. S22 $\ddagger$ ). In contrast, UVvisible stability studies performed at room temperature on a 3 mM toluene solution of $3-\mathbf{P r}^{\mathbf{M e c N}}$ only showed $40 \%$ of the initial absorbance after 24 hours. ${ }^{4}$

A similar high increase in solution stability was observed for the phosphinoxide adduct of $\mathrm{Tb} 4-\mathrm{Tb}^{\mathbf{R}}(\mathrm{R}=\mathrm{Et}, \mathrm{Ph})$ compared to the THF and MeCN complexes. Notably, 3 mM toluene solutions of $4-\mathbf{T b}^{\mathbf{R}}(\mathrm{R}=\mathrm{Et}, \mathrm{Ph})$ display an unchanged absorbance ( $<2 \%$ ) in the UV-vis spectra for up to 24 hours (Fig. S23 and S24廿). After 4 days at room temperature, $80 \%$ of $\mathbf{4 - T b}{ }^{\mathbf{P h}}$ (Fig. S23 $\ddagger$ ) and $70 \%$ of $4-\mathbf{T b}^{\mathbf{E t}}$ (Fig. S24 $\ddagger$ ) are still present in solution.

## Solid state structures of lanthanide(iiI) complexes

The solid-state structures of $\mathbf{1 - L n}(\mathrm{Ln}=\mathrm{Ce}, \mathrm{Nd}$ and Gd) as determined by X-ray diffraction studies present a sixcoordinated $\operatorname{Ln}($ III $)$ metal center in a distorted octahedral geometry (Fig. 1). Four coordination sites are occupied by an oxygen atom from the $-\mathrm{OSiPh}_{3}$ ligands while the two others are filled by oxygen atoms from THF solvent molecules. An
analogous coordination environment of the Ln ion is found in the $\mathbf{1 - L n}\left(\mathrm{Ln}=\mathrm{Ce}, \mathrm{Pr},{ }^{4} \mathrm{Nd}\right.$ and Gd ) complexes. The only difference in these structures is the presence of one THF bound to the potassium counterion in $\mathbf{1 - P r}$ and $\mathbf{1 - G d}$ which is not present in 1-Ce and 1-Nd. Differently, the solid state structure of the 1-Tb complex contains only one Tb-bound THF molecule with a five-coordinated $\mathrm{Tb}($ III $)$ metal center in a distorted trigonal-bipyramidal molecular geometry. ${ }^{5 c}$ The lower coordination number found for the $\mathrm{Tb}($ III) ion is consistent with its decreased ionic radius compared to the $\mathrm{Gd}(\mathrm{III})$ ion. ${ }^{24}$

The decrease of the mean value of the $\mathrm{Ln} 1-\mathrm{O}_{\text {siloxide }}$ bond distances identified along the series 1-Ce (2.30(3) $\AA)$, 1-Pr $(2.27(2) \AA),{ }^{4}$ 1-Nd $(2.27(4) \AA)$ and 1-Gd (2.22(2) $\left.\AA\right)$ (Table 1) is consistent with the difference in Shannon's ionic radii of these 6-coordinate $\operatorname{Ln}($ III) ions ( $\mathrm{Ce}($ III $), 1.01 \AA ; \operatorname{Pr}($ III $), 0.99 \AA ; \operatorname{Nd}($ III $)$, $0.983 \AA$ and $\operatorname{Gd}($ III $), 0.938 \AA) .{ }^{25}$ The mean value of the Ce1$\mathrm{O}_{\text {siloxide }}$ bond distances in 1-Ce is larger than the one found in the six-coordinate tris-complex $\left[\mathrm{Ce}^{\mathrm{III}}\left(\mathrm{OSiPh}_{3}\right)_{3}(\mathrm{THF})_{3}\right]$ (2.222 (4) $\AA$ ). ${ }^{8}$ The observed bond elongation is explained both in terms of the increased steric hindrance and increased negative charge provided by the additional siloxide ligand bound to the cerium ion in the tetrakis 1-Ce complex compared to the THF ligand in the tris-complex. In all the 1-Ln complexes ( $\mathrm{Ln}=\mathrm{Ce}, \mathrm{Pr}, \mathrm{Nd}, \mathrm{Gd}$ and Tb ), the $\mathrm{K}^{+}$ion lies in close proximity of the lanthanide center ( 3.5143 (12)-3.6921 (7) $\AA$; see Table 2) and connects the siloxide ligands through the anionic oxygen and through cation $-\pi$ interactions with the phenyl groups.

The complex 2-Nd crystallizes in the monoclinic $P 2_{1} / n$ space group. The solid-state structure of 2 -Nd shows a fivecoordinated $\mathrm{Nd}($ III $)$ metal center bound by five monodentate triphenylsiloxide ligands in a distorted trigonal bipyramidal geometry (Fig. 2). The triphenylsiloxide ligands bind the two potassium in close proximity to the $\mathrm{Nd}($ (III $)$ metal center through the anionic oxygen and cation $-\pi$ interactions with the phenyl groups. The two potassium cations lie at a distance from the $\mathrm{Nd}($ III) center of $3.6808(8) \AA$ for K1 and 3.7774(8) Å for K2. The K1 potassium is at a similar distance from the metal center


Fig. 1 Solid-state molecular structures of 1-Ln (50\% probability ellipsoids); hydrogen atoms, phenyl groups not bound to the potassium ions and residual solvent molecules are omitted for clarity. (a) 1-Ce, selected distances ( $\AA$ ): Ce1- $\mathrm{O}_{\text {siloxide }}$ range $=2.268(5)-2.351(5)$; mean $\mathrm{Ce} 1-\mathrm{O}_{\text {siloxide }}=$ $2.30(3) ; \mathrm{Ce} 1-\mathrm{O}_{\text {THF }}=2.598(6)-2.605(6) ; \mathrm{Ce} 1-\mathrm{K} 1=3.6305(16)$. (b) $1-\mathrm{Nd}$, selected distances $(\AA)$ : Nd1- $\mathrm{O}_{\text {siloxide }}$ range $=2.231(2)-2.3300(19) ;$ mean $\mathrm{Nd} 1-\mathrm{O}_{\text {siloxide }}=2.27(4) ; \mathrm{Nd} 1-\mathrm{O}_{\mathrm{THF}}=2.573(2)-2.574(2)$ : $\mathrm{Nd} 1-\mathrm{K} 1=3.6925(6)$. (c) $1-\mathrm{Gd}$, selected distances $(\AA)$ : $\mathrm{Gd} 1-\mathrm{O}_{\text {siloxide }}$ range $=2.212(3)-$ 2.253(3); mean $\mathrm{Gd} 1-\mathrm{O}_{\text {siloxide }}=2.22(2) ; \mathrm{Gd} 1-\mathrm{O}_{\mathrm{THF}}=2.488(3)-2.491(3) ; \mathrm{Gd} 1-\mathrm{K} 1=3.5143(12)$.

Table 1 Comparison of the $\mathrm{Ln}-\mathrm{O}_{\text {siloxide }}$ bond lengths $(\AA \AA)$ in the complexes $1-\mathrm{Ln}\left(\mathrm{Ln}=\mathrm{Ce}, \mathrm{Pr},{ }^{4} \mathrm{Nd}, \mathrm{Gd}\right.$ and $\left.\mathrm{Tb}{ }^{5 c}\right), 2-\mathrm{Nd}, 3-\mathrm{Ln}{ }^{\mathrm{THF}}(\mathrm{Ln}=\mathrm{Ce}, \mathrm{Tb})$, $3-\mathrm{Ln}^{\mathrm{MeCN}}\left(\mathrm{Ln}=\mathrm{Pr},{ }^{4} \mathrm{~Tb}^{5 \mathrm{c}}\right), 4-\mathrm{Pr}^{\mathrm{Ph}}$ and $4-\mathrm{Tb}^{\mathrm{R}}(\mathrm{R}=\mathrm{Ph}, \mathrm{Et})^{5 \mathrm{c}}$


Table 2 Comparison of the Ln1-K distances ( $A \AA$ ) in the complexes 1-Ln ( $\mathrm{Ln}=\mathrm{Ce}, \mathrm{Pr}^{4}{ }^{4} \mathrm{Nd}, \mathrm{Gd}$ and $\mathrm{Tb}^{5 c}$ ) and $2-\mathrm{Nd}$

|  | 1-Ce |  | 1-Pr | 1-Nd |
| :--- | :--- | :--- | :--- | :--- |
| Ln1-K | $3.6305(16)$ |  | $3.67(14)^{a}$ | $3.6925(6)$ |
|  | 1-Gd | 1-Tb | 2-Nd |  |
| Ln1-K | $3.5143(12)$ | $3.6921(7)$ | $3.6808(8), 3.7774(8)$ |  |

${ }^{a}$ Average value of two Ln1-K distances due to the disordered $\mathrm{K}^{+}$over two positions.


Fig. 2 Solid-state molecular structure of $2-\mathrm{Nd}$ (50\% probability ellipsoids). Hydrogen atoms, phenyl groups not bound to the potassium ions and residual solvent molecules are omitted for clarity. Selected distances $(\AA \AA)$ : Nd1 $-\mathrm{O}_{\text {siloxide }}$ range $=2.206(2)-2.336(2)$; mean $\mathrm{Nd} 1-\mathrm{O}_{\text {siloxide }}=2.28(4) ; \mathrm{Nd} 1-\mathrm{K} 1=3.6808(8) ; \mathrm{Nd} 1-\mathrm{K} 2=3.7774(8)$.
compared to the Nd1-K1 distance in 1-Nd (3.6925(6) A). However, the second potassium (K2) is found at a larger distance, suggesting a weaker binding. The presence of five triphenylsiloxide ligands around the $\mathrm{Nd}(\mathrm{III})$ ion in 2-Nd does not significantly affect the $\mathrm{Nd}-\mathrm{O}_{\text {siloxide }}$ distances $\left(\left(\mathrm{Nd} 1-\mathrm{O}_{\text {siloxide }}\right)_{\text {avg }}\right.$ $=2.28(4) \AA)$ compared to those found in the tetrakis complex 1 Nd $\left(\left(\mathrm{Nd} 1-\mathrm{O}_{\text {siloxide }}\right)_{\text {avg }}=2.27(4) \AA\right)$.

## Solid state structures of lanthanide(iv) complexes

The complexes 3 -Ce ${ }^{\mathrm{THF}}$ and $3-\mathrm{Tb}^{\mathrm{THF}}$ are isomorphous and crystallize in the monoclinic $C 2 / c$ space group while $3-\mathbf{L n}^{\text {MeCN }}$ ( $\mathrm{Ln}=\mathrm{Pr}, \mathrm{Tb}$ ) was reported to crystallize in the orthorhombic $P c a 2_{1}$ space group. ${ }^{1,5 c}$ Their molecular structures are presented in Fig. 3 and show $\operatorname{Ln}(\mathrm{v})$ metal centers coordinated by two THF molecules and four triphenylsiloxide ligands in a distorted octahedral geometry. The Ce1-O $\mathrm{O}_{\text {siloxide }}$ distances in 3 - $\mathrm{Ce}^{\text {THF }}$ (2.109(3)-2.154(4) $\AA$ ) are consistent with the $\mathrm{Ce} 1-\mathrm{O}_{\text {siloxide }}$ bond length found in the previously reported DME analogue $\left[\mathrm{Ce}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}(\mathrm{DME})\right](2.10(1)-2.13(1) \AA) .{ }^{12}$ Furthermore, the shorter Ce1-O ${ }_{\text {siloxide }}$ bond distances in $3-\mathrm{Ce}^{\mathrm{THF}}$ (mean value $=$ $2.13(3) \AA \AA)$ compared to those of the $\mathrm{Ce}(\mathrm{III})$ complex 1-Ce (2.30(3) A) corroborate the +IV oxidation state of the cerium metal center. This shortening can also be identified in the $\mathrm{Ce} 1-\mathrm{O}_{\text {thF }}$ bond distances (2.501(4) $\AA$ in 3 - $\mathrm{Ce}^{\mathrm{THF}} ; 2.598(6)$ and $2.605(6) \AA$ in 1-Ce). The presence of THF instead of MeCN coordinated to the $\mathrm{Tb}(\mathrm{Iv})$ metal center does not significantly affect the mean Tb1$\mathrm{O}_{\text {siloxide }}$ bond distance in $3-\mathrm{Tb}^{\mathrm{THF}}(2.07(2) \AA$ ) compared to 3 $\mathrm{Tb}^{\text {MeCN }}(2.06(2) \AA)^{5 c}{ }^{5 c}$ The $0.03 \AA$ difference in the mean Ln1$\mathrm{O}_{\text {siloxide }}$ bond distances between 3-Ce ${ }^{\mathrm{THF}}$ and 3-Pr ${ }^{\mathrm{MeCN}}$ (Table 1) is consistent with the difference in the 6 -coordinate Shannon radii of $\operatorname{Ce}(\mathrm{vv})(0.87 \AA)$ and $\operatorname{Pr}(\mathrm{Iv})(0.85 \AA)$ ions. ${ }^{25}$ However, the $0.06-0.07 \AA$ difference in $\mathrm{Ln} 1-\mathrm{O}_{\text {siloxide }}$ bond distances between


Fig. 3 (a) Solid-state molecular structure of $3-\mathrm{Ce}^{\text {THF }}$ (a) and $3-\mathrm{Tb}^{\text {THF }}$ (b) $(50 \%$ probability ellipsoids). Hydrogen atoms, phenyl groups and one residual solvent molecule are omitted for clarity. Selected distances $(A ̊): C e 1-O_{\text {siloxide }}$ range $=2.109(3)-2.154(3)$; mean $\mathrm{Ce} 1-$ $\mathrm{O}_{\text {siloxide }}=2.13(3) ; \mathrm{Ce} 1-\mathrm{O}_{\mathrm{THF}}=2.501(4) ; \mathrm{Tb} 1-\mathrm{O}_{\text {siloxide }}$ range $=$ 2.047(2)-2.087(2); mean Tb1-O siloxide $=2.07(2) ; \mathrm{Tb} 1-\mathrm{O}_{\text {THF }}=2.394(3)$. Symmetry transformation used to generate equivalent atoms: ${ }^{1} 1-x$, $+y ; \frac{1}{2}-z$.
$3-\mathrm{Ce}^{\text {THF }}$ and $3-\mathbf{T b}^{\text {MeCN }}$ or $\mathbf{3 - T b}{ }^{\text {THF }}$ (Table 1) is smaller than what is expected from the difference in the Shannon ionic radii of $\mathrm{Ce}(\mathrm{rv})(0.87 \AA)$ and $\mathrm{Tb}(\mathrm{rv})(0.76 \AA) .{ }^{25}$ A smaller than expected difference was already noticed for $3-\mathbf{P r}^{\mathbf{M e C N}}$ compared to 3 $\mathbf{T b}^{\mathbf{M e C N}}$ and an increased covalency in the $\operatorname{Ln}(\mathrm{rv})-\mathrm{O}$ bonds in 3$\mathbf{P r}^{\mathbf{M e C N}}$ was proposed on the basis of DFT studies. ${ }^{4}$ This can be correlated with studies of the tetravalent inorganic $\mathrm{Cs}_{2} \mathrm{RbLnF}_{7}$ materials ( $\mathrm{Ln}=\mathrm{Ce}, \mathrm{Pr}, \mathrm{Nd}, \mathrm{Tb}, \mathrm{Dy}$ ) where an increases in the $4 \mathrm{f}-$ ligand-p orbital hybridization was discovered in the order $\mathrm{Tb}<$ Dy $<\mathrm{Ce}<\mathrm{Pr}<\mathrm{Nd}^{26}{ }^{26}$

The complex 4-Pr ${ }^{\mathbf{P h}}$ crystallizes in the monoclinic $P 2_{1} / c$ space group and shows a six-coordinated $\operatorname{Pr}(\mathrm{Iv})$ metal center in a distorted octahedral geometry (Fig. 4). The $\operatorname{Pr}(\mathrm{Iv})$ ion is coordinated to four monodentate triphenylsiloxide ligands, one acetonitrile molecule and one triphenylphosphinoxide. Compared to 3$\mathbf{P r}^{\mathbf{M e C N}}$, one MeCN moiety is replaced with a phosphinoxide ligand while the second acetonitrile molecule remains bound retaining the distorted octahedral geometry of the complex and the same coordination number $(\mathrm{CN}=6)$. This is different from what was previously observed with the smaller $\mathrm{Tb}(\mathrm{rv})$ ion. Indeed, the addition of 1 equivalent $\mathrm{OPPh}_{3}$ to $3-\mathbf{T b}^{\mathbf{M e C N}}$ resulted in the removal of the two bound acetonitrile molecules. As such, the solid-state structure of the phosphinoxide complex $4-\mathbf{T b}^{\mathbf{R}}(\mathrm{R}$ $=\mathrm{Ph}, \mathrm{Et})$ does not retain the distorted octahedral geometry found in 3-Tb ${ }^{\text {MeCN }}$ but, instead, displays a distorted trigonal bipyramidal molecular geometry with a five-coordinated Tb (iv) metal center. ${ }^{5 c}$ The mean value of the Pr1-O siloxide bond length in $4-\mathbf{P r}^{\mathbf{P h}}(2.15(3) \AA)$ is slightly larger than in $3-\mathbf{P r}^{\mathbf{M e C N}}$ (2.10(1) $\AA$ ). ${ }^{4}$ This can be rationalized in terms of the increased steric hindrance provided by the bulky triphenylphosphine oxide ligand compared to the MeCN molecule. In the case of the smaller $\mathrm{Tb}(\mathrm{Iv})$ ion, the lower coordination number of the phosphinoxide complex $4-\mathbf{T b}^{\mathbf{P h}}(2.06(5) \AA$ ) compared to the 3$\mathbf{T b}^{\mathbf{M e C N}}$ results in similar values of the $\mathrm{Tb}(\mathrm{rv})-\mathrm{O}_{\text {siloxide }}$ bond distance in the two complexes $\left(4-\mathbf{T b}^{\mathbf{P h}}: 2.06(5) \AA\right.$ and 3 $\left.\mathbf{T b}^{\mathbf{M e C N}}: 2.06(2) \AA\right)^{5 c}$ Anyhow, the mean value of the $\operatorname{Pr}(\mathrm{Iv})-$


Fig. 4 Solid-state molecular structure of $4-\operatorname{Pr}^{\mathrm{Ph}}(50 \%$ probability ellipsoids). Hydrogen atoms and phenyl groups of the siloxide ligands are omitted for clarity. Selected distances (Å): Pr1-O siloxide $^{\text {range }}=$ $2.113(2)-2.180(2) ;$ mean $\operatorname{Pr} 1-\mathrm{O}_{\text {siloxide }}=2.15(3) ; \operatorname{Pr} 1-\mathrm{O} 1=2.280(2)$; Pr1-N1 = 2.636(3).
$\mathrm{O}_{\text {siloxide }}$ bond length in $4-\mathbf{P r}^{\mathbf{P h}}(2.15(3) \AA$ ) is considerably smaller than the mean $\operatorname{Pr}(\mathrm{III})-\mathrm{O}_{\text {siloxide }}$ bond length in 1-Pr (2.27(2) $\AA$ ). Furthermore, the $\operatorname{Pr} 1-\mathrm{O}_{\text {phosphinoxide }}$ bond distance in $\mathbf{4}-\mathbf{P r}^{\mathbf{P h}}$ (2.280(2) $\AA)$ is shorter than those found in the literature for $\operatorname{Pr}($ III $)-\mathrm{OPPh}_{3}$ bond lengths (2.333(3)-2.409(5) $\left.\AA \AA^{; 7,28}\right)$ and is consistent with a Pr metal center in the +IV oxidation state.

## Infrared spectra

The IR spectra are reported in Fig. S18 and S19. $\ddagger$ The IR spectra of the free ligand $\mathrm{HOSiPh}_{3}$, the deprotonated ligand $\left(\mathbf{K L}^{\mathbf{P h}}\right)$ and of the lanthanide complexes show similar features with aromatic $\mathrm{C}=\mathrm{H}$ stretching and bending absorptions at 3060$2995 \mathrm{~cm}^{-1}$ and $700 \mathrm{~cm}^{-1}$, respectively, and the Si-O stretching at $1121 \mathrm{~cm}^{-1}$, as reported in the literature. ${ }^{29}$ Upon deprotonation or binding to a Ln cation, the peak at $855 \mathrm{~cm}^{-1}$ assigned to the $\mathrm{Si}-\mathrm{O}-\mathrm{H}$ stretching disappears ${ }^{30}$ and a new intense band appears at $964 \mathrm{~cm}^{-1}$ for the $\operatorname{Ln}$ (III) complexes 1- $\mathrm{Ln}(\mathrm{Ln}=\mathrm{Ce}, \mathrm{Pr}$, $\mathrm{Gd})$ which is assigned to the Ln-O-Si linkage vibration. This value compares well with the range $960-1000 \mathrm{~cm}^{-1}$ reported for tin(iv) aryl- and alkyl siloxyl complexes. In the $\operatorname{Ln}(\mathrm{Iv})$ complexes 3-Ce ${ }^{\text {THF }}, \mathbf{3 - P r}{ }^{\mathbf{M e C N}}$ and $\mathbf{4 - \mathbf { P r } ^ { \mathbf { P h } }}$ the $\mathrm{Ln}-\mathrm{O}-$ Si stretch is significantly shifted to lower frequencies ( $883 / 899 \mathrm{~cm}^{-1}$ ) compared to $\operatorname{Ln}(\mathrm{III})$ complexes but reasonably consistent with the values reported for the tetrakis(triphenylsiloxide)-titanium(rv) complex $\left(926 \mathrm{~cm}^{-1}\right)^{29 a}$ and with the M-O-Si stretch observed for group IV and group V metals ( $920-906 \mathrm{~cm}^{-1}$ )..$^{29 b}$ Two weak features are present at 2276 and $2304 \mathrm{~cm}^{-1}$ for $\mathbf{3 - \mathbf { P r } ^ { \mathbf { M e C N } }}$ and $\mathbf{4 - \mathbf { P r } ^ { \mathbf { P h } }}$ that are assigned to the $\nu_{2}$ and $\left(\nu_{3}+\nu_{4}\right)$ modes of coordinated acetonitrile molecules, as reported previously ${ }^{31}$ (2281 and $2310 \mathrm{~cm}^{-1}$, respectively, for $\left.\left[\mathrm{Nd}^{\mathrm{II}}\left(\mathrm{ClO}_{4}\right)_{x}(\mathrm{MeCN})_{y}\right]\right)$. This observation confirms the complex formulae reported above.

In inorganic matrices, four f-f transitions have been reported for $\mathrm{Ce}($ (III $)$ in the range $2100-3800 \mathrm{~cm}^{-1}$, in line with the crystal field splitting of ${ }^{2} \mathrm{~F}_{7 / 2}$ (ref. 32) while transitions to the $\operatorname{Pr}^{\mathrm{IV}}$ ( ${ }^{2} \mathrm{~F}_{7 / 2}$ ) level have been detected around $2900 \mathrm{~cm}^{-1}$ for fluorides. ${ }^{27}$ Close scrutiny of the IR spectra up to $6000 \mathrm{~cm}^{-1}$ for the $\operatorname{Pr}(\mathrm{rv})$ complexes did not reveal any feature that could be assigned to an f-f transition. This might be due to very weak oscillator strengths and/or overlap with ligand vibrations, particularly in the $2800-3000 \mathrm{~cm}^{-1}$ range. On the other hand, the spectrum of a more concentrated sample of 1-Ce revealed a very weak and sharp band at $2163 \mathrm{~cm}^{-1}$ (see insert, Fig. S18 $\ddagger$ ) not seen in any other spectra including the blank, that might possibly be assigned to the most intense $\mathrm{f}-\mathrm{f}$ transition of $\mathrm{Ce}^{\text {III }}$ identified at $2128 \mathrm{~cm}^{-1}$ in $\mathrm{YGdO}_{3}: \mathrm{Ce}^{\text {III }}{ }^{32}$

## UV-visible absorption spectra

The UV-vis spectra of $\mathbf{K L}{ }^{\mathbf{P h}}, \mathbf{1}-\mathbf{L n}(\mathrm{Ln}=\mathrm{Ce}, \mathrm{Pr}, \mathrm{Gd})$ and $\mathbf{3 - \mathbf { P r } ^ { \mathbf { M e C N } }}$ were recorded in THF (Fig. S20 $\ddagger$ ). The spectrum of $\mathrm{KOSiPh}_{3}$ features two main bands located at 206 and 223 nm and assigned to $\pi \rightarrow \pi^{*}$ transitions; in addition, a very weak feature is observed in the range $250-275 \mathrm{~nm}$. The same bands are present for all the complexes, with negligible red shifts of 12 nm . The ligand absorptions overlap with the $4 \mathrm{f}-5 \mathrm{~d}$ absorptions of $\mathrm{Ce}($ III), usually in the range $200-270 \mathrm{~nm}$, and have much larger oscillator strengths so that these $4 \mathrm{f}-5 \mathrm{~d}$ transitions could
not be identified in 1-Ce. For the $\operatorname{Ln}(\mathrm{rv})$ complexes $\mathbf{3 - P r}{ }^{\mathbf{M e C N} 4}$ and $3-\mathbf{T b}^{\mathbf{M e C N}},{ }^{5 c}$ an additional broad signal has been reported with maximum at $363 \mathrm{~nm}\left(\varepsilon=3800 \mathrm{M}^{-1} \mathrm{~cm}^{-1}\right)$ and $386 \mathrm{~nm}(\varepsilon=$ $3000 \mathrm{M}^{-1} \mathrm{~cm}^{-1}$ ), respectively. These bands can be assigned to ligand-to-metal charge transfers (LMCT) on the basis of previous UV studies on a range of $\operatorname{Ln}(\mathrm{Iv})$ oxides. ${ }^{2,33}$ Since $\mathbf{4}$-Pr ${ }^{\mathbf{P h}}$ reverts to the $3-\mathbf{P r}^{\text {THF }}$ complex in THF, the UV-visible spectrum of $4-\mathbf{P r}^{\mathbf{P h}}$ was measured in toluene (Fig. S21 $\ddagger$ ), as well as the spectra of $4-\mathbf{T b}^{\mathrm{R} 5}$ and of the previously reported $3-\mathbf{P r}^{\mathbf{M e C N}}$ complex ${ }^{4}$ for comparison. All compounds display a similar broad absorption peak, assigned to a LMCT transition, around 374-379 nm with $\varepsilon=4500$ for $\mathbf{4 - P r}{ }^{\mathbf{P h}}, 3300$ for $4-\mathbf{T b}^{\mathbf{P h}},{ }^{5 c}$ and $3150 \mathrm{M}^{-1} \mathrm{~cm}^{-1}$ for $\mathbf{4 - T b} \mathbf{b}^{\mathbf{E t}, 5 c}$ while data for $3-\mathbf{P r}^{\mathbf{M e C N}}$ are similar to those in THF: $\lambda_{\text {max }}=365 \mathrm{~nm}$ and $\varepsilon=4140 \mathrm{M}^{-1} \mathrm{~cm}^{-1} .{ }^{4} \mathrm{As}$ reported in the synthesis description above, time-dependent spectra for $\mathbf{4 - P r}{ }^{\mathbf{P h}}$ and $\mathbf{4 - T b} \mathbf{b}^{\mathbf{R}}$ evidence a slow decomposition of the complexes with time. To ensure reliable photophysical data, samples were measured immediately after their preparation.

## Ligand-centered luminescence of $\mathrm{KL}^{\mathbf{P h}}$ and 1-Gd

Data are reported in Table S1. $\ddagger$ In toluene at room temperature, the $\mathrm{KOSiPh}_{3}\left(\mathbf{K L}^{\mathbf{P h}}\right)$ ligand has one excitation band with maximum at 289 nm and one structured emission band with several components and main maxima at 325 and 339 nm (Fig. S25 $\ddagger$ ). In frozen solution, the excitation is blue shifted at 276 nm while the emission band at $325 / 339 \mathrm{~nm}$ disappears to the benefit of a broad feature with complicated vibrational structure (Fig. 5 and S25 $\ddagger$ ). The luminescence decay in frozen solution could be fitted with a biexponential model resulting in an average lifetime of $1.8 \pm 0.1 \mu$ (Table 3 and S $2 \ddagger$ ). These data are consistent with this band corresponding to emission from the triplet state, with the 0 -phonon component located at $403 \mathrm{~nm}\left(24815 \mathrm{~cm}^{-1}\right)$.

The energy of the lowest lying excited f-level of $\operatorname{Gd}\left(\right.$ III) $\left({ }^{6} \mathrm{P}_{7 / 2}\right)$ is $32150 \mathrm{~cm}^{-1}$, therefore at higher energy than the highest component of the ligand states so that negligible ligand-tometal energy transfer is expected. The complex is then useful for identifying the singlet and triplet levels of the coordinated ligands. In toluene at room temperature (Fig. S25 $\ddagger$ ), 1-Gd has excitation and emission envelopes with maxima close to those recorded for the ligand, although their components have different intensities. In particular, the component at 317 nm ( $31545 \mathrm{~cm}^{-1}$ ), assigned to the 0-phonon transition of the lowest lying singlet state ${ }^{1} \pi \pi^{*}$ has a much larger intensity. As for the

| Table 3 <br> Gd), <br> 3- $-\mathrm{Pr}^{\mathrm{MeCN}}$ <br> and <br> ane |
| :--- | :--- | :--- | :--- | :--- |
| 4- $\mathrm{Pr}^{\mathrm{Ph}}$ in toluene frozen solutions $(77 \mathrm{~K})$ |

${ }^{a}$ Biexponential decay, amplitude average lifetime: $\tau_{\mathrm{av}}=\Sigma a_{\mathrm{i}} \tau_{\mathrm{i}}$.
ligand, in frozen solution this band disappears and a structured feature emerges with well resolved vibrational components; the 0 -phonon component is identified at $401 \mathrm{~nm}\left(24940 \mathrm{~cm}^{-1}\right)$. The average lifetime, $0.90 \pm 0.2 \mu \mathrm{~s}$ is compatible with the assignment to ${ }^{3} \pi \pi^{*}$ emission, and shorter than the ligand alone average lifetime due to the paramagnetic effect of $\operatorname{Gd}(\mathrm{III})$.

Metal centered luminescence of 1- $\mathbf{L n}(\mathbf{L n}=\mathbf{C e}, \mathbf{P r}), 3-\mathrm{Pr}^{\mathrm{MeCN}}$ and $4-\mathrm{Pr}^{\mathrm{Ph}}$
$\operatorname{Pr}($ Iv $)$ having the same electronic configuration than $\mathrm{Ce}(\mathrm{III}), 4 \mathrm{f}^{1}$, we set out to identify transitions specific to this configuration, for instance d-f transitions that are well known for Ce(III). We therefore compare the luminescence spectra obtained for the $\operatorname{Pr}($ Iv $)$ complexes with those of the $\mathrm{Ce}($ III $)$ and $\operatorname{Pr}($ III $)$ complexes. 3$\mathbf{P r}^{\mathbf{M e C N}}$ being not stable enough in toluene at room temperature the following discussion mainly involves spectra recorded at 77 K, see Fig. 5, S26-S29 and Tables 3 and S2 $\ddagger$ list the corresponding lifetimes.

Upon excitation in the ligand levels at 276 nm , the emission spectrum of 1-Ce (Fig. S26 $\ddagger$ ) features a weak emission band at 360 nm and an intense, well-structured band between 400 and 535 nm with a vibrational progression of $\sim 1500 \mathrm{~cm}^{-1}$; the lifetime determined by monitoring the 410 nm component, 1.5 $\mu \mathrm{s}$, is in line with a ${ }^{3} \pi \pi^{*}$ emission. Excitation at 354 nm results in a composite spectrum with the same structured band featuring a 0-phonon transition at $395 \mathrm{~nm}\left(25320 \mathrm{~cm}^{-1}\right)$ superimposed onto a broad emission centered around 471 nm . We assign this broad band to d-f transitions of Ce(iII) which are the basis of its spectroscopy with tuneable bands ranging from the blue to the red. ${ }^{34,16 a, 15 c, 15 g}$

The $\operatorname{Pr}($ III $)$ ion has two emitting levels, ${ }^{3} \mathrm{P}_{0}\left(21390 \mathrm{~cm}^{-1}\right)$ and ${ }^{1} \mathrm{D}_{2}\left(17335 \mathrm{~cm}^{-1}\right)$, that generate blue and red lines. ${ }^{35}$ In 1-Pr, the triplet state energy level is higher than that of the ${ }^{3} \mathrm{P}_{0}$ level, so that both emitting levels might be populated. However, as shown on Fig. S28, $\ddagger$ upon excitation at 352 nm , a spectrum identical to that of 1-Ce is obtained, corresponding again to ${ }^{3} \pi \pi^{*}$ emission from the ligand. No f-f transitions specific to $\operatorname{Pr}($ III $)$ are detected, which implies negligible energy transfer from the ligand and/or efficient back transfer and quenching. Indeed, it is known that $\operatorname{Pr}($ III $)$ complexes tend to have very low quantum yields. ${ }^{36}$

Upon excitation at 307 or $365 \mathrm{~nm}, 3-\mathbf{P r}^{\mathbf{M e C N}}$ (Fig. 5 and S29 $\ddagger$ ) exhibits one broad and symmetrical emission band centered at 611 nm as well as a very weak band at 500 nm and a shoulder at $\sim 550 \mathrm{~nm}$. The luminescence decay at 611 nm can be fitted with a single exponential, leading to a lifetime of $3.4 \mu \mathrm{~s}$. This differs markedly from the emission spectrum of 1-Pr and, as anticipated, the excitation spectrum is also different with, in addition to the ligand-centered transitions, a broad band around 350 nm corresponding to the LMCT identified in the absorption spectrum. These spectra represent the unique signature of the tetravalent complex.

The luminescence spectrum of $\mathbf{4}-\mathbf{P r}^{\mathbf{P h}}$ is similar to the one of $3-\mathbf{P r}^{\mathbf{M e C N}}$ (Fig. S29 $\ddagger$ ) with a maximum at 608 nm , except that 4$\mathbf{P r}^{\mathbf{P h}}$ is much less luminescent and that the two weak features at 500 and 545 nm are proportionally more intense. No residual


Fig. 5 Excitation and emission spectra of $\mathrm{KOSiPh}_{3}, 1-\mathrm{Ln}(\mathrm{Ln}=\mathrm{Gd}, \mathrm{Ce}, \mathrm{Pr}), 3-\mathrm{Pr}^{\mathrm{MecN}}$ and $4-\mathrm{Pr}^{\mathrm{Ph}}$ in 5 mM toluene solution at 77 K . Spectra are normalized and their intensities cannot be directly compared.
signal attributed to the triphenylphosphine oxide was detected even when both ligands were excited at 365 nm . The average lifetime corresponding to the 608 nm band is $3.45 \mu \mathrm{~s}$ and its onset is estimated to be at $19000 \mathrm{~cm}^{-1}$. The origin of the broad emission band of the $\operatorname{Pr}(\mathrm{rv})$ complexes is discussed in the section below.

## Electronic structure of the $\operatorname{Pr}(\mathrm{Iv})$ complexes

The $\approx 600 \mathrm{~nm}$ emission of the two $\operatorname{Pr}(\mathrm{Iv})$ complexes, could originate either from d-f transitions red-shifted with respect to 1-Ce or from emission from the ${ }^{3} \pi \pi^{*}$ state of the coordinated ligand. Although in a few cases longer lifetimes have been reported, ${ }^{37}$ lifetimes of d-f transitions are usually in the ns range. The measured decays for the $\operatorname{Pr}(\mathrm{rv})$ complexes are in the $\mu \mathrm{s}$ range, which could favour the second explanation. In order to substantiate this point, we have performed calculations at the DFT level (B3PW91 functional) for both $\mathrm{Ce}($ (ii) and $\operatorname{Pr}(\mathrm{Iv})$ complexes. In order to look for the d-f excitation, UV-visible spectra were computed using the TDDFT method. As expected for $\mathrm{Ce}($ (III), the d-f excitation appears as the first significant absorption band whereas no trace of such an absorption was found for $\operatorname{Pr}(\mathrm{iv})$ (see ESI! $\ddagger$ ).

Therefore, the hypothesis of an emission derived from a d-f transition in the $\operatorname{Pr}(\mathrm{rv})$ complex is quite unlikely and we thus investigated the possibility of an emission from the ligand
triplet state. In order to do so, both the ground state and an excited state were optimized. To make sure that the excitation arises from the ligand, a doublet and a quartet spin states were computed for the two complexes and we verified that in both cases the quartet excited state is located on the ligand. One outcome is that the quartet state of 1-Ce is calculated as lying at $27420 \mathrm{~cm}^{-1}$, well in line with the observed value of $24630 \mathrm{~cm}^{-1}$. The corresponding calculated level for $\operatorname{Pr}(\mathrm{rv})$ is largely red shifted by $12100 \mathrm{~cm}^{-1}$; the shift observed


Fig. 6 Partial electronic diagram for $1-\mathrm{Ce}, 3-\mathrm{Pr}^{\mathrm{MecN}}$, and $1-\mathrm{Gd}$. Calculated levels in blue, experimental 0 -phonon levels in red or rose when estimated from the onset of the emission band. Energies of the transitions in $\mathrm{cm}^{-1}$.
experimentally in going from $\mathrm{Ce}(\mathrm{III})$ to $\operatorname{Pr}(\mathrm{Iv})$, is smaller, but remains sizable at $\approx 5100 \mathrm{~cm}^{-1}$. The results are sketched in the electronic diagram of Fig. 6. It should be noted that the calculated absorption spectra (see ESI $\ddagger$ ) also show a LMCT transition at 506 nm but the associated energy of $2.44 \mathrm{eV}\left(19680 \mathrm{~cm}^{-1}\right)$ is larger than that of the triplet state at $1.9 \mathrm{eV}\left(15325 \mathrm{~cm}^{-1}, 625\right.$ nm ) so that we have ruled out a potential contribution to the emission spectrum.

## Conclusions

In conclusion, we have synthesised and crystallographically characterized tetrakis-complexes of several $\operatorname{Ln}(\mathrm{III})$ ions (Ce, Pr , $\mathrm{Nd}, \mathrm{Gd}, \mathrm{Tb}), 1-\mathrm{Ln}$, with the triphenylsiloxide ligand and the first example of a siloxide pentakis-complex of $\mathrm{Nd}(\mathrm{III}), 2-\mathrm{Nd}$.

Oxidation of the 1-Ce complex with silver tetraphenylborate allowed the synthesis of the $\mathrm{Ce}(\mathrm{rv})$ analogue $3-\mathrm{Ce}^{\mathrm{THF}}$. The oxidation of 1-Nd or $2-\mathrm{Nd}$ to yield $\mathrm{Nd}(\mathrm{Iv})$ was proven impossible, but the oxidation of $\mathbf{1 - P r}$ and $1-\mathbf{T b}$ with the strong oxidizing agent "magic blue" allowed the synthesis of the $\operatorname{Ln}(\mathrm{iv})$ complexes $\mathbf{3 - T b}{ }^{\mathbf{M e C N}}$ and $3-\mathbf{P r}^{\mathbf{M e C N}}$. The two molecules of bound
 replaced by THF or phosphinoxides $\mathrm{OPR}_{3}(\mathrm{R}=\mathrm{Et}, \mathrm{Ph})$ leading to isolation of $3-\mathbf{T b}^{\mathbf{T H F}}$ and the phosphinoxide adducts $\left[\mathrm{Tb}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}\left(\mathrm{OPR}_{3}\right)\right], 4-\mathrm{Tb}^{\mathrm{R}}(\mathrm{R}=\mathrm{Et}, \mathrm{Ph})$ and $\left[\mathrm{Pr}^{\mathrm{IV}}\left(\mathrm{OSiPh}_{3}\right)_{4}(-\right.$ $\left.\left.\mathrm{OPPh}_{3}\right)(\mathrm{MeCN})\right], \mathbf{4}-\mathbf{P r}^{\mathbf{P h}}$. Complex $\mathbf{4}-\mathbf{P r}^{\mathbf{P h}}$ is the second ever reported example of a $\operatorname{Pr}(\mathrm{Iv})$ molecular complex and shows significantly increased solution stability compared to $3-\mathbf{P r}^{\mathbf{M e C N}}$ suggesting that the synthesis of other $\operatorname{Pr}(\mathrm{Iv})$ molecular complexes should be in reach. The infrared, UV-visible and luminescence spectra of the isolated complexes were measured and the assignment of the luminescence spectra was corroborated by computational studies. The luminescence spectra of $\operatorname{Gd}($ III $), \operatorname{Pr}$ (III) and Ce (III) all show emission from the coordinated ligand ${ }^{3} \pi \pi^{*}$ state which is only slightly displaced with respect to the free ligand. The $\mathrm{Ce}(\mathrm{III})$ ion also shows a broad emission assigned to d-f transition. The first luminescence spectrum of the $\operatorname{Pr}(\mathrm{Iv})$ was measured and shows a broad intense emission. On the basis of the computational studies and of the measured lifetime in the microsecond range the emission is assigned to the ${ }^{3} \pi \pi^{*}$ state of the coordinated ligand. The coordination of the ligands to the $\operatorname{Pr}(\mathrm{rv})$ ion results in an unprecedented large shift of the ligand triplet state probably due to an increased covalency in the $\operatorname{Ln}(\mathrm{Iv})-\mathrm{O}$ bonds compared with $\operatorname{Ln}(\mathrm{III})-\mathrm{O}$ bonds. The observed large Stokes' shift anticipates the possibility of applications of $\operatorname{Ln}(\mathrm{Iv})$ in material science.

## Author contributions

M. M. conceived the concepts and supervised the project. A. R. W. performed the majority of the experiments. A.-S. C. conceived and measured the luminescence experiments. A.-S. C. and J.-C. G. B. analysed the luminescence data. I. D. and L. M. carried out and analysed the computational data. F. F.-T. carried out and analysed the X-ray data. All authors contributed to the analysis and writing.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

We acknowledge support from the Swiss National Science Foundation and the Ecole Polytechnique Fédérale de Lausanne (EPFL). We thank Dr R. Scopelliti for important contributions to the X-ray single crystal structure analyses. L. M. is a senior member of the Institut Universitaire de France. CalMip is acknowledged for a generous grant of computing time.

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    $\ddagger$ Electronic supplementary information (ESI) available. CCDC 2104729-2104735. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d1sc05517h

