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Introduction

The redox chemistry of the actinide elements has recently undergone a significant change: the range of oxidation states available in crystallographically-characterizable molecular complexes has been extended to +2. The discovery of the first molecular example of $U(\Pi)$ involved potassium graphite reduction of the tris(cyclopentadienyl) complex $\text{Cp}_3^{\'} \text{U}^{\text{III}}$ $(\text{Cp}^{\'} = \text{C}_5 \text{H}_4 \text{SiM} \text{e}_3)$ to $[\text{K}(\text{crypt})][\text{Cp}_3^{\'} \text{U}^{\text{II}}]$ $(crypt = 2.2.2-cryptand).$ ¹ Subsequently, the tris(cyclopentadienyl) complexes Cp_3'' An $(\text{Cp}'' = \text{C}_5\text{H}_3(\text{SiMe}_3)_2, \text{An} = \text{actinide})$ proved to be good precursors for the first examples of crystallographically-characterizable molecular compounds containing Th(II),² Np(II),³⁻⁵ and Pu(II),⁶ eqn (1). Examples of U(II) are now known in different coordination environments beyond the

tris(cyclopentadienyl) ligand sets of eqn $(1).^{7-9}$ Despite the rapid development of synthetic $An(\pi)$ chemistry, there have been few electrochemical studies of these low valent systems, although extensive electrochemistry has been reported for the higher oxidation states of the actinides.¹⁰–¹³ This is due in part to the high reactivity of the divalent and trivalent complexes. In addition, actinide electrochemical studies have been challenging because the +3 and +4 metal precursor complexes can react with supporting electrolytes. For example,

Electrochemical studies of tris(cyclopentadienyl) thorium and uranium complexes in the +2, +3, and +4 oxidation states†

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Electrochemical measurements on tris(cyclopentadienyl)thorium and uranium compounds in the $+2$, $+3$, and +4 oxidation states are reported with $C_5H_3(SiMe₃)_2$, $C_5H_4SiMe₃$, and $C_5Me₄H$ ligands. The reduction potentials for both U and Th complexes trend with the electron donating abilities of the cyclopentadienyl ligand. Thorium complexes have more negative An(III)/An(II) reduction potentials than the uranium analogs. Electrochemical measurements of isolated Th(II) complexes indicated that the $Th(III)/Th(II)$ couple was surprisingly similar to the $Th(IV)/Th(III)$ couple in Cp"-ligated complexes. This suggested that Th(II) complexes could be prepared from Th(IV) precursors and this was demonstrated synthetically by isolation of $[K(crown)(THF)_2][Cp_3''Th^{II}]$ directly from $Cp_3''Th^{IV}Br.$ UV-visible spectroelectrochemical measurements and reactions of ${Cp_3}''\text{Th}^{\text{IV}}$ Br with elemental barium indicated that the thorium system undergoes sequential one electron transformations. **EDGE ARTICLE**
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 **Electrochemical studies of tris(cyclopentadienyl)

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Inman and Cloke found problems studying $(C_5Me_5)Th^IV[C_8]$ $H_6(SiMe_2^{\ t}Bu)_2]$ Cl using [ⁿBu₄N][PF₆] as supporting electrolyte¹⁴ as well as with Cp_3 "Th^{IV}Cl using $\binom{n}{0}$ BU₄N][B(C₆F₅)₄] as supporting electrolyte.¹⁵

Although electrochemical data have been reported on two $U(n)$ systems,^{9,16} analogous studies on Th(π) complexes and on the tris(cyclopentadienyl) systems that led to the first molecular examples of $U(n)$ have been absent. Meyer and coworkers identified the U(III)/U(II) couple in $[(\text{Ad}, \text{Me} A r \text{O})_3 \text{me} s] U^{III}$ at -2.495 V vs. Fc^{+/0},¹⁶ that guided synthetic efforts and allowed isolation of $[K(crypt)]\{[(A^{Ad,Me}ArO)_3mes]U^{II}\}$.⁷ More recently, Layfield and coworkers reported the U(m)/U(n) couple of (C_5 - Pr_5 ₂U^{II} to be -2.33 V vs. Fc^{+/0}.⁹ Inman and Cloke studied Th(_{IV})/ Th(m) redox couples and found that $\text{[^{n}\text{Bu}_{4}\text{N}][\text{B} \text{Ph}_{4}]}$ was a good supporting electrolyte for their complexes.^{15,17} Encouraged by their results, we utilized this supporting electrolyte to obtain electrochemical data in this study and on $\text{Cp}_3\text{'Th}^{\text{IV}}\text{Cl}.^{\text{18}}$

Due to the importance of the tris(cyclopentadienyl) ligand set in the development of low oxidation state actinide chemistry,^{19,20} the electrochemistry of a variety of tris(cyclopentadienyl) uranium and thorium complexes using Cp'' , Cp' , and Cp^{tet} ligands ($Cp^{tet} = C_5Me_4H$), Scheme 1, is reported here as well as the first reported electrochemical measurements on isolated $Th(u)$ complexes.² Also reported are spectroelectrochemical studies on the Th (n) compounds that led to the discovery of new synthetic routes to $Th(II)$ compounds. The results are compared with cyclopentadienyl ligand effects previously examined electrochemically with titanium and zirconium complexes 21 and with rare-earth metal reaction chemistry.²²–²⁴

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Results

Electrochemical protocol

All data were collected in THF using 100 mM $\rm [^{\it n}Bu_{4}N][BPh_{4}]$ or 200 mM $\left[^{n}Bu_{4}N\right][PF_{6}]$ supporting electrolyte concentrations. Both $\rm \lbrack^{n}Bu_{4}N\rbrack\lbrack BPh_{4}\rbrack$ and $\rm \lbrack^{n}Bu_{4}N\rbrack\lbrack PF_{6}\rbrack$ were recrystallized three times prior to use. The low polarity of THF leads to large internal resistance in the electrochemical cell with peak separations over 200 mV often observed.^{15,16} Unless specifically stated, all potentials are referenced to the ferrocenium/ ferrocene couple with $(C_5Me_5)_2Fe$ as an internal standard, Fig. S12 and S13.† All electrochemical data were collected with a glassy carbon disc working electrode, platinum wire counter electrode, and silver wire pseudo-reference electrode. All scans were recorded in the cathodic direction except for the isolated $U(\Pi)$ and Th (Π) compounds which were recorded in the anodic direction. Representative cyclic voltammograms are shown in Fig. 1–6 and complete details are in the ESI.†

Uranium complexes

Initially, $U(m)$ complexes known to undergo chemical reduction and oxidation were examined to determine if both the $U(w)/i$ $U(m)$ and $U(m)/U(n)$ redox events could be observed electrochemically. Indeed, both redox couples were observed in the voltammograms for the U($\scriptstyle\rm III$) complexes Cp₃[']U^{III},²⁵ Cp₃["]U^{III},²⁶ and $\text{Cp}_3^{\text{tet}} \text{U}^{\text{III}}$,²⁶ and for the isolated U(II) complexes $[K(crown)(THF)₂][Cp₃''U^{II}](crown = 18 - crown - 6)²⁷$ and

 a 100 mM $[^n\text{Bu}_4\text{N}][\text{BPh}_4]/\text{THF}.$ b 50 mM $[^n\text{Bu}_4\text{N}][\text{BPh}_4]/\text{THF}.$ c 130 mM $[^n\text{Bu}_4\text{N}][\text{PF}_6]/\text{THF}.$

Table 2 Reduction potentials assigned to $U(III)/U(II)$ couples in this study and the literature

 $[K(crypt)][Cp_3'U^{II}]^1$ These values are summarized in Tables 1 and 2 and highlights are described in the following paragraphs.

 Cp'' . With the bis(trimethylsilyl)cyclopentadienyl ligand, redox couples assigned to $U(w)/U(m)$ and $U(m)/U(n)$ are observed at -0.94 V and -2.73 V, respectively, for $Cp_3^{\prime\prime}U^{\text{III}}$, Fig. 1 and S14.† In comparison, the isolated $U(\Pi)$ complex $[K(\text{crown})(\text{THF})_2][\text{Cp}_3{''}\text{U}^{\text{II}}]^{27}$ displays two redox events at -0.73 V and -2.71 V, Fig. 1 and S25.[†] The $E_{1/2}$ values for the $U(m)/U(n)$ couple are nearly identical in both systems and the event centered at -2.71 V only appears when scanning anodically for $[K(crypt)][Cp_3''U^{II}],$ which supports the assignment as the $U(m)/U(n)$ couple.

Cp'. Similar reproducible data were obtained with the mono(trimethylsilyl)cyclopentadienyl ligand with $U(w)/U(m)$ and $U(m)/U(n)$ couples at -1.26 V and -2.26 V, respectively, for $\text{Cp}_{3}^{\prime} \text{U}^{\text{III}}$, Fig. 2 and S17.† Likewise, the U($\text{I}\text{V}}\text{U}(\text{II})$ and U($\text{II}\text{U}(\text{II})$ couples were observed at -1.28 V and -2.27 V for the U(II) complex $[K(crypt)][Cp_3'U^{II}],$ Fig. 2 and S24.† These data were obtained with 50 mM $\binom{n}{k}$ Bu₄N][BPh₄] because decomposition occurred at higher electrolyte concentrations. The event at

 -2.27 V for $[K(crypt)][Cp_3'U^{II}]$ only appears when scanning anodically. The -2.27 V $E_{1/2}$ value for $[K(crypt)][Cp_3'U^{\text{II}}]$ was less negative than the -2.71 V value for $[K(crypt)][Cp_3''U^{II}],$ but it is similar to the two previously reported $U(m)/U(n)$ couples for $[(A^{d,Me}ArO)₃mes]U^{III}$ and $(C₅ⁱPr₅)₂U^{II}_{9,16}$ The minor unassigned events at about -1.9 V in Fig. 2 and S24 \dagger attest to the complexity of the system. They were observed across multiple runs and do not disappear after repeated recrystallization of substrate and electrolyte.

 Cp^{tet} . With the tetramethylcyclopentadienyl ligand, the U($\frac{V}{V}$) U(m) and U(m)/U(n) couples in Cp^{tet}U^{III} were more negative than in Cp₃ $^{\prime\prime}$ U^{III} and Cp₃ $^{\prime}$ U^{III}: -1.46 V and -3.11 V, Fig. 3 and S20.† However, data could not be obtained from the isolated $U(n)$ compound $[K(crypt)][Cp_3^{\text{tet}}U^{II}]$ because contact with the supporting electrolyte led to immediate decomposition. The voltammogram obtained from the resulting solution displayed at least five redox events, Fig. S29.† This reactivity is consistent with the more strongly reducing nature of the Cp^{tet} complexes as shown by the data in Tables 1 and 2. A third, minor event at -1.7 V was present and cannot be assigned with confidence.

Fig. 1 Voltammogram of 4.6 mM Cp3"U^{III} (solid) and 3.0 mM $[K(crown)(THF)_2][Cp_3"U^I]$ (dashed) at $\nu=$ 200 mV s $^{-1}$, in 100 mM ["Bu₄N][BPh₄]/ THF. The event centered at -0.495 V is due to internal standard (C₅Me₅)₂Fe.

Fig. 2 Voltammogram of $\mathsf{Cp}_3'U^{\parallel\parallel}$ (solid) and $[\mathsf{K}(\mathsf{crypt})][\mathsf{Cp}_3'U^{\parallel}]$ (dashed) at $\nu = 200$ mV s⁻¹, in 50 mM [ⁿBu₄N][BPh₄]/THF. The event centered at -0.495 V is due to internal standard (C $5Me5$)

Thorium complexes

Electrochemical data were collected on all the thorium compounds in this study using both $\binom{n}{1}$ \Pr_6 and $\binom{n}{1}$ \Pr_4 despite multiple reports that electrochemical data on organothorium complexes are difficult to obtain using $\binom{n}{1}B\mu_A N$ $[PF_6]$.^{11,14,15,30-32} Since the voltammograms do not differ drastically between electrolytes, only the data using $[^{n}Bu_{4}N][BPh_{4}]$, Table 3, are discussed below (data with $\binom{n}{1}$ Bu₄N][PF₆] are in Table S1†).

Thorium (w) complexes

 \mathbf{Cp}'' . Initially, $\mathbf{Cp}_{3}^{''}\mathbf{Th}^{\text{IV}}\mathbf{Cl}$ was examined to compare with the values previously reported by Cloke et al .¹⁵ The cyclic voltammogram of Cp_3 "Th^{IV}Cl under our conditions shows the Th (iv) / Th(III) couple at -2.93 V, Fig. S34, \dagger which is close to the value of

-2.96 V reported for Cp_3 "Th^{IV}Cl and Cp_3 "Th^{III}.¹⁵ Similarly, the cyclic voltammogram of Cp_3 "Th^{IV}Br (ref. 2) shows a Th(iv)/Th(iii) redox couple at -2.89 V, Fig. 4 and S30.[†] This suggests that the identity of halide does not significantly affect the reduction potential in this system. This is also consistent with bulk synthetic studies that show that Cp_3 "Th III can be synthesized from both Cp_{3} "Th^{IV}Cl and Cp_{3} "Th^{IV}Br.^{2,33,34}

Cp' and Cp^{tet}. Cp₃'Th^{IV}Cl³⁵ and Cp₃^{tet}Th^{IV}Br (ref. 36) were also examined as each these complexes can be chemically reduced to form tris(cyclopentadienyl)Th(III) species.^{18,36} The cyclic voltammogram of $\text{Cp}_{3}^{\ \prime} \text{Th}^{\text{IV}} \text{Cl}$, 35 Fig. S38,† exhibited a cathodic event at -3.14 V that is 0.21 V more negative than that of Cp_{3} "Th^{IV}Cl. Similarly, the voltammogram of Cp_{3} [']Th^{IV}Br had a cathodic event at -3.17 V, Fig. 4 and S63.[†] This event was determined to be a one electron process by comparing the current passed to that of the internal standard, Fig. S65.† The voltammogram of Cp^{tet}Th^{IV}Br had a cathodic event at -3.48 V, Fig. 4 and S44.† The events in the voltammograms of $\text{Cp}_3\text{/Th}^{\text{IV}}\text{Br}$ and $\mathsf{Cp}_3^\mathsf{tet}$ Th $^\mathsf{IV}\!\mathsf{Br}$ are practically irreversible even at scan rates up to 2000 mV s^{-1} . These results, along with the uranium studies above in Table 1, clearly show that the reduction potential of the actinide complex trends with the electron donation strength of the ligand in the order of $Cp^{tet} > Cp' > Cp''$. Chemical Science

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In addition to the $Th(w)/Th(m)$ couple, the voltammograms of the Th (w) compounds showed an irreversible anodic process that could be a cyclopentadienide oxidation, based on the electrochemical data collected on the cyclopentadienyl salts, KCp', KCp'', and KCp^{tet}, Fig. S66.† These irreversible anodic events were not found in the uranium systems. This difference in Th and U electrochemistry has been previously observed.^{11,15,37,38} Clearly, the Lewis acidity of the metal influences the potential for these cyclopentadienide oxidations.

Fig. 3 $\:$ Voltammogram of 7.2 mM Cp $_3^{\rm tet}$ U^{III} (solid, 100 mM [n Bu $_4$ N][BPh $_4$]/THF) compared to voltammograms of 4.6 mM Cp $_3$ $^{\prime\prime}$ U^{III} (dashed, 100 mM $[^nBu_4N][BPh_4]/THF$) and 11 mM Cp₃[']U^{|||} (dotted, 50 mM [ⁿBu₄N][BPh₄]/THF) at $\nu = 200$ mV s⁻¹. The events centered at -0.495 V are due to internal standard $(\mathsf{C}_5\mathsf{Me}_5)_2\mathsf{Fe}^{\mathsf{II}}$.

Fig. 4 Voltammogram of 7.4 mM Cp₃"Th^{IV}Br (solid), 15 mM Cp₃'Th^{IV}Br (dashed), and 12 mM Cp^{tet}Th^{IV}Br (dotted) at ν = 200 mV s⁻¹, in 100 mM [n Bu4N][BPh4]/THF.

Cyclopentadienyl rings bound to K^+ , $[K(\text{chelate})]^+$, or Anⁿ⁺ could have different oxidation potentials as evidenced by the differing voltammograms of KCp", $[K(crown)][Cp'']$, and $[K(crypt)][Cp'']$, Fig. S67.†

$Th(m)$ complexes

 Cp'' . There are fewer Th(III) options to study since there are only five crystallographically-characterized tris(cyclopentadienyl) Th(III) complexes, Cp_3 "Th^{III},^{33,34} $[\text{C}_5\text{H}_3(\text{SiMe}_2^{\text{t}}\text{Bu})_2]_3$ - $\mathrm{Th}^{\mathrm{III}},$ 34 $\mathrm{Cp}_3^\mathrm{tet} \mathrm{Th}^{\mathrm{III}},$ 36 $(\mathrm{C}_5{}^t \mathrm{Bu}_2 \mathrm{H}_3)_3 \mathrm{Th}^{\mathrm{III}},$ 39 and $(\mathrm{C}_5 \mathrm{Me}_5)_3 \mathrm{Th}^{\mathrm{III}},$ 40 Other $Th(m)$ compounds have been isolated with different ligand environments,⁴¹⁻⁴⁵ but our initial attempts to collect electrochemical data on $(\mathrm{C_5Me_5})_2 \mathrm{Th}^{\mathrm{III}}[^{i}\mathrm{PrNC}(\mathrm{Me})\mathrm{N}^{\mathrm{i}}\mathrm{Pr}]^{\mathrm{44}}$ led to immediate decomposition. Inman and Cloke found that scanning anodically on Cp_{3} "Th $^{\text{III}}$ gave a process at -2.96 V that matched

the reduction of Cp_{3} "Th^{IV}Cl described above and established the Th $(w)/Th(m)$ couple.¹⁵ In our hands, scanning cathodically on $\mathbb{C} p_3^{\prime\prime} \mathbb{C} \mathbb{H}^{\mathrm{III}}$ showed a voltammogram with a redox process centered at -2.85 V, Fig. 5 and S40.[†] A second cathodic event appears after the first cycle at -2.29 V, or when scanning anodically from the open circuit potential, Fig. S40.† The event at -2.29 V was also observed by Cloke and was attributed to a ligand-based event.

 \overline{C} p' and \overline{C} p^{tet}. Since \overline{C} p₃'Th^{III} has only been generated *in* situ,¹⁸ it was not studied under the present conditions. The voltammogram of Cp₃^{tet}Th^{III} at $\nu = 200$ mV s⁻¹ displays only a cathodic event, but at $\nu \geq 400$ mV s⁻¹, a return oxidation appears and the $Th(m)/Th(n)$ redox couple is centered at 3.28 V, Fig. 5 and S48.† This value matches the trend observed for the uranium systems in that Cp^{tet} complexes of thorium are more difficult to reduce than the silyl-cyclopentadienyl analogs.

Fig. 5 Voltammogram of 4.9 mM Cp3"Th^{III} (solid) at $\nu =$ 200 mV s $^{-1}$ and 6.7 mM Cp $_3^{\rm tet}$ Th^{III} (dashed) at $\nu =$ 400 mV s $^{-1}$ in 100 mM ["Bu4N][BPh4]/ THF.

An anodic event at -1.87 V is present and is attributed to a Cp^{tet}based process.

$Th(_{II})$ complexes

The only isolated Th(II) compounds $[K(crown)(THF)₂][Cp₃''Th^H]$ and $[K(crypt)][Cp₃''Th^H]$ exhibited nearly identical voltammograms. Scanning anodically, $[\mathrm{K}(\mathrm{crown})(\mathrm{THF})_2][\mathrm{Cp_{3}}''\mathrm{Th}^{\mathrm{II}}]$ showed a redox process centered at -2.84 V, which is assigned as the Th($\text{m}/\text{Th}(\text{n})$ redox couple, and a second irreversible anodic event at -1.38 V, attributed to ligand-based oxidation, Fig. 6 and S52.† The voltammogram of this Th (n) compound was practically identical over 5 cycles, Fig. S54.† $[K(crypt)][Cp_3''Th^H]$ similarly showed a reversible event centered at -2.85 V and a second anodic event at -1.43 V, Fig. 6, S57 and S61.†

Thorium spectroelectrochemistry

The data on isolated $[\text{Cp}_{3}^{\ \prime\prime}\text{Th}^{\text{II}}]^{1-}$ complexes suggested that the $Th(m)/Th(n)$ redox process occurs at about the same potential as the Th $(w)/Th(m)$ potential of Cp_3 "Th^{IV}Br. To investigate this further, spectroelectrochemical UV-visible measurements were obtained. A potential of -2.90 V was applied to a solution of $\mathrm{Cp}_3^{\prime\prime}\mathrm{Th}^{\mathrm{IV}}$ Br in 200 mM $\mathrm{[^{n}\mathrm{Bu}_4\mathrm{N}][\mathrm{PF}_6]}$ THF and the UV-visible spectrum was recorded approximately every 5 seconds during electrolysis. The formation of Cp_{3} "Th III is clearly shown by the growth of four bands at roughly 360, 500, 580, and 680 nm, Fig. 7, which correspond to the absorption spectrum of Cp_{3} "Th $^{\text{III}}$.^{33,34} No further reduction to the $[Cp_3^{\prime\prime}Th^{II}]^{1-}$ was observed,² although it cannot be ruled out as the absorbance spectrum reached the maximum of the detector.

Fig. 6 Voltammogram of 4.6 mM $[K(crown)(THF)_2][Cp_3''Th^{\parallel}]$ (solid) and 3.1 mM $[K(crypt)][Cp_3''Th^{\parallel}]$ (dashed) with internal standard (C₅Me₅)₂Fe at $\nu = 200 \text{ mV s}^{-1}$ in 100 mM [ⁿBu₄N][BPh₄]/THF.

Fig. 7 UV-visible spectrum of Cp₃"Th^{IV}Br (black, solid) converting to Cp_3 "Th^{III} (black, dashed) during electrolysis at -2.90 V with a starting concentration of 7.0 mM in 200 mM $[^nBu_4N][PF_6]/THF$. The growth of four bands at 365, 510, 590, and 655 nm is indicative of $\mathsf{Cp}_3{}^{\prime\prime}\mathsf{Th}^{\mathsf{III}}$ $(red).³⁴$

Fig. 8 UV-visible spectrum of Cp_3 "Th^{III} (black) converting to $[Cp₃''Th^{II}]^{1–}$ (blue) during electrolysis at -2.90 V with a starting concentration of 1.1 mM in 200 mM $[^{n}Bu_{4}N][PF_{6}]/THF$. The growth of the band at 650 nm is indicative of $[\text{Cp}_3" \text{Th}^{\text{II}}]^{1-}$ (red).²

Electrolysis of a solution of Cp_3 "Th $^\mathrm{III}$ in 200 mM $\mathrm{[^{n}\text{Bu}_4\text{N}]}$ $[PF_6]/THF$ at -2.90 V shows clean conversion to the Th(II) species $[Cp_3^{\prime\prime}Th^{II}]^{1-}$,² as indicated by the growth of the large absorption at 650 nm and the concomitant decrease in absorptions at 360, 500, 580, and 680 nm, Fig. 8. Although the absorption spectrum of $\mathbb{C}p_{3}$ "Th $^{\text{III}}$ had disappeared, the absorption at 650 nm, indicative of Th (n) ,² decreased in intensity as the electrolysis continued. The $Th(\pi)$ species appears to be unstable under the electrolysis conditions.

Chemical synthesis of Th (n) complexes from Th (w) precursors

The similarity of the Th $(w)/Th(m)$ couple in Cp_3 "Th^{IV}Br and Th(III)/Th(II) couple in $[Cp_3^{\prime\prime}Th^{II}]^{1-}$ suggested that Th(IV) compounds could be used as the precursors to $Th(II)$ compounds as well as the known Th (m) precursor, Cp_3 "Th $^{\mathrm{III}}$. Indeed, reaction of 2.2 equivalents of $KC₈$ to a THF solution of $\mathrm{Cp_{3}}''\mathrm{Th}^{\mathrm{IV}}\mathrm{Cl}$ and crown afforded $[\mathrm{K}(\mathrm{crown})(\mathrm{THF})_{2}][\mathrm{Cp_{3}}''\mathrm{Th}_{_}]$ in 50% crystalline yield, with a significant amount of $\mathrm{Cp}_3^{ \prime\prime} \mathrm{Th}^{\mathrm{III}}$ as a byproduct. Previously, Lappert reported that prolonged

stirring of a solution of Cp_3 "Th^{IV}Cl over excess NaK alloy developed a green color, 34 which was later confirmed to be the color of $Th(II).²$

Conversion of Th (w) to Th (n) was also studied with $\mathrm{Cp_{3}}''\mathrm{Th}^{\mathrm{IV}}\mathrm{Br}$. Reaction of $\mathrm{Cp_{3}}''\mathrm{Th}^{\mathrm{IV}}\mathrm{Br}$ with 2 equivalents of KC₈ in THF generated a dark green solution characteristic of $Th(II)$ within 5 minutes, as did reaction of Cp_3 ^{*"*}Th^{IV}Br with excess Na and with excess Li. The UV-visible spectra of these solutions have a strong absorption at 650 nm, identical to the previously reported spectra of $[K(crypt)][Cp_3"Th^H]$ and $[K(crown)(THF)_2][Cp_3"Th^H]_{...}$ ² but the spectra also show a non-negligible amount of Cp_{3} "Th $^{\text{III}}$.³⁴ Formation of the Th (m) complex is reasonable based on the fact that $[Na(\kappa^6\text{-}crown)(\kappa^2\text{-}crown)][Cp_3''\text{Th}^II]$ (see below) reacts with Cp_3 "Th^{IV}Br in THF to immediately form Cp_3 "Th^{III} in near quantitative yield.

These results show that a chelating agent is not necessary for the chemical synthesis of $Th(II)$ species in solution. However, the chelating agent appears necessary for efficient separation of the Th (n) product from the Th (m) starting material, as pure samples of $[M(THF)_x][Cp_3^TTh^II]$ $(M = Li, Na, K)$ were not isolated even though it is possible to isolate chelate-free examples of $[Cp_3''U^{II}]^{1-.46}$ Further support for the importance of alkali metal chelates is that addition of 18-crown-6 to the reaction of Cp_3 "Th^{IV}Br and excess Na provided X-ray quality crystals that were identified as $[Na(\kappa^6\text{-}crown)(\kappa^2\text{-}crown)]$ $[Cp_3$ "Th^{II}], only the third reported crystal structure of a Th (n) complex, Scheme 2, Fig. 9.

Similarly, the reaction of Cp_{3} "Th $^{\text{III}}$, Rb, and crypt in THF afforded dichroic blue/red crystals of $[Rb(crypt)][Cp_3''Th^H]$, isolated in 61% crystalline yield and identified by X-ray crystallography, Scheme 2, Fig. S69.† In addition, the reaction of Cp₃"Th^{III}, Cs, and crypt afforded dark blue/red crystals of $[Cs(crypt)][Cp_3''Th^H]$ in 54% crystalline yield, Scheme 2, Fig. S70.† The $[Rb(crypt)]^{1+}$ and $[Cs(crypt)]^{1+}$ compounds are isomorphous with the $[K(crypt)]^{1+}$ analog² and can be easily separated from the Cp_3 "Th $^{\text{III}}$ starting material, which was difficult without the use of a chelate. The reaction of $\mathrm{Cp}_3^{\ \prime\prime}\mathrm{Th}^{\mathrm{III}},$ Li, and crypt formed dark blue-green needles of $[Li(crypt)][Cp₃''Th^{II}]$ in 83% yield, but the crystals were not suitable for X-ray diffraction, Scheme 2.

Since the reaction chemistry and the spectroelectrochemistry suggested that the Th (n) complexes were generated from a Th(w) precursor through a Th(m) intermediate, reactions with the two-electron reductant Ba were studied. The $Ba(n)/Ba(0)$ reduction potential is nearly identical to that of $K(n)/B$ $K(0).47$ Surprisingly, prolonged stirring of a THF solution of Cp_{3} "Th^{IV}Br and excess Ba afforded only Cp_{3} "Th^{III}. When chelates were added, the reaction of $Cp_3^{\ \prime\prime}Th^WBr$ and crown or $\mathrm{Cp}_3^{\ \ \prime\prime}\mathrm{Th}^{\mathrm{IV}}\mathrm{Br}$ and crypt over excess Ba formed $\mathrm{Cp}_3^{\ \ \prime\prime}\mathrm{Th}^{\mathrm{III}}$ and then the dark green color of $Th(\pi)$ with UV-visible spectra consistent with $[Cp_3^{\ \prime\prime}Th^{II}]^{1-}$. Addition of elemental Hg did not appear to affect the rate of formation of the Th (n) species. These results, coupled with the spectroelectrochemical measurements, strongly suggest that the $Th(w)/Th(u)$ redox couple is not observed experimentally in these systems and that instead two one-electron processes occur.

Fig. 9 Thermal ellipsoid plot of $[Na(\kappa^6\text{-}crown)(\kappa^2\text{-}crown)][Cp_3''Th^I]$ plotted at the 35% probability level. Hydrogen atoms and disorder in the $\kappa^2\text{-}c$ crown unit have been removed for clarity.

Discussion

$An(w)/An(m)$ processes

The trends observed in the $U(w)/U(m)$ and Th $(w)/Th(m)$ redox couples in Tables $1-3$ indicate that Cp^{tet} is more electron donating than Cp $^{\prime}$, which is more electron donating than Cp $^{\prime\prime}$. This follows the electron-donating ability of the ligands previously found in studies of $(C_5R_5)_2Zr(CO)_2$ complexes²¹ and yttrium compounds.22,24 For the zirconium complexes, the CO stretching frequency and the reduction potentials were analyzed to determine electron-donation strength of the cyclopentadienyl ligand. Generally in these $An(w)/An(m)$ studies, the thorium complexes showed less reversible processes than the uranium compounds. In the Cp_{3} "Th $^{\text{IV}}$ Br case, UV-visible spectroelectrochemistry measurements show that this compound is reduced under electrochemical conditions to $\mathbf{Cp}_3{''}\mathbf{Th}^{\text{III}}$, which requires loss of Br^{1-} and geometric reorganization. In the $\text{Cp}_{3}^{}\text{Th}^{\text{IV}}\text{Br}$ case, density functional theory calculations have shown that the putative initial reduction product, $[Cp_3'Th^{III}Br]$ ¹; would be unstable with respect to

 $\text{Cp}_3^{\prime} \text{Th}^{\text{III}}$ and Br^{1-18} These results are consistent with the electrochemical irreversibility of the system.

$An(m)/An(n)$ processes

To our knowledge, only two other $U(m)/U(n)$ couples have been assigned via electrochemistry: $[(\text{Ad}, \text{Me} A\text{rO})_3\text{mes}]U^{\text{III}}$ at -2.495 V using $\binom{n}{b}u_4N[\Pr_6]^{16}$ and $(C_5{}^{i}Pr_5)_2U^{\text{II}}$ at -2.33 V using $\binom{n}{b}u_4N$ [BPh₄].⁹ The -2.26 V value for Cp₃[']U^{III} matches well with these two data points, even though $[Cp_3'U^{II}]^{1-}$ and $(C_5{}^{i}Pr_5)_2U^{II}$ have been assigned $5f^36d^1$ electron configurations,^{1,9} while $\{[(\text{Ad}, \text{Me} \text{ArO})_3 \text{mes}] \text{U}^{\text{II}} \}^{\text{1}-}$ is best described as 5f⁴.⁷ The -2.73 V reduction potential for Cp_{3} "U^{III} is unexpectedly more reducing than those of these other three complexes. This is also unusual in that solutions of $[Cp_3''U^{II}]^{1-}$ have longer lifetimes than solutions of $[Cp_3'U^{II}]^{1-.27}$ The U(III)/U(II) reduction potential for $Cp_3^{\text{tet}}U^{\text{III}}$ was determined to be -3.11 V, which is the most negative reduction potential for these compounds and matches the trend observed for the $An(w)/An(m)$ couples.

Th (n) complexes were investigated for the first time via electrochemistry and the $E_{1/2}$ values for the Th(III)/Th(II) couple

observed in the isolated Th (n) compounds matched the value observed in Cp_3 "Th^{III}. Surprisingly, the Th(IV)/Th(III) couple of Cp_3 "Th^{IV}Br appears to be about the same as the value for the Th(III)/Th(II) couple of $[Cp_3$ "Th^{II}]¹⁻. This result was tested chemically and it was found that reduction of $Th(w)$ with excess reducing agent would form $\text{Th}(\text{II})$ compounds directly with KC_8 , Na, Li, and Ba both with and without the use of a chelating agent. Blue Cp_{3} "Th $^{\text{III}}$ is observed as an intermediate in these reactions which indicates formation of the Th (n) products arises from two one-electron reductions. Furthermore, the $E_{1/2}$ values for $\text{Th}(\text{III})/\text{Th}(\text{II})$ match the expected trend compared to uranium based on previously calculated $An(m)/An(n)$ reduction potentials.⁴⁸–⁵⁰ Edge Article

observed in the isolated Distribution on production and the state of the stat

The thorium electrochemistry was also unusual in that electrochemical data were obtained using $\rm \lbrack^{n}Bu_{4}N\rbrack\lbrack PF_{6}]$ as supporting electrolyte on isolated Th(iv), Th(m), and Th(u) compounds. This electrolyte has proven to be more reactive than $\binom{n}{1}$ Bu₄N][BPh₄] with some complexes^{11,15} and it may have been expected that $Th(II)$ would react with it. The fact that the $Th(m)/Th(n)$ reduction potentials vary slightly depending on the specific electrolyte highlights the fact the reduction potentials of these systems are very sensitive to experimental conditions.

Conclusion

Electrochemical data on three series of tris(cyclopentadienyl) An(IV), An(III), and An(II) (An = Th, U) complexes, including the first data on $Th(\pi)$ complexes, complimented by UV-visible spectroelectrochemical measurements, show a direct correlation between reduction potential and the electron-donating ability of the cyclopentadienyl ring. The studies indicate that Th(III) is a stronger reductant than U(III), but the reduction potential of $U(\Pi)$ is similar to that of Th (Π) . Two unexpected results should stimulate further studies. The $U(m)/U(n)$ reduction potential of $\text{Cp}_{3}^{\prime} \text{U}^{\text{III}}$ is similar to the two previously reported $U(m)/U(n)$ values, but it is significantly less negative than the Cpⁿ analog. The reduction potentials of Th $(w)/Th(m)$ and $Th(m)/Th(n)$ couples are sufficiently similar that $Th(n)$ complexes can be made directly from $\text{Th}(w)$ precursors without the need to isolate the $Th(m)$ intermediate.

Author contributions

J. C. W. synthesized all compounds and performed the cyclic voltammetry experiments. J. C. W. and J. M. B. performed the spectroelectrochemistry experiments. J. W. Z. analyzed the X-ray diffraction data. All authors analyzed the electrochemistry data. J. C. W. and W. J. E. wrote the manuscript with input from all authors.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 M. R. MacDonald, M. E. Fieser, J. E. Bates, J. W. Ziller, F. Furche and W. J. Evans, Identification of the $+2$ Oxidation State for Uranium in a Crystalline Molecular Complex, $[K(2.2.2-Cryptand)][(C_5H_4SiMe_3)_3U]$, *J. Am. Chem.* Soc., 2013, 135, 13310–13313, DOI: 10.1021/ja406791t.
- 2 R. R. Langeslay, M. E. Fieser, J. W. Ziller, F. Furche and W. J. Evans, Synthesis, Structure, and Reactivity of Crystalline Molecular Complexes of the ${[C_5H_3(SiMe_3)_2]_3Th}^{1-}$ Anion Containing Thorium in the Formal +2 Oxidation State, Chem. Sci., 2015, 6, 517–521, DOI: 10.1039/C4SC03033H.
- 3 J. Su, C. J. Windorff, E. R. Batista, W. J. Evans, A. J. Gaunt, M. T. Janicke, S. A. Kozimor, B. L. Scott, D. H. Woen and P. Yang, Identification of the Formal +2 Oxidation State of Neptunium: Synthesis and Structural Characterization of ${Np}^{II}[C_5H_3(SiMe_3)_2]_3\}^{1-}$, J. Am. Chem. Soc., 2018, 140, 7425-7428, DOI: 10.1021/jacs.8b03907.
- 4 M. S. Dutkiewicz, J. H. Farnaby, C. Apostolidis, E. Colineau, O. Walter, N. Magnani, M. G. Gardiner, J. B. Love, N. Kaltsoyannis, R. Caciuffo and P. L. Arnold, Organometallic Neptunium(m) Complexes, Nat. Chem., 2016, 8, 797–802, DOI: 10.1038/nchem.2520.
- 5 M. S. Dutkiewicz, C. Apostolidis, O. Walter and P. L. Arnold, Reduction Chemistry of Neptunium Cyclpentadienide Complexes: From Structure to Understanding, Chem. Sci., 2017, 8, 2553–2561, DOI: 10.1039/c7sc00034k.
- 6 C. J. Windorff, G. P. Chen, J. N. Cross, W. J. Evans, F. Furche, A. J. Gaunt, M. T. Janicke, S. A. Kozimor and B. L. Scott, Identification of the Formal +2 Oxidation State of Plutonium: Synthesis and Characterization of $\{Pu^{II}[C_5H_3(SiMe_3)_2]_3\}^{-}$, J. Am. Chem. Soc., 2017, 139, 3970-3973, DOI: 10.1021/jacs.7b00706.
- 7 H. S. La Pierre, A. Scheurer, F. W. Heinemann, W. Hieringer and K. Meyer, Synthesis and Characterization of a Uranium (n) Monoarene Complex Supported by d Backbonding, Angew. Chem., Int. Ed., 2014, 53, 7158– 7162, DOI: 10.1002/anie.201402050.
- 8 B. S. Billow, B. N. Livesay, C. C. Mokhtarzadeh, J. Mccracken, M. P. Shores, J. M. Boncella and A. L. Odom, Synthesis and Characterization of a Neutral $U(n)$ Arene Sandwich Complex, J. Am. Chem. Soc., 2018, 140, 17369–17373, DOI: 10.1021/jacs.8b10888.
- 9 F. S. Guo, N. Tsoureas, G. Z. Huang, M. L. Tong, A. Mansikkamäki and R. A. Layfield, Isolation of a Perfectly

Linear Uranium(II) Metallocene, Angew. Chem., Int. Ed., 2020, 59, 2299–2303, DOI: 10.1002/anie.201912663.

- 10 K. C. Jantunen, C. J. Burns, I. Castro-Rodriguez, R. E. Da Re, J. T. Golden, D. E. Morris, B. L. Scott, F. L. Taw and J. L. Kiplinger, Thorium(iv) and Uranium(IV) Ketimide Complexes Prepared by Nitrile Insertion into Actinide-Alkyl and -Aryl Bonds, Organometallics, 2004, 23, 4682–4692, DOI: 10.1021/om0343824.
- 11 D. E. Morris, R. E. Da Re, K. C. Jantunen, I. Castro-Rodriguez and J. L. Kiplinger, Trends in Electronic Structure and Redox Energetics for Early-Actinide Pentamethylcyclopentadienyl Complexes, Organometallics, 2004, 23, 5142–5153, DOI: 10.1021/om049634v.
- 12 E. J. Schelter, J. M. Veauthier, J. D. Thompson, B. L. Scott, K. D. John, D. E. Morris and J. L. Kiplinger, 4f-5f Heterotrimetallic Complexes Exhibiting Electrochemical and Magnetic Communication, J. Am. Chem. Soc., 2006, 128, 2198–2199, DOI: 10.1021/ja057808+.
- 13 K. A. Erickson, B. D. Kagan, B. L. Scott, D. E. Morris and J. L. Kiplinger, Revisiting the Bis(Dimethylamido) Metallocene Complexes of Thorium and Uranium: Improved Syntheses, Structure, Spectroscopy, and Redox Energetics of $(C_5Me_5)_2An(NMe_2)_2$ (An = Th, U), Dalton Trans., 2017, 46, 11208–11213, DOI: 10.1039/c7dt02373a.
- 14 Z. E. Button, J. A. Higgins, M. Suvova, F. G. N. Cloke and S. M. Roe, Mixed Sandwich Thorium Complexes Incorporating Bis(Tri-Isopropylsilyl)Cyclooctatetraenyl and Pentamethylcyclopentadienyl Ligands: Synthesis, Structure and Reactivity, Dalton Trans., 2015, 44, 2588–2596, DOI: 10.1039/c4dt02362e.
- 15 C. J. Inman and F. G. N. Cloke, The Experimental Determination of $Th(w)/Th(m)$ Redox Potentials in Organometallic Thorium Complexes, Dalton Trans., 2019, 48, 10782–10784, DOI: 10.1039/c9dt01553a.
- 16 H. S. La Pierre, H. Kameo, D. P. Halter, F. W. Heinemann and K. Meyer, Coordination and Redox Isomerization in the Reduction of a Uranium (m) Monoarene Complex, Angew. Chem., Int. Ed., 2014, 53, 7154–7157, DOI: 10.1002/ anie.201402048.
- 17 J. A. Hlina, J. R. Pankhurst, N. Kaltsoyannis and P. L. Arnold, Metal-Metal Bonding in Uranium-Group 10 Complexes, J. Am. Chem. Soc., 2016, 138, 3333–3345, DOI: 10.1021/ jacs.5b10698.
- 18 J. C. Wedal, S. Bekoe, J. W. Ziller, F. Furche and W. J. Evans, In Search of Tris(Trimethylsilylcyclopentadienyl) Thorium, Dalton Trans., 2019, 48, 16633–16640, DOI: 10.1039/ C9DT03674A.
- 19 W. J. Evans, Tutorial on the Role of Cyclopentadienyl Ligands in the Discovery of Molecular Complexes of the Rare-Earth and Actinide Metals in New Oxidation States, Organometallics, 2016, 35, 3088–3100, DOI: 10.1021/ acs.organomet.6b00466.
- 20 D. H. Woen and W. J. Evans, Expanding the +2 Oxidation State of the Rare-Earth Metals, Uranium, and Thorium in Molecular Complexes, in Handbook on the Physics and Chemistry of Rare Earths, Elsevier B.V., 2016, pp. 1–57, DOI: 10.1016/bs.hpcre.2016.08.002.
- 21 C. E. Zachmanoglou, A. Docrat, B. M. Bridgewater, G. Parkin, C. G. Brandow, J. E. Bercaw, C. N. Jardine, M. Lyall, J. C. Green and J. B. Keister, The Electronic Influence of Ring Substituents and Ansa Bridges in Zirconocene Complexes as Probed by Infrared Spectroscopic, Electrochemical, and Computational Studies, J. Am. Chem. Soc., 2002, 124, 9525–9546, DOI: 10.1021/ja020236y. Chemical Science

Lines United Netallowski, Article. Published on 07 May 2021. The Science Article is a National Common Com
	- 22 J. F. Corbey, D. H. Woen, C. T. Palumbo, M. E. Fieser, J. W. Ziller, F. Furche and W. J. Evans, Ligand Effects in the Synthesis of Ln^{2+} Complexes by Reduction of Tris(Cyclopentadienyl) Precursors Including C–H Bond Activation of an Indenyl Anion, Organometallics, 2015, 34, 3909–3921, DOI: 10.1021/acs.organomet.5b00500.
	- 23 S. A. Moehring, M. Beltran-Leiva, D. Paez-Hernandez, R. Arratia-Perez, J. W. Ziller and W. J. Evans, Rare-Earth Metal(II) Aryloxides: Structure, Synthesis, and EPR Spectroscopy of $[K(2.2.2-Cryptand)][Sc(OC₆H₂^tBu₂-2,6-Me-$ 4)3], Chem.–Eur. J., 2018, 24, 18059–18067, DOI: 10.1002/ chem.201803807.
	- 24 S. A. Moehring and W. J. Evans, Evaluating Electron Transfer Reactivity of Rare-Earth Metal (n) Complexes Using EPR Spectroscopy, Organometallics, 2020, 39, 1187–1194, DOI: 10.1021/acs.organomet.9b00837.
	- 25 J. G. Brennan, R. A. Andersen and A. Zalkin, Chemistry of Trivalent Uranium Metallocenes: Electron-Transfer Reactions with Carbon Disulfide. Formation of $[(RC5H₄)₃U]₂[\mu-\eta¹,\eta¹-CS₂],$ Inorg. Chem., 1986, 25, 1756– 1760, DOI: 10.1021/ic00231a007.
	- 26 M. del Mar Conejo, J. S. Parry, E. Carmona, M. Schultz, J. G. Brennann, S. M. Beshouri, R. A. Andersen, R. D. Rogers, S. Coles and M. Hursthouse, Carbon Monoxide and Isocyanide Complexes of Trivalent Uranium Metallocenes, Chem.–Eur. J., 1999, 5, 3000–3009.
	- 27 C. J. Windorff, M. R. MacDonald, K. R. Meihaus, J. W. Ziller, J. R. Long and W. J. Evans, Expanding the Chemistry of Molecular U^{2+} Complexes: Synthesis, Characterization, and Reactivity of the ${[C_5H_3(SiMe_3)_2]_3U}^-$ Anion, Chem.–Eur. J., 2016, 22, 772–782, DOI: 10.1002/chem.201503583.
	- 28 C. Clappe, D. Leveugle, D. Hauchard and G. Durand, Electrochemical Studies of Tricyclopentadienyl Uranium Iv Chloride Complexes: $(RCp)_3UCl (RCp = RC_5H_4 with R = H;$ Me: CH_3 ; t Bu: $(CH_3)_3C$; TMS: $(CH_3)_3Si$) Evidence of a Disproportionation Mechanism in Oxidation, J. Electroanal. Chem., 1998, 448, 95–103, DOI: 10.1016/S0022- 0728(98)00029-1.
	- 29 D. Hauchard, M. Cassir, J. Chivot and M. Ephritikhine, Electrochemical Study of Uranium (w) and Uranium (w) Organometallic Compounds in Tetrahydrofuran by Means of Conventional Microelectrodes and Ultramicroelectrodes. Part I. Application to the Na(Hg) Reduction of Cp_3UCl (Cp $= \eta - C_5H_5$, *J. Electroanal. Chem.*, 1991, 313, 227-241, DOI: 10.1016/0022-0728(91)85182-O.
	- 30 R. G. Finke, G. Gaughan and R. Voegeli, Organoactinide Electrochemistry. A Cyclic Voltammetric and Coulometric Study of $(C_5Me_5)_2UCl_2$, $[(C_5Me_5)_2UCl_2 \cdot THF]^-Na^+$, $(C_5Me_5)_2UCl$ THF and $(C_5Me_5)_2ThCl_2$, J. Organomet. Chem., 1982, 229, 179–184, DOI: 10.1016/S0022-328X(00)90280-8.
- 31 P. L. Watson, T. H. Tulip and I. Williams, Defluorination of Perfluoroolefins by Divalent Lanthanoid Reagents: Activating C–F Bonds, Organometallics, 1990, 9, 1999–2009, DOI: 10.1021/om00157a006.
- 32 O. Maury, M. Ephritikhine, M. Nierlich, M. Lance and E. Samuel, Chloride Ion Transfer during the $U(w)/U(m)$ Reduction of UCl₄ in Tetrahydrofuran (THF), Studied by Cyclic Voltammetry; Synthesis and Molecular Structure of ["Bu₄][UCl₅(THF)], Inorg. Chim. Acta, 1998, 279, 210-216, DOI: 10.1016/s0020-1693(98)00126-1.
- 33 P. C. Blake, M. F. Lappert, J. L. Atwood and H. Zhang, The Synthesis and Characterisation, Including X-Ray Diffraction Study, of $[Th{n-C₅H₃(SiMe₃)₂}]$; The First Thorium(III) Crystal Structure, J. Chem. Soc., Chem. Commun., 1986, 453, 1148–1149, DOI: 10.1039/ C39860001148.
- 34 P. C. Blake, N. M. Edelstein, P. B. Hitchcock, W. K. Kot, M. F. Lappert, G. V. Shalimoff and S. Tian, Synthesis, Properties and Structures of the Tris(Cyclopentadienyl) Thorium(iii) Complexes $\left[\text{Th}\{\eta^5-\text{C}_5\text{H}_3\text{(SiMe}_2\text{R})_2\text{-}1,3\}_3\right]$ $\left(\text{R} =$ Me or ^tBu), *J. Organomet. Chem.*, 2001, **636**, 124-129, DOI: 10.1016/S0022-328X(01)00860-9.
- 35 C. J. Windorff, M. R. MacDonald, J. W. Ziller and W. J. Evans, Trimethylsilylcyclopentadienyl (Cp′) Uranium Chemistry: Synthetic and Structural Studies of $Cp_4'U$ and $Cp_3'UX$ $(X =$ Cl, I, Me), Z. Anorg. Allg. Chem., 2017, 643, 2011–2018, DOI: 10.1002/zaac.201700323.
- 36 N. A. Siladke, C. L. Webster, J. R. Walensky, M. K. Takase, J. W. Ziller, D. J. Grant, L. Gagliardi and W. J. Evans, Actinide Metallocene Hydride Chemistry: C–H Activation in Tetramethylcyclopentadienyl Ligands to Form $[\mu-\eta^5]$ $C_5Me_3H(CH_2)-\kappa C]^2$ ⁻ Tuck-over Ligands in a Tetrathorium Octahydride Complex, Organometallics, 2013, 32, 6522– 6531, DOI: 10.1021/om4008482. Edge Article

11 P. L. Watson, T. I. Tulip and I. Wilhelms, Dellucionismic of Supercence (Creeks), Theoretical Creative Commons Article is likensed under the model on 12.

10. L. May 2021. Downloaded under a common and th
	- 37 C. J. Kuehl, R. E. Da Re, B. L. Scott, D. E. Morris and K. D. John, Toward New Paradigms in Mixed-Valency: Ytterbocene-Terpyridine Charge-Transfer Complexes, Chem. Commun., 2003, 3, 2336–2337, DOI: 10.1039/ b306484k.
	- 38 R. E. Da Re, C. J. Kuehl, M. G. Brown, R. C. Rocha, E. D. Bauer, K. D. John, D. E. Morris, A. P. Shreve and J. L. Sarrao, Electrochemical and Spectroscopic Characterization of the Novel Charge-Transfer Ground State in Diimine Complexes of Ytterbocene, Inorg. Chem., 2003, 42, 5551–5559, DOI: 10.1021/ic030069i.
	- 39 A. Formanuik, A.-M. Ariciu, F. Ortu, R. Beekmeyer, A. Kerridge, F. Tuna, E. J. L. Mcinnes and D. P. Mills, Actinide Covalency Measured by Pulsed Electron Paramagnetic Resonance Spectroscopy, Nat. Chem., 2016, 9, 578–583, DOI: 10.1038/nchem.2692.
	- 40 R. R. Langeslay, G. P. Chen, C. J. Windorff, A. K. Chan, J. W. Ziller, F. Furche and W. J. Evans, Synthesis,

Structure, and Reactivity of the Sterically Crowded $Th³⁺$ Complex $(C_5Me_5)_3$ Th Including Formation of the Thorium Carbonyl, $[(C_5Me_5)_3Th(CO)][BPh_4]$, *J. Am. Chem. Soc.*, 2017, 139, 3387–3398, DOI: 10.1021/jacs.6b10826.

- 41 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, Chemical Structure and Bonding in a Thorium (m) -Aluminum Heterobimetallic Complex, Chem. Sci., 2018, 9, 4317–4324, DOI: 10.1039/c8sc01260a.
- 42 D. N. Huh, S. Roy, J. W. Ziller, F. Furche and W. J. Evans, Isolation of a Square-Planar $Th(m)$ Complex: Synthesis and Structure of $[\text{Th}(\text{OC}_6\text{H}_2^{\text{t}}\text{Bu}_2\text{-}2, 6\text{-Me-4})_4]^{1-}$, J. Am. Chem. Soc., 2019, 141, 12458–12463, DOI: 10.1021/jacs.9b04399.
- 43 J. S. Parry, F. G. N. Cloke, S. J. Coles and M. B. Hursthouse, Synthesis and Characterization of the First Sandwich Complex of Trivalent Thorium: A Structural Comparison with the Uranium Analogue, *J. Am. Chem. Soc.*, 1999, 121, 6867–6871, DOI: 10.1021/ja9903633.
- 44 J. R. Walensky, R. L. Martin, J. W. Ziller and W. J. Evans, Importance of Energy Level Matching for Bonding in Th^{3+} Am³⁺ Actinide Metallocene Amidinates, $(C_5Me_5)_2[^{i}$ PrNC(Me)NⁱPr]An, *Inorg. Chem.*, 2010, 49, 10007-10012, DOI: 10.1021/ic1013285.
- 45 R. R. Langeslay, M. E. Fieser, J. W. Ziller, F. Furche and W. J. Evans, Expanding Thorium Hydride Chemistry Through Th^{2+} , Including the Synthesis of a Mixed-Valent Th^{4+}/Th^{3+} Hydride Complex, *J. Am. Chem. Soc.*, 2016, 138, 4036–4045, DOI: 10.1021/jacs.5b11508.
- 46 D. N. Huh, J. W. Ziller and W. J. Evans, Chelate-Free Synthesis of the U(II) Complex, $[(C_5H_3(SiMe_3)_2)_3U]^{1-}$, Using Li and Cs Reductants and Comparative Studies of $La(II)$ and Ce(II) Analogs, *Inorg. Chem.*, 2018, 57, 11809-11814, DOI: 10.1021/acs.inorgchem.8b01966.
- 47 CRC Handbook of Chemistry and Physics, ed. W. M. Haynes, D. R. Lide and T. J. Bruno, CRC Press, 97th edn, 2016.
- 48 L. R. Morss, Comparative Thermochemical and Oxidation-Reduction Properties of Lanthanides and Actinides, in Handbook on the Physics and Chemistry of Rare Earths, ed. K. A. Gschneider Jr, L. Eyring, G. R. Choppin and G. H. Lander, Elsevier Science, Amsterdam, 1994.
- 49 S. G. Bratsch and J. J. Lagowski, Actinide Thermodynamic Predictions. 3. Thermodynamics of Compounds and Aquo Ions of the 2+, 3+, and 4+ Oxidation States and Standard Electrode Potentials at 298.15 K, J. Phys. Chem., 1986, 90, 307–312, DOI: 10.1021/j100274a021.
- 50 Q.-Y. Wu, J.-H. Lan, C.-Z. Wang, Z.-P. Cheng, Z.-F. Chai, J. K. Gibson and W.-Q. Shi, Paving the Way for the Synthesis of a Series of Divalent Actinide Complexes: A Theoretical Perspective, Dalton Trans., 2016, 45, 3102–3110, DOI: 10.1039/C5DT04540A.