

A Crystallizable Dinuclear Tuck-In-Tuck-Over Tuck-Over Dialkyl Tren Uranium Complex and Double Dearylation of BPh_4^- To Give the BPh_2^- -Functionalized Metallocycle $[\text{U}\{\text{N}(\text{CH}_2\text{CH}_2\text{NSiMe}_3)_2(\text{CH}_2\text{CH}_2\text{NSiMe}_2\text{CHBPh}_2)\}(\text{THF})](\text{THF})]$

Benedict M. Gardner, Jonathan McMaster, William Lewis, Alexander J. Blake, and Stephen T. Liddle*

School of Chemistry, University of Nottingham, University Park, Nottingham, NG7 2RD, U.K.

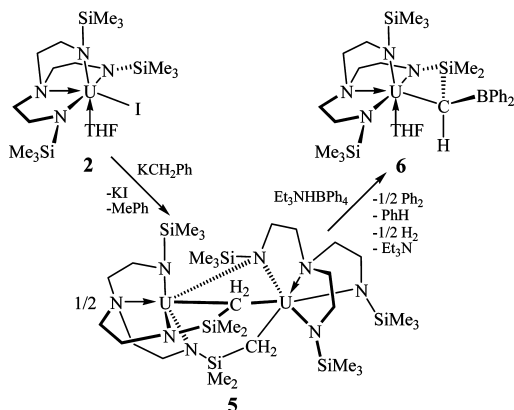
Received June 2, 2009; E-mail: stephen.liddle@nottingham.ac.uk

Cyclopentadienyl-based ligands, and the pentamethylcyclopentadienyl ligand in particular, have been enormously successful at supporting novel reactivity patterns in actinide chemistry.¹ In recent years a number of groups have investigated the use of nonmetallocene ligands with uranium(III/IV) chemistry and novel and diverse reactivity profiles have emerged.²

Recently, we have investigated the capacity of the Tren^{TMS} ligand $\{\text{N}(\text{CH}_2\text{CH}_2\text{NSiMe}_3)_3\}$ to support novel uranium(IV)–metal bonds. We have reported the syntheses of $[(\text{Tren}^{\text{TMS}})\text{U}(\text{X})(\text{THF})]$ ($\text{X} = \text{Cl}$, **1**;³ $\text{X} = \text{I}$, **2**⁴) and demonstrated their utility in the preparation of the first structurally authenticated uranium–gallium³ and –rhenium bonds.⁴ However, in preliminary reactions with some transition metal anions, we have noted that KX elimination is not straightforward, and thermolysis is required.

We targeted $[(\text{Tren}^{\text{TMS}})\text{U}(\text{THF})_2][\text{BPh}_4^-]$ (**3**) as a precursor since we reasoned the BPh_4^- anion would be more labile than coordinated halides, and KBPh_4 elimination is a proven synthetic method in f-element chemistry.⁵ Since KBPh_4 does not react with **1** or **2** in THF, we anticipated that treatment of **2** with $\text{KCH}_2\text{C}_6\text{H}_5$ would give the metallocycle $[\text{U}\{\text{N}(\text{CH}_2\text{CH}_2\text{NSiMe}_3)_2(\text{CH}_2\text{CH}_2\text{NSiMe}_2\text{CH}_2)\}(\text{THF})]$ (**4**)⁶ which would undergo protonolysis with $\text{Et}_3\text{NHBPh}_4$ to afford **3**. Herein, we show that this superficially straightforward chemistry is far more complex, as evidenced by the unprecedented formation of a dinuclear tuck-in-tuck-over tuck-over dialkyl Tren–uranium(IV) complex, and the first example of *double* dearylation of BPh_4^- in a molecular context to give a BPh_2^- -functionalized metallocycle.

Scheme 1. Synthesis of **5** and **6**



Reaction of **2** with $\text{KCH}_2\text{C}_6\text{H}_5$ proceeds cleanly in toluene to reproducibly give complex **5**, Scheme 1, isolated as yellow crystals from hexane in 54% crystalline yield. A variable-temperature ^1H NMR study and X-ray crystallography enabled us to conclusively identify **5** as a dinuclear tuck-in-tuck-over tuck-over dialkyl,⁷ which is further supported by FTIR and CHN data.⁸ Monitoring the reaction by ^1H NMR spectroscopy showed the smooth conversion of **2** to **5** with concomitant formation of toluene

within minutes. Intermediates were not observed, suggesting the decomposition of the putative benzyl derivative of **2** to **4** and the H^+ transfer/dimerization to form **5** are rapid.⁹

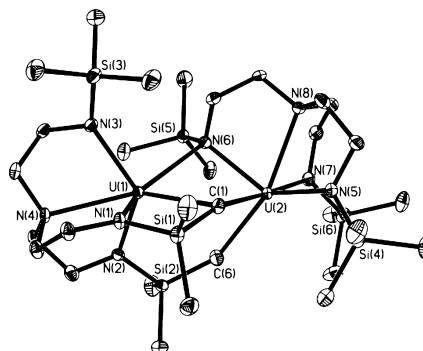


Figure 1. Molecular structure of **5**. Thermal ellipsoids set at 30% probability; hydrogen atoms omitted for clarity. Selected bond lengths (Å): U(1)–N(1) 2.239(3), U(1)–N(2) 2.257(3), U(1)–N(3) 2.292(3), U(1)–N(4) 2.677(3), U(1)–N(6) 2.738(3), U(1)–C(1) 2.667(5), U(2)–N(5) 2.296(4), U(2)–N(6) 2.381(3), U(2)–N(7) 2.267(3), U(2)–N(8) 2.616(3), U(2)–C(1) 2.669(4), U(2)–C(6) 2.493(5).

The molecular structure of **5** is illustrated in Figure 1 with selected bond lengths. The U(2)–Tren ligand is coordinated normally, except for the bridging N(6) center. Bridging Tren amides are known but usually result from alkali metal occlusion.¹⁰ However, the coordination of the U(1)–Tren ligand is unprecedented. In addition to the three anionic amides, two trimethylsilyl groups are metalated. The C(1) center bridges U(1) and U(2) in a tuck-in-tuck-over coordination mode, with essentially identical U–C bond distances of 2.667(5) and 2.669(5) Å. In contrast, terminal C(6) binds in a tuck-over manner with a significantly shorter C(6)–U(2) bond length of 2.493(5) Å. The U–C bond distances compare well to the small number of related metallocyclic uranium(IV)–alkyls.^{6b,c} The U– N_{amide} and N_{amine} bond distances are typical of U(IV)–N bond lengths¹¹ and are commensurate with their binding modes.

Treatment of **5** with 2 equiv of $\text{Et}_3\text{NHBPh}_4$ does not give **3**. Instead, the tuck-in metallocycle **6** was isolated as pale green crystals in 52% crystalline yield, Scheme 1, and the characterization data support its formulation.⁸

The molecular structure of **6** is depicted in Figure 2 with selected bond lengths. The U(1)–C(1) bond distance of 2.644(9) Å compares well to **5** and related metallocyclic uranium(IV)–alkyls.⁶ The U– N_{amido} and U– N_{amine} bond lengths of 2.244(6) (av.) and 2.573(6) Å are typical of U(IV)–N bond distances.¹¹ The U(1)–N(1) bond is short at 2.193(6) Å, and the bite angle of the tuck-in arm is acute at $68.7(2)^\circ$ [cf. $86.41(17)^\circ$ in **5**]. The boron center is trigonal planar [$\Sigma\angle = 360^\circ$] and the B(1)–C(1) distance of 1.493(11) Å is short, suggesting B–C multiple bond character. For comparison, average B– CH_2 and B– C_{Ph} bond lengths of 1.444 and 1.576 Å were reported for $[\text{Mes}_2\text{BCH}_2][\text{Li}(12\text{-crown-4})_2]$ ¹² and BPh_3 .¹³ To further

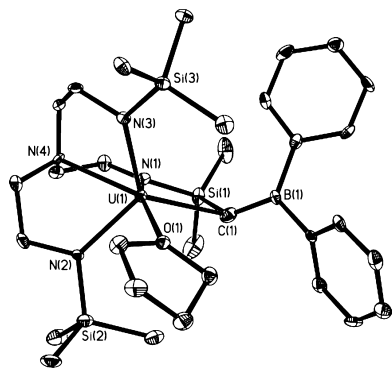
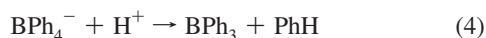
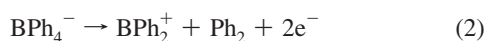
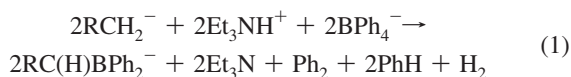


Figure 2. Molecular structure of **6**. Thermal ellipsoids set at 30% probability; hydrogen atoms omitted for clarity. Selected bond lengths (Å): U(1)–N(1) 2.193(6), U(1)–N(2) 2.262(6), U(1)–N(3) 2.277(6), U(1)–N(4) 2.573(6), U(1)–C(1) 2.644(9), U(1)–O(1) 2.565(5), B(1)–C(1) 1.493(11), B(1)–C(16) 1.596(11), B(1)–C(22) 1.591(12).

validate **6** and probe the B(1)–C(1) bond we carried out DFT calculations on a full model of **6**.⁸ The calculation reproduced the metrical parameters and inspection of the Kohn–Sham orbitals, and Mayer bond orders (B–C = 1.33) confirm the manifestation of a B(1)–C(1) π -bond perturbed by the polarizing uranium center.



To shed light on the formation of **6**, we analyzed the reaction mother liquor using GC–MS, which revealed the presence of benzene and biphenyl.⁸ Thus, the overall reaction can be represented by eq 1. Monitoring the reaction by variable temperature ¹H NMR spectroscopy showed conversion of **5** to **6**, and no intermediates were observed.^{8,14} The stoichiometry of eq 1 suggests that eqs 2–5¹⁵ should be considered (S = solvent): (i) formation of BPh₂⁺, eq 2, appears unlikely but could be facilitated by a redox active uranium center, and this would account for the generation of Ph₂ and BPh₂; (ii) eq 3 is known for BPh₄[−] and accounts for the formation of Ph₂,¹⁶ (iii) attack of BPh₃ by a carbanion center with extrusion of Ph[−] (or PhH) seems unlikely on steric grounds, but this cannot be ruled out;¹⁷ (iv) formation of C₆H₆ may be accounted for with eq 4, point (iii), or direct extrusion of Ph[−] from BPh₄[−] which then abstracts H⁺ from Et₃NH⁺ or the cyclometalated arm in an acid–base reaction;¹⁸ (v) previous electrochemical studies have demonstrated that eq 5 is viable,¹⁵ which would sustain eqs 3 and 4, generate a BPh₂⁺ of sufficient reactivity to allow nucleophilic attack by a carbanion center, and regenerate BPh₄[−] which is a potential source of Ph[−].

The formation of **6** is remarkable and is, as far as we are aware, the first example of double dearylation of BPh₄[−] in a molecular context.¹⁵ The reason why the use of BPh₄[−] as a counteranion is avoided in homogeneous catalysis is open to debate. It is usually assumed that BPh₄[−] can block incoming substrates by weak coordination.¹⁹ The BPh₄[−] anion can also become metalated.²⁰ Monodearylation of BPh₄[−] (eq 3) has been recognized as another potentially detrimental role for BPh₄[−].¹⁶ The double dearylation reactivity of BPh₄[−] described here adds to the growing list of possible reactions that should be contemplated when using BPh₄[−].

To conclude, the unprecedented dinuclear tuck-in-tuck-over tuck-over dialkyl Tren-uranium(IV) complex **5** extends the palate of novel chemistry which may be achieved with uranium and nonmetallocene ligands, and the BPh₂-functionalized complex **6** reveals a new double dearylation reaction for the BPh₄[−] anion.

Acknowledgment. We thank the Royal Society, the EPSRC, the University of Nottingham, and the NSCCS for support and Dr. R. Bourne (Nottingham) for obtaining GC–MS data.

Supporting Information Available: Experimental, X-ray and computational data for **5** and **6**. This material is free of charge via the Internet at <http://pubs.acs.org>.

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