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Light-induced splitting of P–C bonds in a lanthanum(III) hemiphosphinal complex†

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Insertion of fluorenone into the lanthanum–phosphorus bond of terminal phosphido complexes $(\text{PN})_2\text{La}(\text{PHR})$ ($\text{R} = \text{Mes}$ (**1a**), $\text{R} = \text{Ph}$ (**1b**) and $\text{R} = {^t}\text{Bu}$ (**1c**)) results in the formation of light-sensitive hemiphosphinal complexes **2a–c**. Upon exposure to daylight, the P–C bonds in **2a–c** break resulting in the formation of diphosphines and a lanthanum bound fluorenonyl-radical. The scope and mechanism of this reaction are experimentally and theoretically evaluated.

Introduction

In the past decades, early transition metal and f-element phosphanido complexes have been found to adopt a rich and versatile insertion chemistry, which allows the construction of exotic and rare types of phosphorus functionalized molecules.^{1–4} From a thermodynamic point of view, the main driving force for this versatile insertion chemistry is the exchange of a rather weak M–P interaction for more ionic M–N and M–O bonds, alongside the formation of a new P–C bond.⁵ Thus, in the 1990s a large number of insertion reactions were reported by the groups of Hey-Hawkins and Stephan which included the build-up of phosphaguamidates,⁶ phosphathioureates⁷ and phosphaamidates^{8,9} at zirconium(IV). This chemistry also has been widely extended into the f-elements in the past decade.^{1–4,10–15} Furthermore, the insertion of carbodiimides, nitriles and iso(thio)cyanates, olefins or alkynes into M–P bonds (M = early transition metals or f-element) is also considered to be a fundamental step in metal-catalyzed hydrophosphin(yl)ation reactions.^{3,4,10,11,16–20} Despite this plethora of examples, the insertion chemistry of ketones, especially of fluorenone (and related diaryl ketones *e.g.* benzophenone) into M–P bonds, yielding hemiphosphinals, is still scarcely studied and so far only a handful of hemiphosphinal complexes have been reported (Fig. 1, left).^{13,21–26} This is surprising, since diarylketonates hold a variety of promising properties such as their easy reduction to form (stable) ketyl-radicals^{27–40} and to

undergo reductive coupling^{41,42} reactions or 1,2-migratory insertion reactions.^{13,43–46} This makes them very attractive synthons for organometallic chemists, which is further demonstrated by a recent example from Erker and co-workers.^{44,47} Using hemiphosphinal zirconium(IV) complexes, originally described by Stephan and co-workers²⁶ in 1996, they promoted new reactivity in FLP chemistry.^{44,47} Furthermore, when electron accepting ligands, such as bipyridine, are introduced, the Arnold group has recently shown, that terminal thorium phosphanido complexes can undergo reductive coupling of the phosphanido-ligands.⁴⁸

We have recently reported a versatile lanthanum anilidophosphine complex **I**⁴⁹ (Fig. 1, right) which readily undergoes salt metathesis reactions leading to highly functionalized metal complexes, such as the first stable 2-phosphaethynthio-

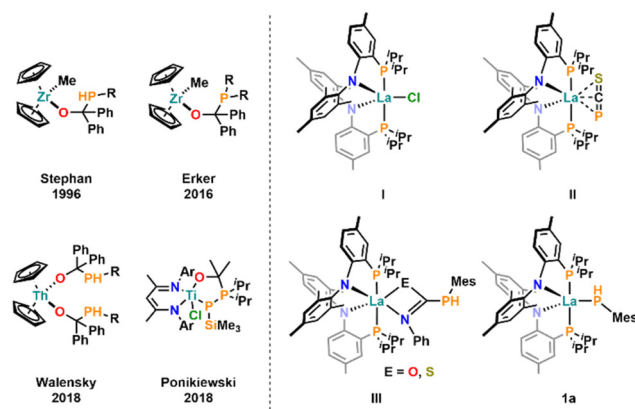


Fig. 1 Left: Selected examples of hemiphosphinal complexes reported in the literature.^{13,25,26,44} Right: Selection of anilidophosphine supported lanthanum complexes showing unexpected and versatile chemistry recently reported by our group.

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late ($[\text{SCP}]^-$) complex⁵⁰ **II** or the first terminal, primary phosphanido complex **1a** of the lanthanides.⁴⁹ The latter was found to be a promising candidate for the design of lanthanum phosphinidenes⁵¹ and readily undergoes insertion chemistry with *e.g.* iso(thio)cyanates to give phospha(thio)ureates of the general structure **III**.⁵² Given the previous mentioned propensity of electropositive metal-phosphorus insertion chemistry, and the lack of follow-up chemistry explored on hemiphosphinals, here we expand the reactivity of **1a** towards fluorenone. To our surprise and in contrast to a previously reported thorium hemiphosphinal complex by the Walensky group,¹³ the resulting hemiphosphinal is light sensitive leading to a P-C bond scission, forming diphosphine and a fluorenonyl radical stabilized on lanthanum(III).

Results and discussion

Reaction between fluorenone and **1a** in the absence of light resulted in the clean formation of the desired hemiphosphinal complex **2a** (Scheme 1). Successful formation of the complexes is evident by the shift of the resonance corresponding to the phosphanido ligand in the $^{31}\text{P}\{^1\text{H}\}$ NMR spectrum from $\delta = -36.4$ ppm in **1a** to $\delta = -32.1$ ppm in **2a** (Fig. S19†). The retention of the proton on the phosphorus atom is confirmed by ^1H NMR spectroscopy showing a doublet at 5.17 ppm ($^1J_{\text{PH}} = 210.6$ Hz). Additionally, the observation of a new doublet in the $^{13}\text{C}\{^1\text{H}\}$ NMR of **2a** at $\delta = 95.3$ ($^1J_{\text{CP}} = 12.3$ Hz) corresponding to the former fluorenone carbonyl carbon atom is indicative for the successful insertion into the La-P bond. X-ray diffraction studies (Fig. 2) unambiguously proved the insertion of the fluorenone moiety into the former La-P bond, forming a new La-O bond (La1–O1 2.1867(13) Å) and a new P-C bond (P70–C80 1.9163(19) Å). The distance of 4.462(2) Å between P70 and La1 proves that no interaction remains between the phosphorus and the lanthanum atom after the insertion of the fluorenone moiety. To our surprise, and in contrast to all previous reported insertion reactions of ketones into polarized M-P bonds,^{13,26} samples of **2a** were not stable towards ambient light (Fig. 3a). Keeping NMR samples exposed to daylight resulted in a colour change from orange to dark red (Fig. 3b). Along with the colour change, the broad hemiphosphinal ^{31}P NMR resonance at -32.1 ppm vanishes

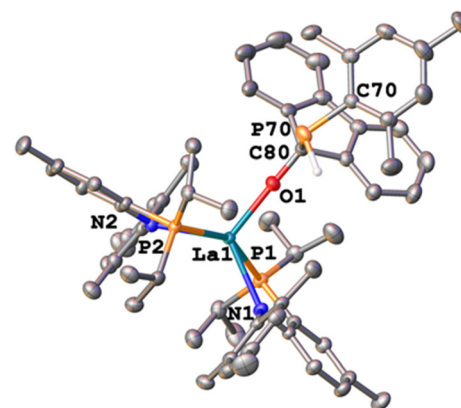
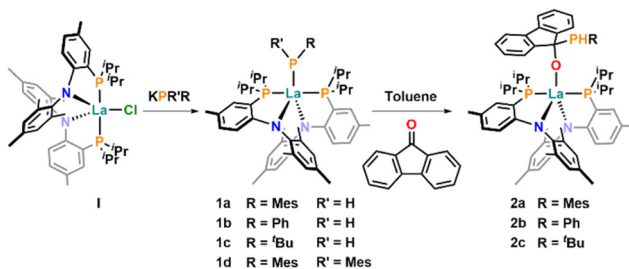


Fig. 2 Molecular structure of the hemiphosphinal complex **2a**. Thermal ellipsoids are shown at a probability level of 50%. Hydrogen atoms (except for H70 on P70) and lattice solvent molecules have been omitted for clarity. Selected bond parameters of the hemiphosphinal moiety: **2a**: La1–O1 2.1867(13) Å, O1–C80 1.401(2) Å, C80–P70 1.9163(19) Å, La1–O1–C80 177.79(12)°, O1–C80–P70 103.22(12)°.



Scheme 1 Synthesis of the phosphanido complexes **1a–d** and the corresponding hemiphosphinal complexes **2a–2c**.

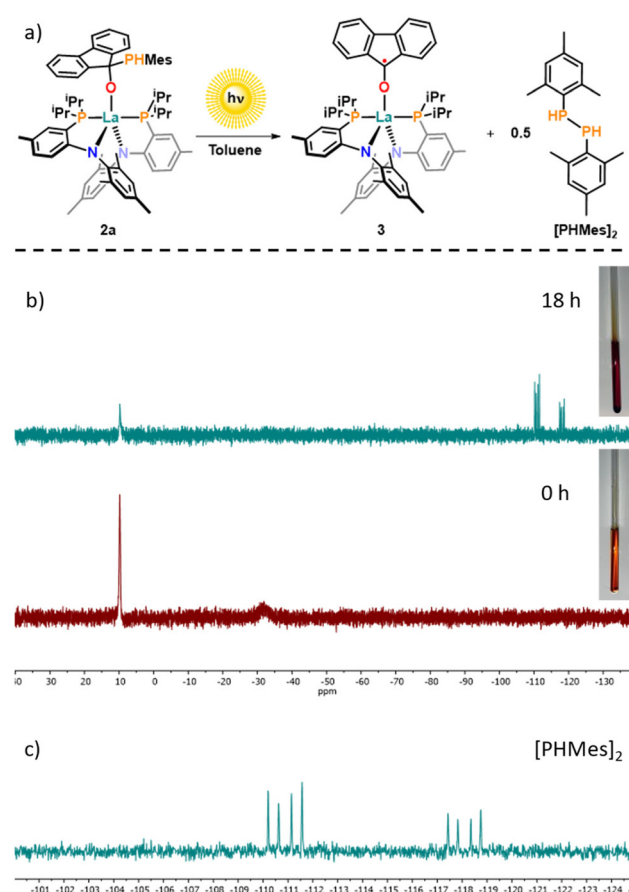


Fig. 3 (a) Net reaction scheme of the photolysis of complex **2a**. (The radical is highlighted in red) (b) ^{31}P NMR spectra of **2a** before and after irradiation with sunlight for 18 h together with the visual colour change observed. (c) Multiplet of [PHMes]₂.



and two new multiplets at -110.8 ppm and -118.2 ppm appear, which correspond to dimesityldiphosphine [PHMes_2] (**Fig. 3b** and **c**).^{48,53}

Intrigued by the distinct colour change from orange to intense red, the disappearance of the PN resonances in the ^{31}P NMR spectra over time, and the fact that diarylketones, especially benzophenone, can be photo-reduced using visible light in the presence of *tert*-butanolate,⁵⁴ we investigated if the reaction might also proceed *via* a fluorenonyl-radical **3**. Indeed, when fresh samples of **2a** were irradiated with light inside the EPR spectrometer, we saw the formation of radical species **3** (**Fig. 4**, top). Simulation of the radical forming is in good agreement with a fluorenyl radical being bound to a lanthanum center with $g_{\text{iso}} = 2.0026$ and hyperfine coupling to both the lanthanum center ($a_{\text{iso}} = 1.7$ MHz) and the four distinct fluorenone hydrogens ($a_{\text{iso}} = 10.1, 8.11, 3.04, 2.18$ MHz) (**Fig. 4**). Complete photolysis of a separate sample of **2a** using a high-intensity photoreactor (5000 K cold-white LED, 36 min, **Fig. S48†**) and subsequent recrystallisation of the reaction mixture by vapour diffusion of hexane into a benzene solution resulted in the formation of red blocks of **3**. EPR of these red blocks confirmed their identity to be the same radical as observed after *in situ* photolysis of complex **2a** in the EPR tube (**Fig. 4**, middle). NMR Evans Method revealed a magnetic moment of $1.42 \mu_{\text{B}}$ which agrees with the presence of one unpaired electron. X-ray diffraction analysis of the new complex **3** unambiguously confirmed the presence of a fluorenonyl radical being bound to the lanthanum center (**Fig. 5**, middle). This is clearly observable from three bond parameters: (I) The La1–O1 bond distances which increases from $2.1867(13)$ Å in **2a** to $2.261(4)$ Å in **3**; (II) the decrease of the O1–C80 bond distance from $1.401(2)$ Å in **2a** to $1.303(6)$ Å in **3** and the change of the La1–O1–C80 bond angle, moving from $177.79(12)^\circ$ to 180.0° in **2a** and **3** respectively. All these parameters indicate a major change of the electronic situation within the fluorenone-moiety affecting the electron density within its carbonyl functionality and are in line with previously reported lanthanide fluorenonyl radicals,^{27–34} e.g. supported by samarium^{9,28,40} or ytterbium.³⁸

To set these values into context (and to rule out the influence of potential steric and crystal packing effects) we synthesized the pure fluorenolate complex **4** and a corresponding cationic fluorenone complex **5**. The “isostructural” series of complexes **3–5** allows the direct comparison between the bond metrics in the fluorenolate, the fluorenonyl-radical and the neutral fluorenone moiety. Synthesis of the fluorenolate complex **4** was achieved by salt metathesis between the chloride precursor **I** and lithium fluorenolate (Scheme 2). The product is isolated in 76% yield as a faint yellow powder. Characteristic NMR features that indicate the successful formation of the fluorenol complex are (I) the presence of the fluorenolate CH resonance at 6.10 ppm in its ^1H NMR spectrum, together with the characteristic fluorenolate multiplets between 7.58 – 6.80 ppm (II) the shift of the characteristic aryl-H of the PN ligand from 5.65 ppm in complex **I** to 5.75 ppm in **4**. Notably in this case the shift of the ^{31}P NMR resonance of

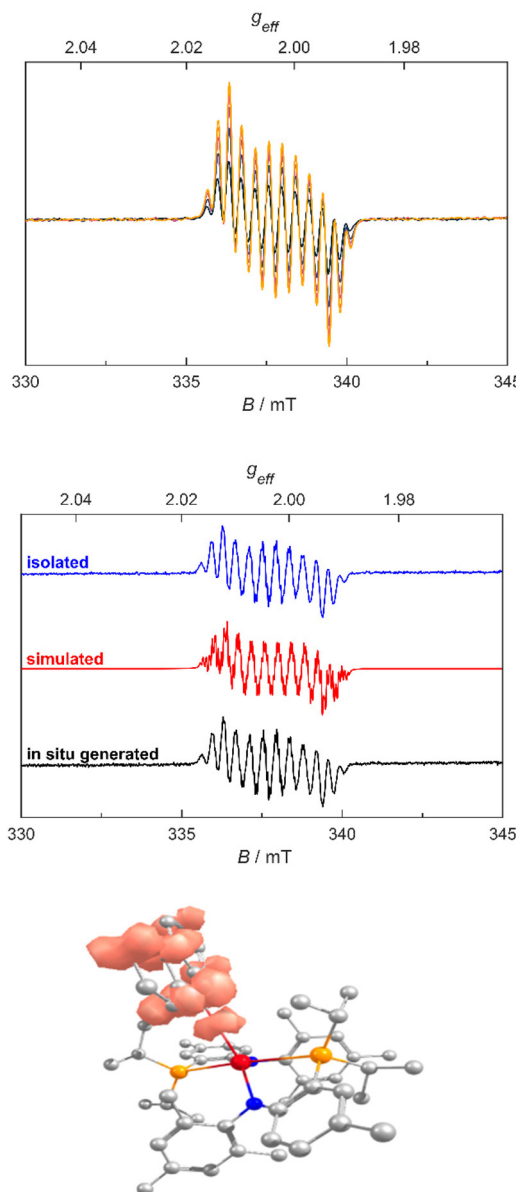


Fig. 4 Top: EPR spectra of the *in situ* irradiated sample of complex **2a**, each shown step corresponds to a 40 s irradiation with 0.1% lamp power. Please note that the initial radical concentration at 0 minutes is not 0 as the sample already decomposes when being transferred to the EPR tube even in the dark. Middle: highly resolved EPR of complex **3** from *in situ* photolysis (black), isolated radical (blue) and simulation (red). The intensities of these spectra have been normalized for clarity. Bottom: calculated Spin density plot of the fluorenyl radical complex **3**.

the PN ligand cannot be viewed as indicative for the formation of the new species **4** as it shifts from 9.20 ppm in **I** to 9.17 ppm in **4**. X-ray quality crystals could be obtained from a concentrated toluene solution of **4** within 3 h at room temperature. Unfortunately, the synthesis of the desired cationic complex **5** proved to be more tricky and despite various attempts (halide abstraction was attempted using AgBArF_{20} , AgBArF_{24} , NaBArF_{24} and $[(\text{Et}_3\text{Si})_2\text{H}][\text{BArF}_{20}]$ in the presence of complex **I** and fluorenone) no defined cationic mono-fluore-



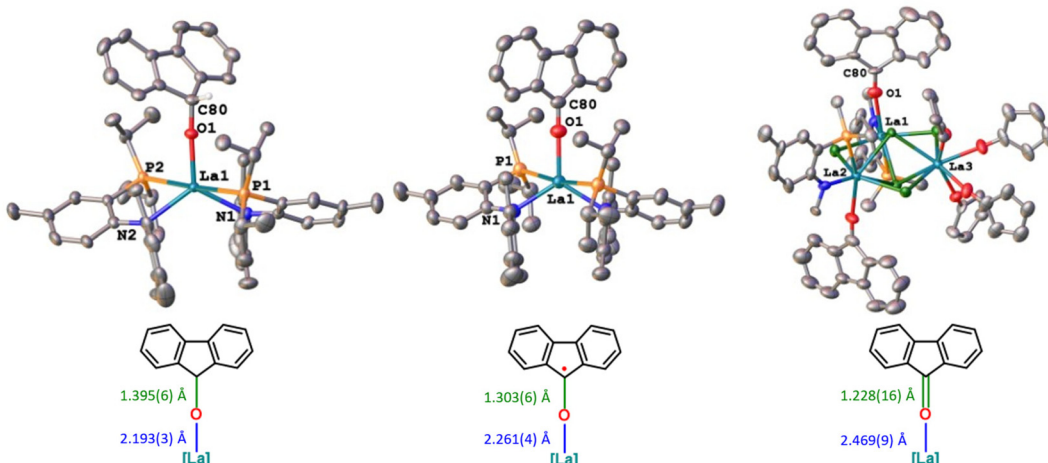
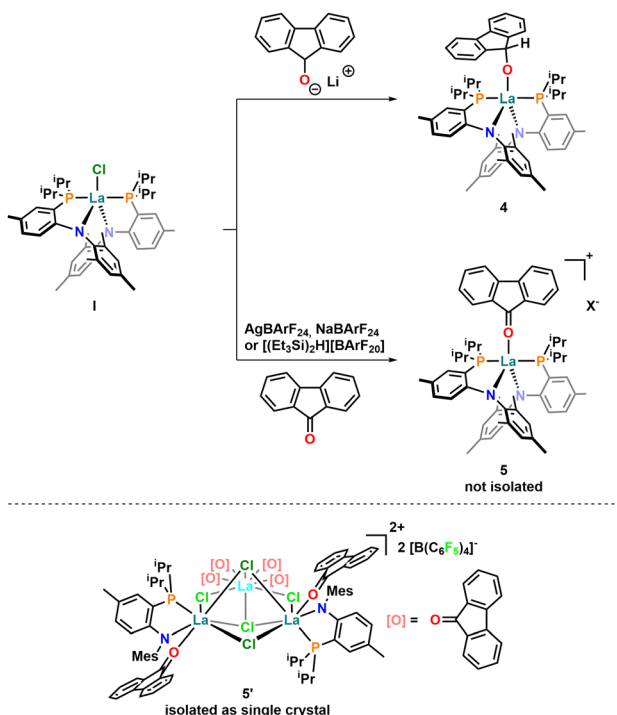


Fig. 5 Molecular structures of the fluorenolate complex **4** (left), the fluorenonyl radical complex **3** (middle) and the dicationic fluorenone cluster **5'** (right). Thermal ellipsoids are shown at a probability level of 50%. Hydrogen atoms (except for the H at C80 in **4**), counter ions and lattice solvent molecules have been omitted for clarity. In the structure of **5'** the PN ligands as well as the majority of the fluorenone ligands have been truncated for clarity. The La1–O1 and the O1–C80 bond distances have been depicted further for clarity and comparison.



Scheme 2 Synthesis of the fluorenolate complex **4** and attempted synthesis of the cationic fluorenone complex **5** as structural comparisons for the radical complex **3**. Please note, that complex **5** has never been isolated and only complex **5'** (X-ray structure see Fig. 5, right) was accessible in single crystal yield.

none complex **5** could be isolated. However, using $[(\text{Et}_3\text{Si})_2\text{H}][\text{BArF}_{20}]$ as a chloride abstraction reagent we observed the formation of the dicationic tri-lanthanum fluorenone cluster **5'** after crystallisation from dichloromethane layered by pentane at -40°C . Although this cluster is not the

desired complex **5** it still holds valuable structural information concerning the interactions between lanthanum and neutral fluorenone. Comparison between the structures of **3**, **4**, and **5'** shows that (as expected) the La1–O1 distance increases from 2.193(3) Å in **4** to 2.261(4) Å in **3** culminating in the longest 2.469(9) Å in **5'**. At the same time the O1–C80 distance decreases from 1.395(6) Å in **4** to 1.303(6) Å in **3** to 1.228(16) Å in **5'** which agrees with an increasing carbonyl double bond character between O1 and C80 (Fig. 5).

To gain further insights into the reaction (Scheme 1) its mechanism was further elucidated by computational methods (DFT, B3PW91, see ESI† for more information) for the hemiphosphinal complex **2a**. For simplification, we modelled the insertion reaction the C=O double bond of benzophenone into the La–P bond, which is predicted to be kinetically facile (barrier of 1.7 kcal mol⁻¹) and thermodynamically favourable (-21.6 kcal mol⁻¹) (Fig. S74†).⁵⁵ Therefore, the irradiation used experimentally is needed in order to break the P–C bond. This was investigated by computing the TDDFT spectrum of **2a** (Fig. S69†), showing a transition at 602 nm that involves a HOMO–3 to LUMO transition. The HOMO–3 displays a bonding P–C interaction while the LUMO has primarily an antibonding P–C one so that populating the latter would imply breaking of the P–C bond (Fig. 6).

Intrigued by this result, we further investigated the possibility of a homolytic cleavage of the P–C bond, and if the second reaction product corresponds to a mesitylphosphinyl radical that quickly recombines to form the observed dimesityldiphosphine. However, performing the P–C bond photolysis in the presence of various radical scavengers (TEMPO, styrene, radical clocks *etc.*) gave no indication of the presence of free phosphinyl radicals. This was further confirmed by X-ray photolysis experiments of single crystals of **2a** showing no signs of decomposition or changes along the C80–P70 bond (Table S2†). Furthermore, no EPR signals were observed if the

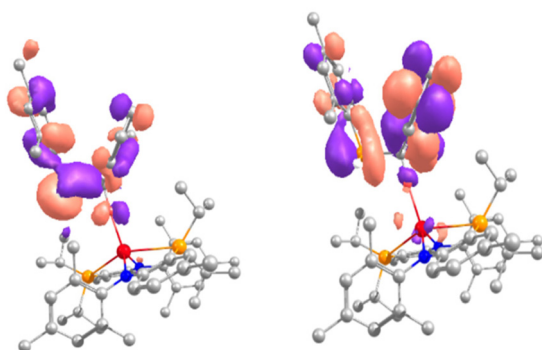


Fig. 6 HOMO-3 and LUMO of complex **2a** calculated by DFT, irradiation at 602 nm induces a transition from a binding P–C bond orbital (HOMO-3) into the antibonding P–C bond orbital (LUMO).

EPR photolysis experiments were performed in a frozen solution of **2a**. Since these experiments suggest that the reaction does not proceed *via* a monomolecular process, an alternative pathway involving a concerted mechanism for the P–P dimerization and the P–C bond splitting seems to be more likely (Scheme 3).

To figure out whether this decomposition reaction is unique to the primary mesitylphosphine moiety, we prepared the phenylphosphanido, the *tert*-butylphosphanido and the dimesitylphosphanido complexes **1b–d**. The complexes of the

primary phosphines **1b** and **1c** were readily accessible in good yields (60 and 82% respectively), following a salt metathesis protocol between complex **I** and the corresponding potassium phosphanides ($\text{KPHR} = \text{R} = \text{Ph}$ or ^tBu). Formation of the desired complexes **1b** and **1c** is indicated by the ^1H and ^{31}P NMR spectra. The ^1H NMR spectra show the characteristic doublets of the primary phosphanido ligand at 4.06 ppm ($^1J_{\text{PH}} = 194.8$ Hz) and 3.47 ppm ($^1J_{\text{PH}} = 200.7$ Hz) for **1b** and **1c**, respectively. These $^1J_{\text{PH}}$ coupling constants can also be extracted from the ^{31}P NMR spectra showing distinct doublets at -10.2 ppm ($^1J_{\text{PH}} = 194.8$ Hz, Fig. S5†) and 97.5 ppm ($^1J_{\text{PH}} = 199.3$ Hz, Fig. S11†) for **1b** and **1c** respectively. Furthermore, the characteristic aryl proton of the PN ligand is shifted from 5.65 ppm in complex **I** to 5.67 ppm and 5.68 ppm in the ^1H NMR spectra of **1b** and **1c**. Accordingly, the resonances of the PN-ligands phosphorus atom in the $^{31}\text{P}\{^1\text{H}\}$ NMR spectra are shifted from 9.2 ppm for complex **I** to 10.4 ppm and 12.3 ppm for **1b** and **1c** respectively.

On the other hand, synthesis of the dimesitylphosphanide complex **1d** was found to be rather messy, producing multiple inseparable products. Apart from a few single crystals (yield less than 1%, structure see Fig. 7) no pure bulk material could be obtained, wherefore no further reactivity (insertion chemistry) with **1d** could be investigated. Single crystals could be obtained by storing a concentrated hexane solution for 5 weeks at -40 °C for complex **1b** or at room temperature over-

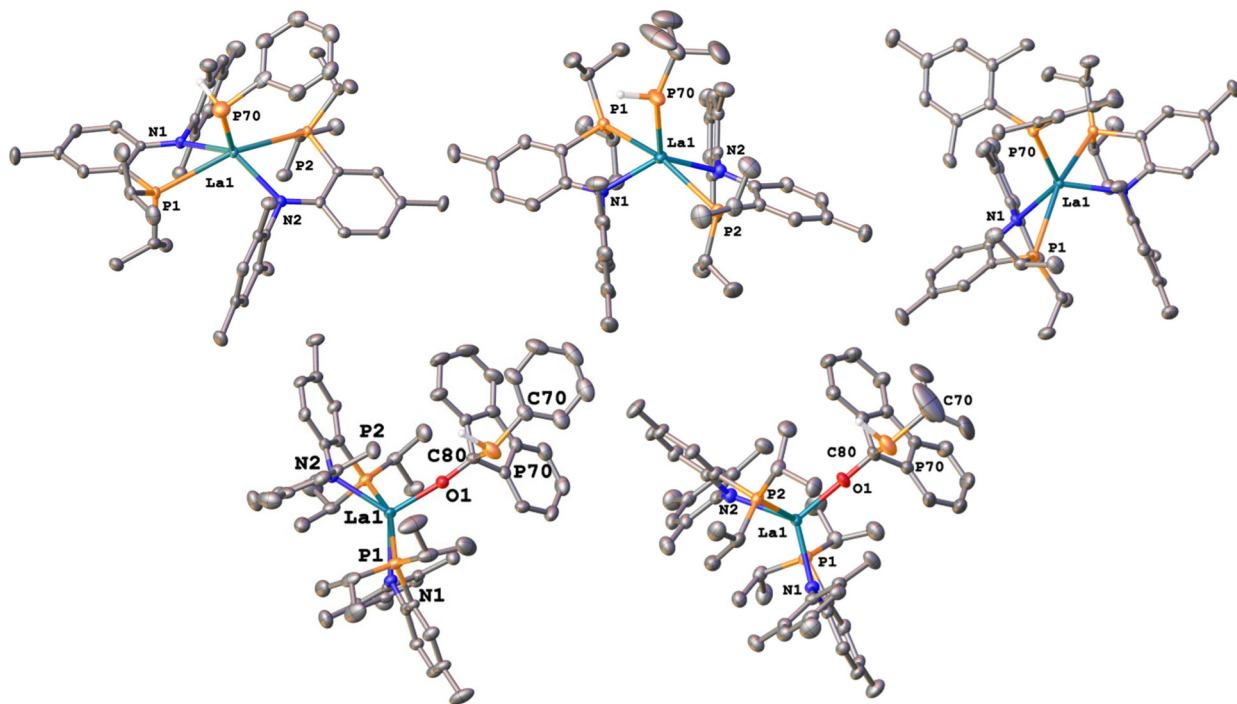


Fig. 7 Molecular structure of the phosphanido complexes **1b–d** (top from left to right). Bottom row shows the insertion complexes **2b** and **2c** after fluorenone insertion into the corresponding phosphanido complexes. Thermal ellipsoids are shown at a probability level of 50%. Hydrogen atoms (except for H70 on P70) and lattice solvent molecules have been omitted for clarity. Selected bond parameters: **1b**: La1–P70 3.009(1) Å, **1c**: La1–P70 2.8364(6) Å, **1d**: La1–P70 2.9216(14) Å. **2b**: La1–O1 2.191(3) Å, O1–C80 1.402(5) Å, C80–P70 1.924(4) Å, La1–O1–C80 167.8(3)°, O1–C80–P70 107.4(3)°. **2c**: La1–O1 2.1776(14) Å, O1–C80 1.413(2) Å, C80–P70 1.895(2) Å, La1–O1–C80 176.90(15)°, O1–C80–P70 103.35(14)°.



night for complex **1c** or by storing a concentrated diethyl ether solution at -40°C over three days for complex **1d**. Structural investigation on the phosphanido complexes **1b-d** (Fig. 7, top

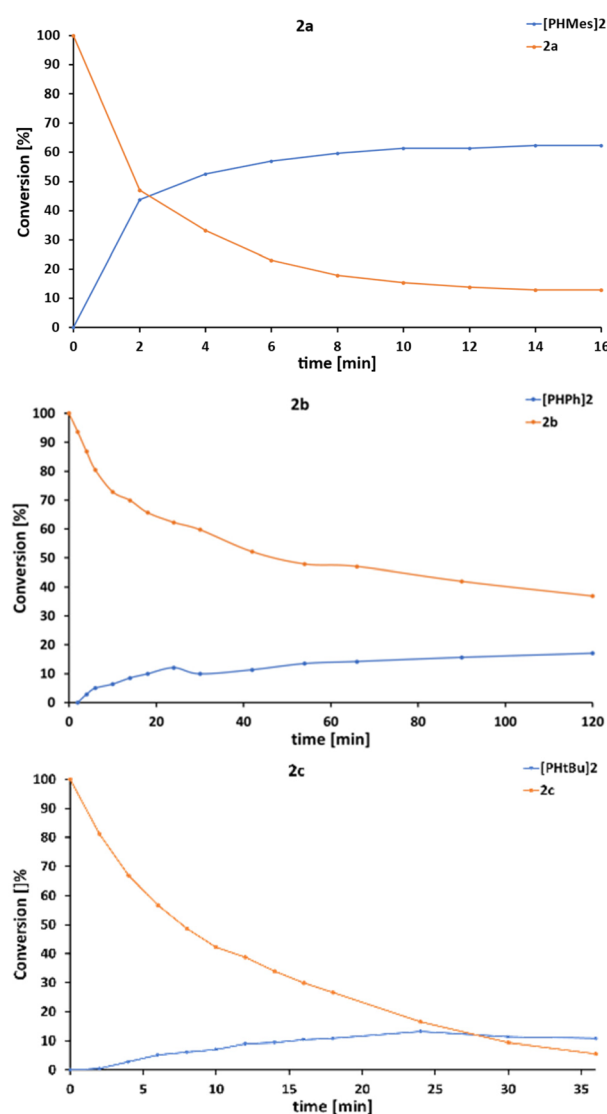
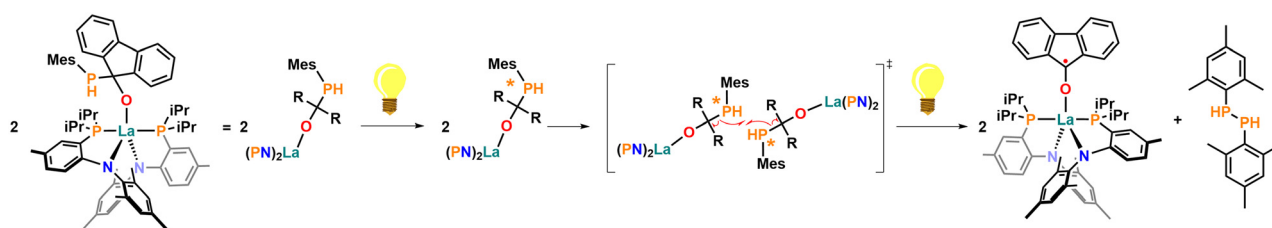


Fig. 8 Time-conversion-plots of the light-promoted decomposition of **2a**, **2b** and **2c**. The blue curves show the amount of diphosphine being formed, while the orange lines show the consumption of the starting hemiphosphinal complex. Conversions were determined via ^{31}P NMR spectroscopy using PPh_3 as an internal standard.

row) revealed that the La-P bond distance is strongly depending on the substitution on the phosphorus donor and a La1-P70 distance of $3.009(1)$ Å was measured for **1b** ($3.053(1)$ Å in **1a**), which shortened to $2.8364(6)$ Å in **1c**. Interestingly, the steric congestion/bulk of the phosphanido ligand seems to play a minor role over the electronic properties of the phosphanido ligand towards the La1-P70 distance, since with $2.9216(14)$ Å this distance in **1d** is shorter than in the primary phosphanido complexes **1a** and **1b**. Insertion reactions with fluorenone proceeded similarly to the insertion forming **2a** and the complexes **2b** and **2c** were obtained as orange solids in yields of 86% and 68%. The shift of the PH-resonance in the ^{31}P NMR spectra is again the most indicative feature for the formation of the complexes **2b** and **2c**, similar to **2a**. Conversion from **1b** to **2b** shifts this phosphorus resonance from -10.2 ppm to 5.46 ppm ($^1J_{\text{PH}} = 196.9$ Hz) while **1c** to **2c** results in a shift from 97.5 ppm to 23.6 ppm ($^1J_{\text{PH}} = 191.6$ Hz). Similar to **2a**, **2b** and **2c** were also highly sensitive towards light and a fast colour change was observed when samples were left exposed to daylight. Crystals suitable for X-ray diffraction studies were grown from concentrated toluene solutions of the complexes at room temperature (Fig. 7, bottom row). The structural features of the complexes are similar to **2a** wherefore a detailed discussion is omitted at this place. Further structural information on the complexes can be extracted from the ESI (Tables S1 and S2†).

To determine the influence of the phosphine, we monitored the reaction over time using triphenylphosphine (PPh_3) as an internal standard (Fig. 8). Firstly, this reveals that the reaction is the fastest for complex **2a**. Additionally it shows that the starting complexes are consumed at a much faster rate than the diphosphines are formed and also that the diphosphine concentration does not increase further once a threshold level is reached, which appears to be distinctive for the corresponding diphosphines. While in the case of **2a**, about 60% conversion to dimesityldiphosphine can be reached after 16 minutes, for **2b** and **2c** only 16% and 10% conversions are reached within 120 and 36 min respectively. This further shows that reaction for **2b** is much slower compared to the more electron rich phosphines in **2a** and **2c**. Notably, in the case of **2c**, the concentration of the starting material is further decreasing even after the diphosphine concentration has reached its conversion. This indicates that another decomposition path for **2c** might be possible, which is in agreement



Scheme 3 Postulated mechanism for the photoinduced P-C bond cleavage observed in complex **2a** via a concerted (homolytic) C-P bond scission / P-P bond formation mechanism

with the fact that the ^{31}P NMRs of **2c** show substantial formation of HPN and other phosphorous species (Fig. S50†). Thus, we deduce that aryl phosphines seem to react much more cleanly. To further elucidate if our postulated mechanism from Scheme 3 might be valid, we further performed scrambling experiments using equimolar mixtures of **2a** and **2b**. Indeed, during irradiation we saw the formation of three diphosphine species, being the two symmetrical dimesityldiphosphine ($[\text{PHMes}]_2$) and diphenyldiphosphine ($[\text{PhPh}]_2$), as well as the mixed phenyl-mesityl diphosphine (PhPH-PHMe) (Fig. S51 and S52†). Together with the fact, that no free phosphanyl radicals could be trapped and evidenced during the reaction (*vide supra*) the observation of the scrambling product **PhPh-PHMe** further supports our theory of a concerted reaction mechanism.

Conclusion

We have reported the insertion reaction of fluorenone into labile La-P bonds in **1a–c**, giving access to hemiphosphinal complexes **2a–c**. These complexes are sensitive to light, decomposing to a fluorenonyl radical complex **3** and the corresponding diphosphines $[\text{PHMe}]_2$, $[\text{PhPh}]_2$ and $[\text{Ph}^t\text{Bu}]_2$. The former can be detected by EPR spectroscopy and can even be isolated in (single) crystalline form. On the phosphine side, although calculations suggest a direct P-C bond splitting no free phosphanyl radicals could be detected. Thus, we propose a concerted P-C bond scission / P-P bond formation mechanism which is also indicated experimentally *via* scrambling experiments (Scheme 3).

In summary we have reported a rare case of a light sensitive hemiphosphinal complex, which easily decomposes to give access to synthetically challenging diphosphines. Given the fact, that similar decomposition reactions have not been observed for transition metal^{8,9,21,22,26} and actinide¹³ based hemiphosphinal complexes we believe that this reaction opens new venues in the use of lanthanides in (phosphor-)organic chemistry. Additionally, it gives facile access to multi-spin systems with fluorenone radicals attached to optically and magnetically active lanthanides.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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