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Photolysis-driven bond activation by thorium and uranium tetraosmate polyhydride complexes†

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Transition metal multimetallic complexes have seen intense study due to their unique bonding and potential for cooperative reactivity, but actinide–transition metal (An–TM) species are far less understood. We have synthesized uranium– and thorium–osmium heterometallic polyhydride complexes in order to study An–Os bonding and investigate the reactivity of An–Os interactions. Computational studies suggest the presence of a significant bonding interaction between the actinide center and the four coordinated osmium centers supported by bridging hydrides. Upon photolysis, these complexes undergo intramolecular C–H activation with the formation of an Os–Os bond, while the thorium complex may activate an additional C–H bond of the benzene solvent, resulting in a $\mu\text{-}\eta^1,\eta^1$ phenyl ligand across one Th–Os interaction.

Introduction

Metal hydrides have seen extensive interest due to their importance in processes such as catalytic hydrofunctionalization of olefins,¹ reduction of organic substrates and small molecules,^{2,3} and electrocatalytic redox reactions.⁴ While d-block metal hydrides have been the focus of the majority of these investigations, it was soon discovered that f-block hydrides exhibit unique reactivity that, in some cases (particularly the hydrogenation and polymerization of unsaturated organic substrates), render them even better catalysts than their transition metal (TM) counterparts.⁵ Furthermore, heterometallic hydride complexes—derived from the combination of f-block and transition metals—have exciting potential for cooperative reactivity utilizing the unique properties of each metal center; they also offer the possibility of providing fundamental insight into f-block–TM bonding. Bridging hydrides are well suited to support these interactions, as their minimal steric profile allows for close metal–metal contact. In addition, hydride elimination to form metal–metal bonds is

well-precedented in the transition metal literature, offering a potential route to unsupported metal–metal bonds.⁶ Lastly, heterometallic actinide hydrides are of wider interest due to their possible superconductive properties at high pressures, and intriguingly, for their potential as hydrogen storage materials.⁷

Significant strides have been made in the synthesis of rare earth/transition metal polyhydrides since the first example, a trimetallic Y_2Zr tetrahydride, was synthesized by Evans in 1984.⁸ In subsequent years, several groups have reported the successful syntheses of such complexes using alkane elimination, H_2 elimination, and salt metathesis.^{9–15} Hou has reported extensive studies on Y- and Lu–TM complexes, synthesizing polyhydride clusters with all of the 2nd and 3rd row transition metals between groups 6 and 9 (excluding Tc), several of which display hydrogen addition and release properties.^{16–22}

In contrast, while actinide (An) borohydride and aluminohydride chemistry is well-established,^{5,23–27} multimetallic hydride chemistry of the actinides is underdeveloped. Ephritikhine reported a series of U/Re complexes supported by three bridging hydrides, utilizing potassium rhenate salts to install the rhenium center on uranium pentamethylcyclopentadienyl (Cp^*) halide complexes.^{28–30} The U–Re distance in $[\text{K}(18\text{-crown-6})][(\text{Cp}^*)_2(\text{Cl})\text{U}(\mu\text{-H})_3\text{Re}(\text{H})_3(\text{PPh}_3)_2]$ was long (3.255(8) Å), leading the authors to conclude there was no direct U–Re interaction;²⁹ limited reactivity was observed with all U–Re species described.

Following our report of hydride-supported An–Al bonds (An = Th, U),²⁶ we have recently focused on applying this strategy to stabilize new An–TM interactions. Our efforts have resulted in the syntheses of several multimetallic actinide tetrairidate dodecahydride complexes, $\text{Th}\{[(\mu\text{-H})_2(\text{H})\text{IrCp}^*]_2[(\mu\text{-H})_3\text{IrCp}^*]_2\}$

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(A) and $U\{(\mu-H)_3IrCp^*\}_4$ (B), from the iridium polyhydride starting material $K[Cp^*IrH_3]$ and actinide halides (Scheme 1). These complexes featured the first reported computational evidence for An–Ir interaction.³¹ Given the efficacy of eliminating a bridging ligand to form An–Co bonds,³² we sought to engender An–TM bonds *via* dihydrogen elimination, but both A and B proved resistant to this process, either by thermolysis up to 80 °C, or *via* photolysis with 254 nm light.

We turned to the osmium counterpart, Cp^*OsH_5 , reported by Girolami, to pursue the synthesis of An–Os multimetallic complexes.³³ This species is known to be photoactive, losing multiple hydrides to form the $[Cp^*OsH_2]_2$ dimer. We reasoned that synthesis of actinide tetraosmate complexes analogous to tetrairidate complexes A and B might lead to more productive photolysis, ideally with the loss of hydrides and formation of An–Os bonds. Here, we report the syntheses, calculated bonding character, and photolytic chemistry of these uranium and thorium tetraosmate species.

Results and discussion

In order to install the osmium centers around the actinide center, Cp^*OsH_5 was converted *in situ* to the potassium osmate species $K[Cp^*OsH_4]$ (1) with the addition of benzylpotassium in THF, forming a pale-yellow solution. ¹H NMR confirms the stoichiometric deprotonation of Cp^*OsH_5 , resulting in formation of 1 and toluene (Fig. S7†). Compared to Cp^*OsH_5 , the hydride resonance of 1 shifts upfield from –11.00 ppm to –17.15 ppm, with a concomitant reduction in integration from five protons to four (relative to the Cp^* resonance). The ¹H NMR spectrum closely resembles that of the $Li[Cp^*OsH_4]$ species reported by Suzuki, which contains a hydride resonance at –17.52 ppm.³⁴

Reaction of four equivalents of 1 with $ThCl_4(DME)_2$ or UCl_4 in THF led to the formation of $Th\{(\mu-H)_4OsCp^*\}_4$ (2-Th) and $U\{(\mu-H)_4OsCp^*\}_4$ (2-U), which can be isolated as colorless and yellow crystals from *n*-hexane in 66% and 68% yield, respectively (Scheme 2).

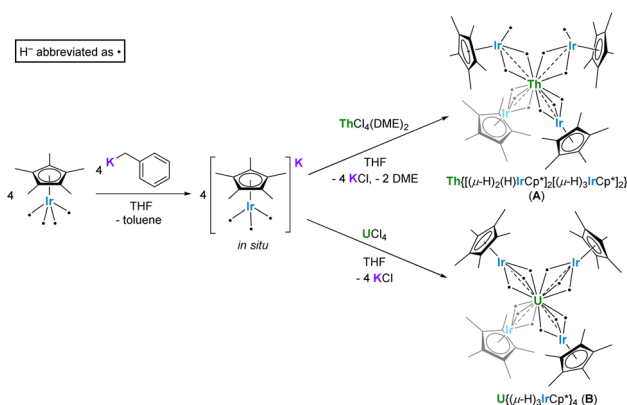
The ¹H NMR spectrum of diamagnetic 2-Th features two resonances for the Cp^* methyl protons and hydrides at

2.13 ppm and –11.01 ppm, respectively. These integrate in a 15 : 4 ratio, indicating that all hydrides in the potassium metalate starting material are preserved. The ¹H NMR spectrum of 2-U contains a sharp resonance at 3.65 ppm for the Cp^* methyl protons, as well as a hydride resonance at 89.8 ppm, shifted significantly downfield due to the paramagnetic influence of the formally uranium(IV) center. Once again, the two peaks integrate in a 15 : 4 ratio.

The solid-state IR spectra of 2-Th and 2-U each feature a single metal-hydride stretching signal at 1993 cm^{-1} and 1990 cm^{-1} , respectively, shifted by about 100 cm^{-1} compared to the reported hydride stretch for Cp^*OsH_5 of 2083 cm^{-1} .³³ These values are significantly higher than the bridging hydride stretch of 1762 cm^{-1} found in $[Cp^*OsH_2]_2$.³⁵ Several other examples of bridging hydrides in Cp^* -substituted osmium multimetallic complexes have been reported, but lack reported hydride stretches for comparison.^{34,36} However, our values are in good agreement with the bridging hydride stretching signals of 1962 and 1951 cm^{-1} for the iridate complexes A and B,³¹ which exhibit a similar decrease in frequency by approximately 100 cm^{-1} from the value for Cp^*IrH_4 of 2150 cm^{-1} .³⁷

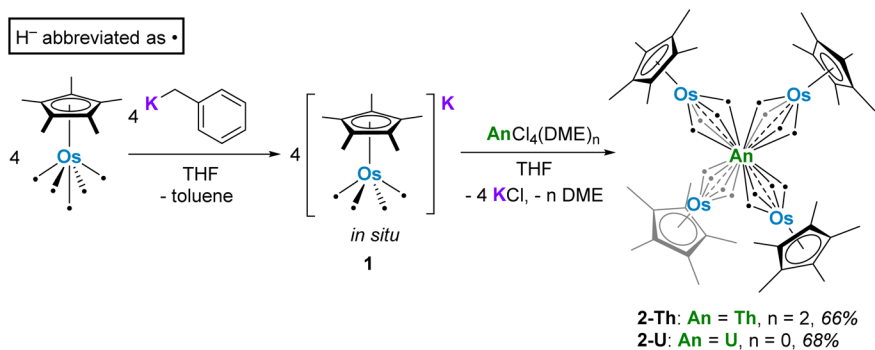
Single crystals of 2-Th and 2-U suitable for X-ray diffraction (XRD) were grown from saturated solutions of cold *n*-pentane, and the solid-state structures of both were crystallographically characterized, confirming that four osmium centers are coordinated around the actinide centers. No hydrides could be located in the difference maps, most likely due to the close proximity of numerous heavy atoms, a phenomenon that we observed previously with the related actinide-iridium complexes.³¹ Complex 2-Th crystallizes in the space group $P2_1/c$ with Th–Os distances ranging from 3.0183(6) Å to 3.0379(6) Å, well within the sum of covalent radii for Th and Os (3.50 Å).³⁸ The Os centers are slightly distorted from an ideal tetrahedral geometry around Th, with a τ_4 value³⁹ of 0.93 (calculated with $\alpha = 115.80(2)^\circ$, $\beta = 112.32(2)^\circ$), slightly less than that of A (0.97).³¹ The Os–Os distances range from 4.7446(7) Å to 5.1395(7) Å, which precludes any Os–Os interactions considering osmium's covalent radius of 1.44 Å.³⁸ Unlike A, complex 2-Th has one consistent hydride binding mode across all osmate moieties. All osmate fragments feature fairly linear Th–Os– $Cp^*_{centroid}$ angles (171.10(15)° to 176.92(14)°). These values are sufficiently close to linearity to indicate that all four hydrides in each $[Cp^*OsH_4]^-$ fragment bridge the Th and Os centers. Therefore, 2-Th is formally 16-coordinate, with 16 bridging hydrides around the thorium center, matching the highest observed coordination number around an atom in an isolated compound;^{40,41} interestingly, higher coordination numbers have been proposed for early actinide-noble gas complexes.⁴² This is reminiscent of the 15-coordinate amino-diborane thorium complex reported by Girolami, which also features thorium with an extraordinarily large number of bridging hydrides, in this case provided by borohydride-based ligands rather than transition metal polyhydrides.²³

Complex 2-U crystallizes in the cubic space group $Pa\bar{3}$, with a single osmate moiety centered on the 3-fold symmetry axis and another which generates the remaining three osmate fragments through symmetry. The on-axis U–Os distance is



Scheme 1 Previously reported syntheses of An–Ir multimetallic complexes A and B.



Scheme 2 Synthetic route to compounds **1** and **2-An**.

2.9490(6) Å, while the off-axis U–Os distances are each 2.9501(4) Å, far shorter than the sum of covalent radii for U and Os (3.40 Å)³⁸ and on average shorter than those of **2-Th** by approximately 0.08 Å (3.027(6) Å vs. 2.950(6) Å). The Os centers are arranged in an essentially tetrahedral manner around uranium, with a τ_4 value of 0.98 (calculated with $\alpha = 111.00(2)^\circ$, $\beta = 111.00(2)^\circ$), significantly more tetrahedral than **B**, which has a τ_4 value of 0.87.³¹ As in **2-Th**, the Os–Os distances are long enough to rule out any possible interaction, varying from 4.7703(8) Å to 4.8615(7) Å. The U–Os–Cp_{centroid}^{*} angles are quite linear, at 175.30(11)° for the on-axis fragment and 176.4(5)° for the off-axis fragments, indicating that as in **2-Th**, all 16 hydrides of **2-U** bridge the osmium centers and uranium. This is the first report of such a high coordination number for uranium, which previously had been limited to 14-coordinate complexes,^{43,44} and a second example of a formally 16-coordinate actinide complex.

To further support our conclusions about the hydride binding modes in **2-Th** and **2-U** as well as investigate potential metal–metal interactions, we turned to density functional theory (DFT) calculations for further insight. All DFT computations employed the B3PW91 functional (full details in the ESI†). Computational modeling of **2-Th** and **2-U** predicts near-tetrahedral configurations for the osmium fragments around the actinide centers, as observed in the solid-state structures, as well as sixteen bridging hydrides for each species (Fig. 1). The calculated An–Os distances are consistently ~0.04 Å shorter than the experimentally determined distances, but this and all other observed disparities between computational and experimental metrics are quite minor (Table 1). In addition, the two highest experimentally observed stretching frequencies for **2-Th** and **2-U** fall within the predicted ranges according to calculations, further evidence that these calculations are accurately modeling the hydrides within these compounds.

Natural Bond Order (NBO) analyses were carried out on **2-Th** and **2-U** as well, to analyze the degree of metal–metal interaction between the actinide and osmium centers. Wiberg Bond Indices (WBIs) of 0.72–0.73 were calculated for the Th–Os interactions in **2-Th**, while values of 0.81–0.83 were found for the U–Os interactions in **2-U**. These values suggest significant bonding interactions between the actinide and osmium centers, and are comparable to previously reported An–TM bonds as well as our

An–Ir systems.^{31,45,46} Complexes **2-Th** and **2-U** are, to the best of our knowledge, the first reported compounds to evidence An–Os covalency. Analysis of the M–H bonds reveals a more covalent hydride interaction with Os than U or Th, with Os–H WBIs of 0.57–0.60 compared to An–H WBIs of 0.18–0.24. Similar to **A** and **B**, the identity of the actinide does not appear to affect hydride covalency in **2-An**.

With these actinide-osmium polyhydrides in hand, we investigated the potential for thermolytic and photolytic H₂

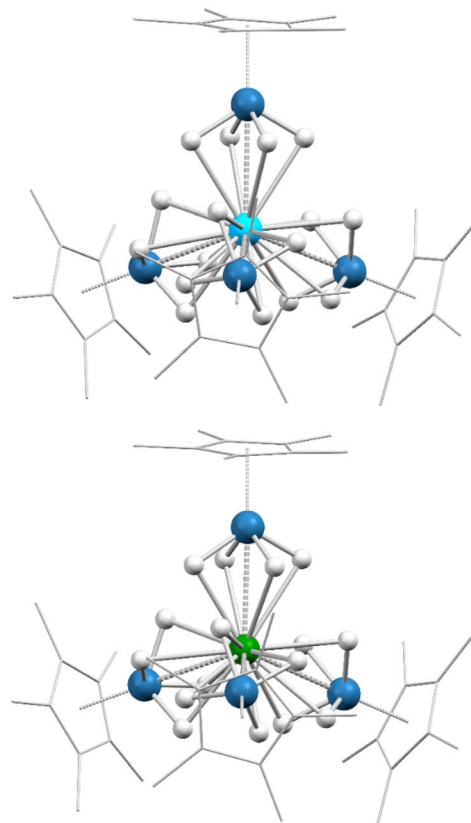


Fig. 1 Computed structures for **2-Th** and **2-U** (single-crystal XRD-derived ORTEP diagrams can be found in Fig. S32 and S33†). Color key: light blue (thorium), green (uranium), blue (osmium), white (hydrogen). Hydrogen atoms on the Cp^{*} rings are omitted and Cp^{*} ligands are wireframed for clarity.



Table 1 Experimentally and computationally derived bond distances, angles, and Os–H stretches for 2-Th and 2-U

	2-Th		2-U	
	Expt.	Comp.	Expt.	Comp.
An–Os dist. (Å)	3.0183(6)–3.0379(6)	2.977–2.978	2.9490(6)–2.9501(4)	2.911–2.913
Os–C _{Cp*} dist. (Å)	2.173(11)–2.320(10)	2.187–2.306	2.18(3)–2.29(3)	2.187–2.300
∠An–Os–C _{P_{centroid}*} (°)	171.10(15)–176.92(14)	178.9–179.3	175.30(11)–176.4(5)	177.1–179.2
Os–H (cm ⁻¹)	1990	2051–1963	1993	2056–1947
	862	947–838	863	939–818

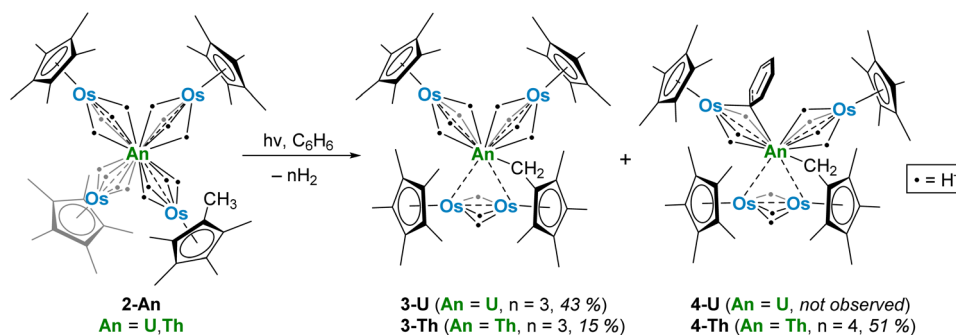
elimination in 2-U and 2-Th. Both complexes were stable in C₆D₆ when heated at 60 °C for 8 hours, as monitored by ¹H NMR. However, upon irradiation by 254 nm light in C₆H₆, complex 2-U undergoes a color change from yellow to dark brown, forming complex 3-U (Scheme 3) *via* the C–H activation of a Cp* methyl group and the loss of hydrides as H₂, as detected by ¹H NMR spectroscopy at 4.47 ppm in C₆D₆ (Fig. S12[†]).⁴⁷

Complex 3-U shows significantly reduced solubility compared to 2-U. It exhibits minimal solubility in *n*-hexane, requiring several drops of benzene to facilitate dissolution of the crude material. Upon workup in this manner, 3-U crystallizes at –40 °C as tiny brown crystals in 43% yield. Larger, X-ray quality crystals were grown from dilute solutions in *n*-hexane without benzene, from which the solid-state structure was determined (Fig. 2, left). As a result of photolysis, two of the Cp*Os moieties in 3-U feature formally Os(II) centers, with a significantly reduced Os–Os distance of 2.4639(4) Å and the formation of a U–C bond between uranium and the activated Cp* methylene. The U–Os distances of the remaining two formally Os(IV) fragments are slightly lengthened compared to 2-U, ranging from 2.9571(7) Å to 3.0264(7) Å. The U(IV) center is disordered across two positions, asymmetrically bridging the diosmium fragment towards either osmium atom with near-50% occupancy. There is some variation in the U–C bond length depending on which direction the asymmetric U atom favors, at 2.603(7) Å and 2.657(9) Å. These values are similar to the U–C bond lengths found for other uranium “tuck-over” complexes.^{48,49} In addition, the two unreacted [Cp*OsH₄][–] fragments in complex 3-U splay further outward from the tetrahedral geometry of 2-U, with an Os–U–Os angle of

118.15(3)° or 120.75(3)° depending on the uranium position. As with the previous complexes, no hydrides could be resolved in the solid-state structure.

As discussed previously, photolytic loss of hydrides has precedence with Cp*OsH₅, which is known to photolyze with the loss of 3 equivalents of H₂ to form the dimer [Cp*OsH₂]₂. This species has an essentially identical Os–Os distance of 2.4568(6) Å to that of 3-U.³⁵ The newly formed [(CH₂)Me₄C₅Os(μ-H)₃OsCp*]₂^{2–} fragment closely resembles the doubly deprotonated dimer, with one deprotonation occurring at a Cp* methyl group and the other occurring from the bridging hydrides, coordinated side-on to the actinide center. Hou previously reported the reaction and side-on coordination of [Cp*OsH₂]₂ with rare earth dialkyl complexes of the form (C₅Me₄SiMe₃)Ln(CH₂SiMe₃)₂(THF) (Ln = Y, Lu) resulting in trinuclear, C–H activated complexes.¹⁷ However, in these species, the geometry of the [Cp*OsH₂]₂ fragment was not preserved due to the migration of several hydrides to bridge the lanthanide and osmium atoms, and reactivity was promoted *via* thermolysis rather than photolysis.

The ¹H NMR spectrum of C_s symmetric 3-U contains six observable resonances, allowing all protons to be assigned besides the hydrides of the diosmium fragment. The mirror plane passes through the Os–Os and U–CH₂ bonds, resulting in the observation of two Cp* resonances (in a 2 : 1 ratio) and two methyl resonances from the activated Cp* (in a 1 : 1 ratio). The activated methylene resonance is shifted far upfield to –89.1 ppm. The hydrides of 3-U are observed at 55.5 ppm, significantly more shielded than the hydrides of 2-U, which are shifted to 89.8 ppm. This downfield signal integrates to about eight protons, and is therefore assigned to the hydrides of the

Scheme 3 Photolysis of 2-U and 2-Th with 254 nm light in C₆H₆ to produce 3-U, 3-Th, and 4-Th.

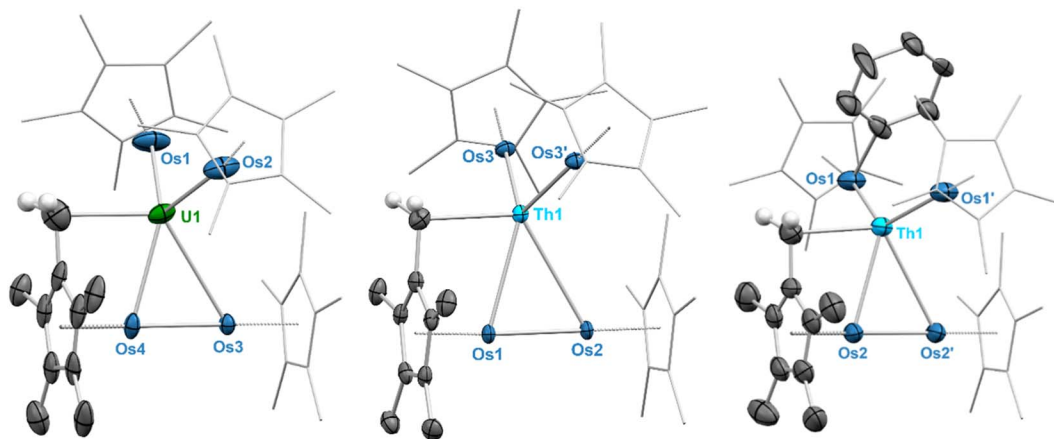


Fig. 2 Single-crystal XRD-derived ORTEP diagrams for **3-U** (left), **3-Th** (middle), and **4-Th** (right), with ellipsoids drawn at the 50% probability level. Non-methylene hydrogen atoms have been omitted and non-cyclometallated Cp* ligands have been wireframed for clarity. Only the major component of the disordered actinide center is shown in **3-U** and **3-Th**. Hydrides were not resolved in the crystal structures due to the numerous heavy metal centers in proximity. Selected bond distances (Å) and angles (°) for **3-U**: U1–Os1 2.9870(12), U1–Os2 3.0267(11), U1–Os3 3.2205(8), U1–Os4 2.7974(8), U1–C6 2.657(9), Os3–Os4, 2.4640(5), Os1–U1–Os2 118.15(3), Os3–Os4–Cp*_{centroid} 179.49(15), Os4–Os3–Cp*_{centroid} 179.45(15). Selected bond distances (Å) and angles (°) for **3-Th**: Th1–Os3 3.0260(3), Th1–Os2 3.2567(5), Th1–Os1 2.9539(4), Th1–C1 2.534(10), Os1–Os2, 2.4689(4), Os3–Th1–Os3' 121.133(14), Os1–Os2–Cp*_{centroid} 179.86(2), Os4–Os3–Cp*_{centroid} 177.133(17). Selected bond distances (Å) and angles (°) for **4-Th**: Th1–Os1 3.0593(9), Th1–Os2' 3.2965(11), Th1–Os2 2.9403(12), Th1–C12 2.703(12), Os2–Os2', 2.4536(9), Os1–Th1–Os1' 135.06(4), Os2–Os2'–Cp*_{centroid} 179.89(16).

two unactivated [Cp*OsH₄][−] fragments. However, the hydrides associated with the diosmium fragment were not observed between −100 to 600 ppm in the ¹H NMR spectrum, perhaps due to the paramagnetic nature of **3-U**. We therefore hypothesized that a diamagnetic Th(IV) photolysis product could serve as a useful analogue to **3-U**.

Complex **2-Th** displays slightly divergent reactivity from **2-U**, photolyzing over four hours with a color change from colorless to orange to produce both **3-Th** and **4-Th** (Scheme 3). As with the photolysis of **2-U**, H₂ formation can be detected by ¹H NMR spectroscopy (Fig. S15[†]). However, complex **3-Th**, the analogous species to **3-U**, is the minor product in this reaction. The major product, **4-Th**, results from the additional activation of one equivalent of the benzene solvent, replacing one bridging hydride from one [Cp*OsH₄][−] fragment with an asymmetrical bridging μ-η¹,η¹-phenyl ligand. This benzene activation appears to lend stability to the photolysis product, as reactions in *n*-hexane instead result in dark brown intractable mixtures and yield no crystalline product. In contrast, the photolysis product **3-U** can be prepared and isolated from *n*-hexane in satisfactory yields.

Compounds **3-Th** and **4-Th** consistently co-crystallize from the crude photolysis mixture, typically in a 22 : 78 **3-Th** to **4-Th** ratio (mol/mol%) as measured by ¹H NMR. Complex **4-Th** could be isolated in sparing (<20%) yield with photolysis times in excess of two days. Tracking of the reaction by ¹H NMR shows that **4-Th** forms in significant quantities with **3-Th** upon photolysis of **2-Th** and gradually increases in concentration over time, such that **3-Th** cannot be isolated with short reaction times. Addition of H₂ gas to a sample of **4-Th** did not lead to hydrogenolysis of the bridging phenyl ligand to form **3-Th**. Separation of **3-Th** from **4-Th** was achieved a single time by

recrystallization of a particularly high-percentage **3-Th**-containing photolysis crop (47% **3-Th** by ¹H NMR) from a 1 : 1 hexane/benzene solution at −40 °C, yielding crystals containing 88% **3-Th**. As a result, the ¹H NMR signals of each compound can be distinctly identified, although quantities sufficient for high-quality 2D NMR spectra were not obtained. Spectroscopic analyses were instead carried out on the crystalline mixture of **3-Th** and **4-Th**. As confirmation that two species are present in this material, Diffusion Ordered Spectroscopy (DOSY) experiments on a nearly 1 : 1 solution of **3-Th/4-Th** demonstrate that the ¹H NMR peaks associated with **3-Th** and **4-Th** have slightly different diffusion coefficients (Fig. S31[†]). This slight difference in diffusion rate is unsurprising given the relatively small difference in molecular weight and steric bulk between the two complexes, and is well outside the margin of error in these experiments.

The ¹H NMR spectrum of **3-Th** contains one more resonance than the spectrum of **3-U**. Analogous to **3-U**, there are four resonances in the methyl region, in addition to a singlet methylene resonance at 1.34 ppm. In contrast to the uranium species, there are two hydride signals, one integrating to eight hydrides and the other integrating to three hydrides, found at −9.74 ppm and −11.06 ppm, respectively. These signals are in turn assigned to the hydrides of the two [Cp*OsH₄][−] fragments and the hydrides of the diosmium fragment, [(CH₂)Me₄C₅Os(μ-H)₃OsCp*]^{2−}. Therefore, we assign three hydrides to the diosmium fragment of **3-U** as well, resulting in 11 total hydrides for both **3-U** and **3-Th**.

The ¹H NMR spectrum of **4-Th** is significantly more complex, as the bridging phenyl ligand renders each osmate fragment distinct. The phenyl signals of **4-Th** are observed between 7.07 and 7.98 ppm. The three unactivated Cp* moieties and four



methyl groups of the activated Cp* each appear as separate signals, and the methylene resonances arising from the Cp* cyclometallation are split as an AX system centered at 1.08 and 1.37 ppm. ¹H, ¹³C HSQC experiments confirm that the protons giving rise to these signals are bonded to the same carbon, which appears at 61.07 ppm in the ¹³C NMR spectrum. In the hydride region, the three hydrides of the [Cp*OsH₃(C₆H₅)][−] fragment are distinct in solution, and splitting can be observed between the three signals. The hydride distal to the activated benzene appears furthest downfield, at −8.16 ppm, and is split into a triplet by the two proximal hydrides, which appear as doublets at −10.54 and −10.84 ppm. The bridging hydrides of the [(CH₂)Me₄C₅Os(μ-H)₃OsCp*]^{2−} and [Cp*OsH₄][−] fragments are observed as two singlets at −9.80 and −11.59 ppm, respectively.

Both **3-Th** and **4-Th** are highly soluble in benzene and toluene while only sparingly so in *n*-hexane. Recrystallization of the aforementioned singular **3-Th** sample from *n*-hexane and minimal benzene afforded single crystals of **3-Th** suitable for diffraction experiments. X-ray quality crystals of **4-Th** were grown from *n*-hexane by recrystallizing the primarily **4-Th** crystalline material isolated from long photolysis times (>1 day).

Complex **3-Th** crystallizes with one equivalent of *n*-hexane in the space group *Pmn*2₁, with half of the molecule generated by a mirror plane (Fig. 2, middle). It is isostructural to **3-U**, but unlike **3-U**, the disorder of the central actinide atom strongly favors one orientation, at over 90% occupancy, indicating a preference for the molecules to align in the same orientation. Further discussion of the structure of **3-Th** will primarily refer to this major component. The diosmium fragment in **3-Th** is nearly identical to that of **3-U**, with a similar Os–Os distance of 2.4689(4) Å, but a shorter Th–C bond measuring 2.534(10) Å (2.729(10) Å in the minor component). The monoosmate fragments feature slightly longer Th–Os bond lengths than the U–Os distances in **3-U** at 3.0260(3) Å, with a similar Os–Th–Os angle of 121.133(14)°.

In comparison, the primary notable feature in the structure of **4-Th** is the presence of an asymmetric bridging phenyl ligand across one Th–Os interaction (Fig. 2, right). The structure is completely asymmetric, but due to disorder **4-Th** crystallizes in the space group *P4*₂/*ncm*, with the thorium center and the bridging phenyl ligand disordered equivalently across two sites, resulting in four possible conformations for the molecule. The diosmium fragment is isostructural to that of **3-Th**, with an Os–Os distance of 2.4536(9) Å and a Th–C bond length of 2.703(12) Å between Th and the activated Cp* methylene.

In the remaining portion of **4-Th**, the two monoosmate fragments splay further outward to accommodate the activated phenyl ring, with an expanded Os–Th–Os angle of 135.06(4)°, compared to 121.133(14)° in **3-Th**. Within this expanded pocket, one equivalent of activated benzene is present as a bridging μ-η¹, η¹-phenyl across one Th–Os interaction, replacing one of the bridging hydrides. This phenyl ring is coordinated asymmetrically, angled at 67.0(14)° relative to the Th–Os bond. It is heavily skewed toward the osmium center, with an Os–C distance of 2.12(5) Å compared to a Th–C distance of 2.96(4) Å. Examples of bridging μ-η¹, η¹-aryl ligands involving osmium are limited to

triosmium carbonyl clusters and feature symmetric coordination modes.^{50–52} Several late transition metal multimetallic species feature similar μ-η¹, η¹-aryls, including two related Au–Ag and Au–Cu polymeric chains,⁵³ as well as Pt–Ag and Pt–Cu complexes.^{54,55} The polymeric complex [Au₂Ag₂(C₆F₅)₄(N≡CCH₃)₂]_n most closely resembles the highly asymmetric phenyl coordination mode of **4-Th**, with Ag–C_{phenyl} distances of 2.508(6) and 2.687(6) Å, Au–C_{phenyl} distances of 2.055(6) and 2.088(6) Å, and a C_{phenyl}–Au–Ag angle of 66.00(15)°. In all these cases, however, the bridging aryl ligand is coordinated to a metal center in the starting material, whereas in **4-Th** it arises from the C–H activation of benzene. In the rare earths, a symmetric η¹-phenyl bridging two scandium centers has been proposed as an intermediate in the activation of benzene by (1,1'-fc(NSi^tBuMe₂)₂)₂ScI(THF)₂ and KC₈, though no bridging aryl species were isolated.⁵⁶

Photolysis reactions of **2-Th** in substituted aryl solvents such as toluene, mesitylene, and fluorobenzene were attempted as well. Photolysis in toluene led to a color change to orange, similar to the reaction in benzene. ¹H NMR analysis reveals a mixture of numerous products, presumably from the activation of the aryl ring at the *ortho*, *meta*, and *para* positions, as well as the formation of **3-Th**, further evidence that **3-Th** is formed without the activation of benzene (or any solvent). There does not appear to be a strong preference for activation at any position, making isolation of a single product untenable. Photolysis was less fruitful in mesitylene, with detection of **3-Th** as the major product, but less evidence of aryl activation products in the ¹H NMR spectrum. In contrast, reactions in fluorobenzene rapidly became brown, but the major identifiable products were Cp*OsH₅ and [Cp*OsH₂]₂, with no isolable C–H or C–F activation products of fluorobenzene.

In order to investigate potential metal–metal bonding interactions in compounds **3-U**, **3-Th**, and **4-Th**, better elucidate the positions of the hydrides in these photolysis products, and gain insight into the benzene activation by **4-Th**, we undertook a similar computational investigation as for complexes **2-U** and **2-Th**, at the same level of theory. Complexes with different numbers of hydrides were computed, considering different spin states for each (see Fig. S34 and Table S5 in ESI†). For **3-U** and **3-Th**, the most stable structures are found with eleven hydrides in a triplet and singlet spin state, respectively, which is in line with the presence of a U(IV) and Th(IV) actinide center (Fig. 3, top and S34†). In **4-Th**, a singlet spin state is once again most stable, indicating a Th(IV) center, while only ten hydrides are found due to the substitution of one hydride with the bridging phenyl ligand (Fig. 3, bottom). It is worth noting that the three optimized structures are the only structures with linear Os–Os–Cp*_{centroid} angles in line with the experimental structures; simulations with greater or fewer hydrides converged to structures with bent Os–Os–Cp*_{centroid} geometries. As with complexes **2-An**, the optimized geometries compare well with the experimental data (Table S6†). Among others, the Os–Os distance is well reproduced computationally at 2.47 Å, *versus* about 2.46 Å experimentally.

NBO analyses were carried out on the most stable optimized structures for the three complexes. WBIs of 0.53–0.79 were



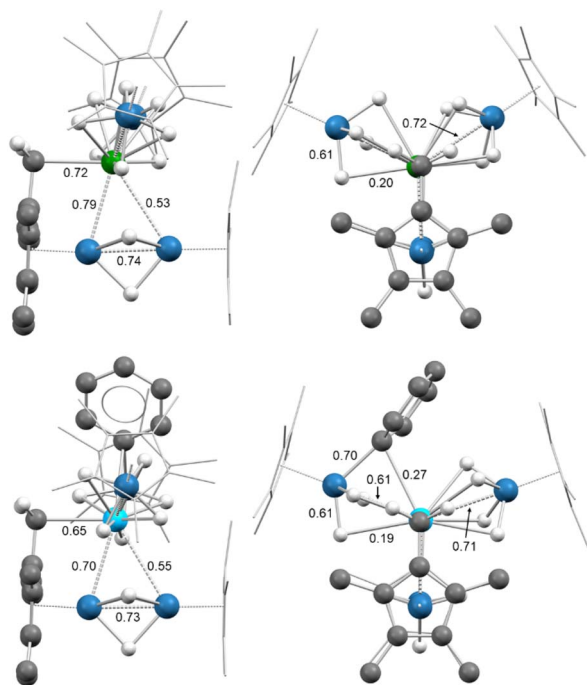


Fig. 3 Calculated structures for **3-U** (top) and **4-Th** (bottom), with WBIs labeled for select bonds. Color key: green (uranium), light blue (thorium), blue (osmium), gray (carbon), white (hydrogen). Methyl hydrogen atoms have been omitted and non-cyclometallated Cp* ligands have been wireframed for clarity.

found for the U–Os interactions in **3-U**, while WBIs of 0.55–0.70 were found for the Th–Os interactions in **3-Th**, with comparable values of 0.55–0.71 in **4-Th**. These values are lower than that found in the parent complexes **2-An** but indicates that a substantial covalent bonding An–Os interaction remains in these complexes. As in complexes **2-U** and **2-Th**, the M–H bonds are more covalent with Os than U or Th, with Os–H WBIs of 0.30–0.60 compared to An–H WBIs of 0.02–0.21 for **3-An** and **4-An**. NBO analysis of the M–C_{phenyl} bonding in **4-Th** shows a covalent Os–C bond (45% Os/55% C) which appears to be delocalized toward Th (second-order perturbation analysis, see Table S9†). It is thus a three-center-two-electron bond, as with the bridging hydrides (Fig. S36†).

Interestingly, in the three complexes an Os–Os WBI of 0.73–0.74 is found in the diosmium fragment. This result is in line with a substantial bonding interaction between the two Os centers. The interaction is further corroborated at the NBO level where a single covalent Os–Os bond is found in **3-U** and **3-Th** (only a second order perturbation was found for **4-Th**). This bond implies a 5d–5d overlap between the two Os centers in an almost non-polarized covalent bond (Table S8†); the hydride-mediated metal–metal interaction can also be observed in the HOMO-3 orbital of **3-Th** and **4-Th** (Fig. 4, left). As previously mentioned, the diosmium fragment in complexes **3-An** and **4-Th** is similar to [Cp*OsH₂]₂, which Girolami and coworkers concluded to contain no Os–Os bond, citing the results of Morokuma on the analogous diruthenium complex.^{34,35,57,58} There are several key differences between the compounds and

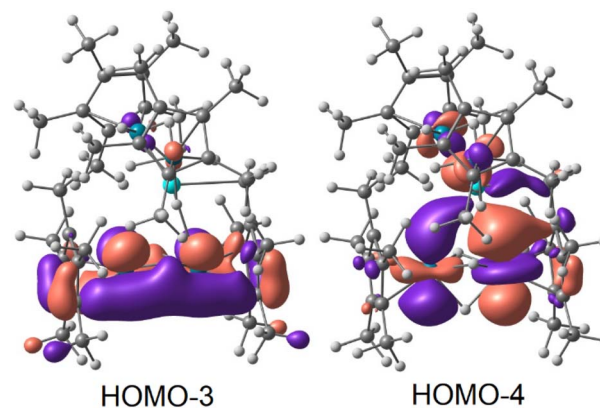


Fig. 4 Kohn–Sham Os–Os bonding orbitals for **3-Th** derived from DFT (isovalue = 0.03). The orbitals for **4-Th** are nearly identical and may be found in Fig. S35.†

computational methods used which may contribute to the discrepancy. First, the calculations in the 1993 paper were done using the *ab initio*, restricted Hartree-Fock method, in contrast to the semi-empirical method employed in this study. The geometry of the antibonding osmium, 5d_{z2} orbitals is different between the thorium compounds and [Cp*RuH₂]₂ as well. In the HOMO-4 orbital of **3-Th** and **4-Th**, the osmium 5d_{z2} orbitals are oriented such that they overlap with a thorium 6d orbital (Fig. 4, right), resulting in δ-antibonding interactions between the osmium orbitals rather than the σ-antibonding overlap predicted for [Cp*RuH₂]₂. This should decrease the repulsive interaction experienced between the osmium centers, and may account for the difference in the calculated Os–Os bonding.

Conclusions

Complexes **3-U**, **3-Th**, and **4-Th** are notable as products of photolysis-driven C–H activation by both actinide and transition metal centers. All feature the intramolecular activation of sp³ C–H bonds, while **4-Th** also features the uncommon intermolecular activation of a benzene sp² C–H bond by an actinide complex, resulting in a bridging Th–C–Os bond. Actinide-promoted benzene activation has been documented in uranium inverse sandwich complexes, resulting in the direct borylation of benzene and naphthalene, but in this instance did not result in the formation of a new An–C bond following activation.⁵⁹ Despite the report of thermolytic benzene C–H activation by thorium neopentyl species in 1981, subsequent examples of benzene activation by f-block metal centers remain few and far between. These reports have been limited to the rare-earth elements and involve thermolysis at high temperatures^{60,61} or strong reductants such as a potassium mirror or KC₈.^{56,62} In contrast, the formation of **4-Th** is photolytically driven, occurring at ambient conditions. The osmium polyhydride OsH₆(P^tPr₃)₂ has been known to activate a variety of polycyclic, *N*-substituted aromatic species upon thermolysis,^{63–67} but no reactivity with smaller, less substituted aromatics such as toluene or benzene has been reported. In addition, the photolytic reactivity of **2-U** and **2-Th** is not solely



dependent on the osmium centers, as both Cp^*OsH_5 and $[\text{Cp}^*\text{OsH}_2]_2$ do not form stable Os–C bonds upon photolysis in benzene, although H/D exchange is observed when Cp^*OsH_5 is photolyzed to form $[\text{Cp}^*\text{OsH}_2]_2$ in C_6D_6 (Fig. S30†). This is particularly exciting, as heterobimetallic cooperative C–H activation is a field of growing interest, yet systems featuring An–TM cooperativity are rare.⁶⁸

In the context of actinide photolysis reactivity, the reactions of 2-U and 2-Th are also unique. In contrast to previously reported photolytic actinide C–H activations, which were driven by irradiation of stable or transient terminal uranium nitrides^{69,70} or Th–C bond cleavage,⁷¹ here the chemistry is driven by the loss of hydrides as H_2 . This contributes to the minimal record of photolytic elimination of hydrides in the actinides,²⁸ as well as photolysis-driven actinide–metal bond formation.³² It is, to the best of our knowledge, the first documented example of the formation of new actinide bonds *via* photolytic hydride elimination. Future work will focus on the synthesis of heterobimetallic hydride-supported An–TM systems, hydride elimination to drive unsupported metal–metal bonding in those systems, and potential cooperative reactivity with small molecules.

Data availability

The experimental and computational data associated with this article are provided in the ESI.†

Author contributions

C. Z. Ye performed the experimental work. S. N. Kelly and I. J. Brackbill recorded and interpreted the XRD data. I. Del Rosal and L. Maron performed and interpreted the computational studies. C. Z. Ye wrote the original draft of the manuscript. J. Arnold and C. Camp conceptualized the research, found the funds, administrated the project, and supervised the work. I. Del Rosal, L. Maron, J. Arnold, and C. Camp revised and edited the manuscript. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 S. W. M. Crossley, C. Obradors, R. M. Martinez and R. A. Shenvi, *Chem. Rev.*, 2016, **116**, 8912–9000.
- 2 M. A. Esteruelas and L. A. Oro, *Chem. Rev.*, 1998, **98**, 577–588.
- 3 A. J. Jordan, G. Lalic and J. P. Sadighi, *Chem. Rev.*, 2016, **116**, 8318–8372.
- 4 N. A. Eberhardt and H. Guan, *Chem. Rev.*, 2016, **116**, 8373–8426.
- 5 M. Ephritikhine, *Chem. Rev.*, 1997, **97**, 2193–2242.
- 6 R. N. Perutz and B. Procacci, *Chem. Rev.*, 2016, **116**, 8506–8544.
- 7 L. Havela, D. Legut and J. Kolorenč, *Rep. Prog. Phys.*, 2023, **86**, 056501.
- 8 W. J. Evans, J. H. Meadows and T. P. Hanusa, *J. Am. Chem. Soc.*, 1984, **106**, 4454–4460.
- 9 D. Jr. Alvarez, K. G. Caulton, W. J. Evans and J. W. Ziller, *J. Am. Chem. Soc.*, 1990, **112**, 5674–5676.
- 10 D. Jr. Alvarez, K. G. Caulton, W. J. Evans and J. W. Ziller, *Inorg. Chem.*, 1992, **31**, 5500–5508.
- 11 M. L. H. Green, A. K. Hughes, D. M. Michaelidou and P. Mountford, *J. Chem. Soc., Chem. Commun.*, 1993, 591–593.
- 12 N. S. Radu, P. K. Gantzel and T. D. Tilley, *J. Chem. Soc., Chem. Commun.*, 1994, 1175–1176.
- 13 D. J. Schwartz, G. E. Ball and R. A. Andersen, *J. Am. Chem. Soc.*, 1995, **117**, 6027–6040.
- 14 M. V. Butovskii, O. L. Tok, F. R. Wagner and R. Kempe, *Angew. Chem., Int. Ed.*, 2008, **47**, 6469–6472.
- 15 A. P. Sobaczynski, T. Bauer and R. Kempe, *Organometallics*, 2013, **32**, 1363–1369.
- 16 T. Shima and Z. Hou, *Chem. Lett.*, 2008, **37**, 298–299.
- 17 T. Shima and Z. Hou, *Organometallics*, 2009, **28**, 2244–2252.
- 18 Y. Takenaka, T. Shima, J. Baldamus and Z. Hou, *Angew. Chem., Int. Ed.*, 2009, **48**, 7888–7891.
- 19 Y. Takenaka and Z. Hou, *Organometallics*, 2009, **28**, 5196–5203.
- 20 T. Shima, Y. Luo, T. Stewart, R. Bau, G. J. McIntyre, S. A. Mason and Z. Hou, *Nat. Chem.*, 2011, **3**, 814–820.
- 21 T. Shima and Z. Hou, *Chem.–Eur. J.*, 2013, **19**, 3458–3466.
- 22 W. W. N. O, X. Kang, Y. Luo and Z. Hou, *Organometallics*, 2014, **33**, 1030–1043.



- 23 S. R. Daly, P. M. B. Piccoli, A. J. Schultz, T. K. Todorova, L. Gagliardi and G. S. Girolami, *Angew. Chem., Int. Ed.*, 2010, **49**, 3379–3381.
- 24 A. C. Dunbar, J. C. Wright, D. J. Grant and G. S. Girolami, *Inorg. Chem.*, 2021, **60**, 12489–12497.
- 25 R. J. Lastowski, C. M. Caroff, K. D. Vogiatzis and G. S. Girolami, *Organometallics*, 2023, **42**, 2839–2848.
- 26 A. B. Altman, A. C. Brown, G. Rao, T. D. Lohrey, R. D. Britt, L. Maron, S. G. Minasian, D. K. Shuh and J. Arnold, *Chem. Sci.*, 2018, **9**, 4317–4324.
- 27 R. J. Ward, P. Rungthanaphatsophon, P. Huang, S. P. Kelley and J. R. Walensky, *Chem. Sci.*, 2023, **14**, 12255–12263.
- 28 D. Baudry and M. Ephritikhine, *J. Organomet. Chem.*, 1986, **311**, 189–192.
- 29 S. M. Cendrowski-Guillaume, M. Lance, M. Nierlich, J. Vigner and M. Ephritikhine, *J. Chem. Soc., Chem. Commun.*, 1994, 1655–1656.
- 30 S. M. Cendrowski-Guillaume and M. Ephritikhine, *J. Chem. Soc., Dalton Trans.*, 1996, 1487–1491.
- 31 C. Z. Ye, I. D. Rosal, M. A. Boreen, E. T. Ouellette, D. R. Russo, L. Maron, J. Arnold and C. Camp, *Chem. Sci.*, 2023, **14**, 861–868.
- 32 A. L. Ward, W. W. Lukens, C. C. Lu and J. Arnold, *J. Am. Chem. Soc.*, 2014, **136**, 3647–3654.
- 33 C. L. Gross, S. R. Wilson and G. S. Girolami, *J. Am. Chem. Soc.*, 1994, **116**, 10294–10295.
- 34 T. Shima and H. Suzuki, *Organometallics*, 2005, **24**, 3939–3945.
- 35 C. L. Gross and G. S. Girolami, *Organometallics*, 2007, **26**, 160–166.
- 36 T. Shima, T. Ichikawa and H. Suzuki, *Organometallics*, 2007, **26**, 6329–6337.
- 37 T. M. Gilbert and R. G. Bergman, *Organometallics*, 1983, **2**, 1458–1460.
- 38 B. Cordero, V. Gómez, A. E. Platero-Prats, M. Revés, J. Echeverría, E. Cremades, F. Barragán and S. Alvarez, *Dalton Trans.*, 2008, 2832–2838.
- 39 L. Yang, D. R. Powell and R. P. Houser, *Dalton Trans.*, 2007, 955–964.
- 40 I. A. Popov, T. Jian, G. V. Lopez, A. I. Boldyrev and L.-S. Wang, *Nat. Commun.*, 2015, **6**, 8654.
- 41 D. Pollak, R. Goddard and K.-R. Pörschke, *J. Am. Chem. Soc.*, 2016, **138**, 9444–9451.
- 42 L. Yang, S. Cooper and N. Kaltsoyannis, *Phys. Chem. Chem. Phys.*, 2021, **23**, 4167–4177.
- 43 E. R. Bernstein, W. C. Hamilton, T. A. Keiderling, S. J. La Placa, S. J. Lippard and J. J. Mayerle, *Inorg. Chem.*, 1972, **11**, 3009–3016.
- 44 A. V. Blake, T. V. Fetrow, Z. J. Theiler, B. Vlasisavljevich and S. R. Daly, *Chem. Commun.*, 2018, **54**, 5602–5605.
- 45 J. A. Hlina, J. R. Pankhurst, N. Kaltsoyannis and P. L. Arnold, *J. Am. Chem. Soc.*, 2016, **138**, 3333–3345.
- 46 G. Feng, K. N. McCabe, S. Wang, L. Maron and C. Zhu, *Chem. Sci.*, 2020, **11**, 7585–7592.
- 47 G. R. Fulmer, A. J. M. Miller, N. H. Sherden, H. E. Gottlieb, A. Nudelman, B. M. Stoltz, J. E. Bercaw and K. I. Goldberg, *Organometallics*, 2010, **29**, 2176–2179.
- 48 W. J. Evans, K. A. Miller, A. G. DiPasquale, A. L. Rheingold, T. J. Stewart and R. Bau, *Angew. Chem., Int. Ed.*, 2008, **47**, 5075–5078.
- 49 B. M. Gardner, J. McMaster, W. Lewis, A. J. Blake and S. T. Liddle, *J. Am. Chem. Soc.*, 2009, **131**, 10388–10389.
- 50 R. D. Adams, V. Rassolov and Q. Zhang, *Organometallics*, 2012, **31**, 2961–2964.
- 51 R. D. Adams, V. Rassolov and Q. Zhang, *Organometallics*, 2013, **32**, 6368–6378.
- 52 R. D. Adams, Z. Luo and Y. O. Wong, *J. Organomet. Chem.*, 2015, **784**, 46–51.
- 53 E. J. Fernández, A. Laguna, J. M. López-de-Luzuriaga, M. Monge, M. Montiel, M. E. Olmos and M. Rodríguez-Castillo, *Organometallics*, 2006, **25**, 3639–3646.
- 54 M.-E. Moret and P. Chen, *Organometallics*, 2008, **27**, 4903–4916.
- 55 M.-E. Moret and P. Chen, *J. Am. Chem. Soc.*, 2009, **131**, 5675–5690.
- 56 W. Huang, F. Dulong, S. I. Khan, T. Cantat and P. L. Diaconescu, *J. Am. Chem. Soc.*, 2014, **136**, 17410–17413.
- 57 H. Suzuki, H. Omori, D. H. Lee, Y. Yoshida and Y. Morooka, *Organometallics*, 1988, **7**, 2243–2245.
- 58 N. Koga and K. Morokuma, *J. Mol. Struct.*, 1993, **300**, 181–189.
- 59 P. L. Arnold, S. M. Mansell, L. Maron and D. McKay, *Nat. Chem.*, 2012, **4**, 668–674.
- 60 M. E. Thompson, S. M. Baxter, A. R. Bulls, B. J. Burger, M. C. Nolan, B. D. Santarsiero, W. P. Schaefer and J. E. Bercaw, *J. Am. Chem. Soc.*, 1987, **109**, 203–219.
- 61 L. Maron, E. L. Werkema, L. Perrin, O. Eisenstein and R. A. Andersen, *J. Am. Chem. Soc.*, 2005, **127**, 279–292.
- 62 M. P. Coles, P. B. Hitchcock, M. F. Lappert and A. V. Protchenko, *Organometallics*, 2012, **31**, 2682–2690.
- 63 M. A. Esteruelas, A. B. Masamunt, M. Oliván, E. Oñate and M. Valencia, *J. Am. Chem. Soc.*, 2008, **130**, 11612–11613.
- 64 M. A. Esteruelas, I. Fernández, A. Herrera, M. Martín-Ortiz, R. Martínez-Álvarez, M. Oliván, E. Oñate, M. A. Sierra and M. Valencia, *Organometallics*, 2010, **29**, 976–986.
- 65 B. Eguillor, M. A. Esteruelas, I. Fernández, M. Gómez-Gallego, A. Lledós, M. Martín-Ortiz, M. Oliván, E. Oñate and M. A. Sierra, *Organometallics*, 2015, **34**, 1898–1910.
- 66 M. A. Esteruelas, C. Larramona and E. Oñate, *Organometallics*, 2016, **35**, 1597–1600.
- 67 L. Cancela, M. A. Esteruelas, A. M. López, M. Oliván, E. Oñate, A. San-Torcuato and A. Vélez, *Organometallics*, 2020, **39**, 2102–2115.
- 68 A. Lachguar, A. V. Pichugov, T. Neumann, Z. Dubrawski and C. Camp, *Dalton Trans.*, 2024, **53**, 1393–1409.
- 69 R. K. Thomson, T. Cantat, B. L. Scott, D. E. Morris, E. R. Batista and J. L. Kiplinger, *Nat. Chem.*, 2010, **2**, 723–729.
- 70 D. M. King, F. Tuna, E. J. L. McInnes, J. McMaster, W. Lewis, A. J. Blake and S. T. Liddle, *Nat. Chem.*, 2013, **5**, 482–488.
- 71 N. S. Settineri and J. Arnold, *Chem. Sci.*, 2018, **9**, 2831–2841.

