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Nitride protonation and NH₃ binding *versus* N-H bond cleavage in uranium nitrides†

Megan Keener, D Rosario Scopelliti D and Marinella Mazzanti **D**

The conversion of metal nitrides to NH_3 is an essential step in dinitrogen fixation, but there is limited knowledge of the reactivity of nitrides with protons (H⁺). Herein, we report comparative studies for the reactions of H⁺ and NH_3 with uranium nitrides, containing different types of ancillary ligands. We show that the differences in ancillary ligands, leads to dramatically different reactivity. The nitride group, in nitride-bridged cationic and anionic diuranium(IV) complexes supported by $-N(SiMe_3)_2$ ligands, is resistant toward protonation by weak acids, while stronger acids result in ligand loss by protonolysis. Moreover, the basic $-N(SiMe_3)_2$ ligands promote the N-H heterolytic bond cleavage of NH_3 , yielding a "naked" diuranium complex containing three bridging ligands, a nitride (N^3-) and two NH_2 ligands. Conversely, in the nitride-bridged diuranium(IV) complex supported by $-OSi(O^tBu)_3$ ligands, the nitride group is easily protonated to afford NH_3 , which binds the U(IV) ion strongly, resulting in a mononuclear $U-NH_3$ complex, where NH_3 can be displaced by addition of strong acids. Furthermore, the $U-OSi(O^tBu)_3$ bonds were found to be stable, even in the presence of stronger acids, such as NH_4BPh_4 , therefore indicating that $-OSi(O^tBu)_3$ supporting ligands are well suited to be used when acidic conditions are required, such as in the H^+/e^- mediated catalytic conversion of N_2 to NH_3 .

Introduction

Uranium nitrides were identified more than 100 years ago as active catalysts in the conversion of dinitrogen (N_2) to ammonia (NH_3) , and very recently, the stoichiometric conversion of N_2 into NH_3 by molecular uranium complexes was reported. Notably, dinuclear uranium(III) complexes 2a,3 or dinuclear U(IV) complexes combined with an external reducing agent, were capable of carrying out the four electron reduction of N_2 . Full cleavage of N_2 to nitrides has also been reported, 2e,4a,b where the resulting hydrazido and nitride ligands could be further reduced and protonated, yielding stoichiometric amounts of NH_3 . 2,4b

The conversion of metal nitrides to NH₃ is an essential step in dinitrogen fixation, where effective nitride protonation is crucial in building catalytic cycles.⁵ A key challenge in developing catalytic N₂ reduction to NH₃, is the simultaneous addition of reducing agent and acid (H⁺), while maintaining the structure of the catalyst, by avoiding ligand protonation and irreversible binding of NH₃. However, studies addressing the reactivity of nitrides with protonating agents⁶ are limited to d-block metals, with few instances of imido (NH²⁻), amido

Institut des Sciences et Ingénierie Chimiques, École Polytechnique Fédérale de Lausanne (EPFL), 1015 Lausanne, Switzerland. E-mail: marinella.mazzanti@epfl.ch † Electronic supplementary information (ESI) available: Synthetic methods, NMR spectra, crystallographic data. CCDC 2097071, 2097072 and 2097366–2097369. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d1sc03957a

(NH₂⁻), and ammonia (NH₃) bound products being isolated and characterized. Protonation of the ligand⁷ or the metal center,⁸ rather than the nitride, is common, and favoring protonation of the nitride can be challenging. The irreversible binding of NH₃, or its reactivity with the metal complex, may also constitute an important disadvantage in catalyst design. Despite the inherent stability of the N-H bonds in NH₃, activation under mild conditions can occur through ligand–metal cooperation,⁹ where the presence of basic ligands can result in the redox-neutral heterolysis of NH₃ and subsequent ligand protonolysis. Facile N-H activation of NH₃ *via* metal-ligand cooperative addition was also reported in one instance for a uranium complex.¹⁰

Since the first examples of uranium nitrides were isolated in the gas phase¹¹ and in solution, ^{4a,12} uranium nitride complexes have been the subject of an increasing number of reports. Mononuclear, dinuclear, polynuclear, mono- and bis-nitride complexes that contain uranium in oxidation states ranging from (III) to (VI) have been synthesized. 13 Several systems have revealed high reactivity towards small molecules, such as CO₂, CO, H_2 , and N_2 , $^{2a,b,3,13f,i,k,14a-q}$ and demonstrated their ability to promote C-H bond activation. 13m, 15a-d Moreover, addition of excess strong acid to terminal or bridging uranium nitrides, derived from azide or N2, has been reported to yield variable amounts of ammonia (20-100%).2c,4b,13i,l,14o,q However, the isolation and characterization of partially protonated, intermediate species have not been reported thus far. Additionally, the site of first protonation, the parameters controlling the yield, or the fate of the uranium complex have not yet been elucidated.

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Scheme 1 Previously reported dinuclear U^{IV}-nitride complexes A, B, and C and their previous and currently described reactivity.

We reasoned that investigating the protonation of uraniumnitrides, supported by different ancillary ligands, and their interaction with the resulting NH3, could provide insight into the parameters controlling the reactivity. This would allow us to harness significant elements for the design of complexes active in the H⁺/e⁻ mediated catalytic conversion of N₂ to NH₃.

We have previously reported that $-OSi(O^tBu)_3$ are versatile ancillary ligands for the synthesis of bridging and terminal uranium nitrides in oxidation states ranging from (III) to (VI). 13g,i,l,14q,16 Moreover, they have allowed the isolation of nitride and oxide bridged diuranium(III) complexes which promote the stoichiometric conversion of N₂ to NH₃.^{2a,3,4b} We also found that using -OSi(O^tBu)₃ (complex A; Scheme 1)^{14h,i,i}

versus -N(SiMe₃)₂ (B and C)^{13k,m,17,18} as supporting ligands leads to significant differences in reactivity of analogous nitride and oxide bridged diuranium complexes with respect to reducing agents (KC₈), electrophilic molecules such as CO₂, CO, and H₂ (Scheme 1), and N2 reduction.2a,18

Herein, we report the comparative studies for the reactions of H⁺ and NH₃ with the diuranium(w) bridging nitrides, A, B and C. We show that the different ancillary ligands lead to dramatically different reactivity, resulting in the isolation of a stable terminal NH₃ complex (2) and a "naked" diuranium complex containing three bridging ligands, a nitride (N³⁻) and two NH₂ ligands, providing important information on the species that can be formed during conversion of nitride to ammonia.

Results and discussion

Reactivity of OSi(O^tBu)₃-containing complexes

First, we were interested in probing the reactivity of the previously reported anionic bridging nitride (A), supported by -OSi(O^tBu)₃ ligands. The nitride link in this complex has been described to be more reactive towards electrophiles than in the analogous -N(SiMe₃)₂ complex C (Scheme 1) but its reactivity with acids was not investigated.13k

Addition of 1.0 equiv. of NH_4BPh_4 to a solution of **A** in d_8 -THF at -40 °C resulted in the partial consumption of A and appearance of new resonances in the ¹H NMR spectrum (Fig. S1†). While some resonances remain unidentified, two sets resonances were assigned to the $[(U^{IV}(OSi(O^tBu)_3)_3(THF))_2(\mu-NH)]$ (1), which could be cleanly produced using the weaker acid, HNEt₃BPh₄ (Scheme 2 and the following section), and $[U^{IV}(OSi(O^tBu)_3)_3(THF)_2(NH_3)][BPh_4]$ (2; see below).

Scheme 2 Reactivity of complex A toward various H⁺ (acid) sources.

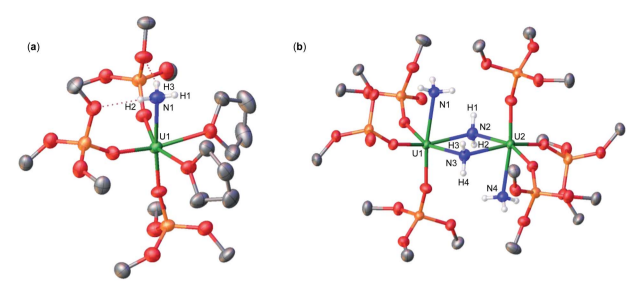


Fig. 1 Molecular structures of (a) $[U^{IV}(OSi(O^tBu)_3)_3(THF)_2(NH_3)][BPh_4]$, 2, and (b) $[(U^{IV}(OSi(O^tBu)_3)_3(NH_3))_2(\mu-NH_2)_2]$, 3, with thermal ellipsoids drawn at the 50% probability level. Hydrogen atoms on the $-OSi(O^tBu)_3$ ligands, methyl groups, and the BPh₄ anion in (a) have been omitted for clarity.

Addition of 3.0 equiv. of NH_4BPh_4 to a solution of A in d_8 -THF resulted in immediate formation of a teal solution and precipitation of a white solid. Crystals of two products were isolated at -40 °C from a concentrated Et₂O solution of the reaction mixture and characterized by XRD analysis as the complexes $[U^{IV}(OSi(O^tBu)_3)_3(THF)_2(NH_3)][BPh_4]$ (2) (Fig. 1a), and $[(U^{IV}(OSi(O^tBu)_3)_3(NH_3))_2(\mu-NH_2)_2]$ (3) (Fig. 1b). Analysis by ¹H NMR spectroscopy of the reaction mixture was consistent with the formation of complexes 2 and 3 in a 1.1:1 ratio (Fig. S2†). Few pure crystals of 3 could be obtained for ¹H NMR spectroscopy, but attempts to isolate larger quantities of pure material remained unsuccessful, resulting in mixtures of both 2 and 3. The 1 H NMR spectrum of complex 3 in d_{8} -THF displays two signals at δ 5.1 and -5.6 ppm, corresponding to the $-OSi(O^tBu)_3$ ancillary ligands, and two resonances at -10.7 and -70.6 ppm assigned to the amido (NH₂¹⁻) and NH₃ resonances, respectively.

Addition of excess (9.0 equiv.) NH_4BPh_4 to a solution of **A** in d_8 -THF, resulted in a pale blue mixture. After 12 hours, the 1H NMR spectrum of this mixture shows only the presence of the signals assigned to complex **2** (Fig. S4†). Large plate, teal crystals suitable for XRD analysis of $[U^{IV}(OSi(O^tBu)_3)_3(THF)_2(NH_3)]$ [BPh₄] (2), were obtained from a concentrated Et₂O solution at -40 °C (Fig. 1b and Scheme 2), in 82% yield. The 1H NMR spectrum of complex **2** in d_8 -THF displays three aromatic BPh₄ phenyl signals and one broad resonance at δ 9.6 ppm, corresponding to the $-OSi(O^tBu)_3$ ancillary ligands. The resonance at -160.1 ppm is assigned to the uranium bound NH_3 .

The clean formation of **2**, from the reaction of **A** with excess NH₄BPh₄ and its isolation in 82% yield, cannot be explained only by the protonation of the nitride, but requires binding of NH₃ released during protonation by NH₄BPh₄. In order to confirm that the NH₃ ligand in **2**, arises both from nitride protonation and from added NH₄BPh₄, we performed the reaction of **A** with isotopically enriched ¹⁵NH₄BPh₄ to yield **2**-¹⁴/

¹⁵N. Adding a solution of HCl in Et₂O to 2-^{14/15}N leads to the formation of both ¹⁵NH₄Cl and ¹⁴NH₄Cl (97% total yield), indicated by a doublet and triplet respectively in the ¹H NMR spectrum. The ratio of ¹⁵NH₄Cl: ¹⁴NH₄Cl is quite large (8:1 ratio), because of the large excess (9.0 equiv.) of ¹⁵NH₄BPh₄ used in the initial synthesis of $\mathbf{A} \rightarrow 2^{-14/15}N$. The NH₃ ligand in 2 binds quite strongly to the U(IV) center, as it could not be removed under dynamic vacuum for a few hours, but is removed upon addition of strong acids.

The isolation of complex 3, when 3.0 equiv. of NH_4BPh_4 are added to **A**, suggests that the formation of **2** proceeds through the bis-amido intermediate **3**, which can be further protonated to yield the mononuclear terminal NH_3 complex. We suggest that the formation of the bis-amido intermediate **3**, involves a putative mono-amido complex, $[(U^{IV}(OSi(O^tBu)_3)_3(NH_3))_2(\mu-NH_2)][BPh_4]$ (**3**_{int}) that undergoes ligand disproportionation to yield **2** and **3** (Scheme S1†).

In order to isolate intermediates with a lower degree of protonation, we investigated the reaction of A with HNEt₃BPh₄, a weaker acid compared to NH₄BPh₄. Addition of 1.0 equiv. of $HNEt_3BPh_4$ to a solution of **A** in d_8 -THF at -40 °C resulted in a green solution and precipitation of a white solid (CsBPh₄). The ¹H NMR spectrum of the reaction mixture reveals the complete consumption of A and the clean formation of a new species (Fig. S5†). Pale green crystals of $[(U^{IV}(OSi(O^tBu)_3)_3)_2(\mu\text{-THF})_2(\mu\text{-}$ NH)] (1) were obtained in 69% yield from a concentrated THF solution at -40 °C (Scheme 2 and Fig. 2). The ¹H NMR spectrum of complex 1 in d_8 -THF displays a broad resonance at δ 0.72, corresponding the $-OSi(O^tBu)_3$ ancillary ligands, and another broad resonance at δ 150.3 ppm assigned to the imido (NH²⁻) group. A similar chemical shift (176.5 ppm) was observed for the imido group in the previously reported bis-imido bridged complex $[K_2{[U(OSi(O^tBu)_3)_3]_2(\mu-NH)_2}].^{13i}$

We also investigated the addition of excess HNEt₃BPh₄ to A, in order to determine if protonation of the imido (NH²⁻) group

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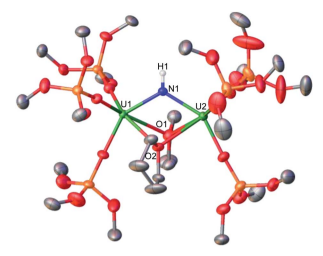


Fig. 2 Molecular structure of $[(U^{IV}(OSi(O^tBu)_3)_3)_2(\mu-THF)_2(\mu-NH)]$ [BPh₄], 1, with thermal ellipsoids drawn at the 50% probability level. Hydrogen atoms and methyl groups on the −OSi(O^tBu)₃ ligands have been omitted for clarity.

in 1, would yield the NH₃ complex, 2. Addition of excess (10 equiv.) HNEt₃BPh₄ to a solution of A in d_8 -THF at -40 °C resulted in the formation of a pale green solution with precipitation of a white solid (CsBPh₄) (Scheme 2). The reaction mixture was brought to room temperature and analysis by ¹H NMR spectroscopy revealed resonances consistent with the presence of 1 and unreacted HNEt₃BPh₄. Further stirring of the solution for an additional 2 days at room temperature resulted in a blue solution, with partial consumption of 1. After 8 days, ¹H NMR spectroscopy indicated the complete consumption of 1 with appearance of the resonance at δ 9.6 ppm, assigned to the NH₃ complex (2) in 40% yield, together with unidentified signals (Fig. S6a-d†). The absence of the resonance at δ -160.1 ppm, assigned to the uranium bound NH₃, is attributed to the fast exchange of the NH3 ligand in the presence of excess HNEt₃BPh₄. We were able to confirm this by addition of 10.0 equiv. $HNEt_3BPh_4$ to a solution of 2 in d_8 -THF. Over 12 hours, we see that the signal corresponding to the uranium bound NH₃ disappears (Fig. S7†). These results indicate that 1 is further protonated to the -NH₃ containing species by excess HNEt₃BPh₄, but requires longer reaction times to yield 2. The formation of 2 (40% yield) in this reaction arises from cleavage

of the U=N=U bridge, which requires the formation of other U-containing products that we were unable to identify.

Structural characterization of OSi(O^tBu)₃-containing complexes

The solid-state molecular structures of complexes 1, 2, and 3 were determined by X-ray diffraction studies. The metrical parameters are presented in Table 1, including those previously reported for complex A.

Complex 1 crystalizes in the space group C2/c with one molecule per asymmetric unit. The solid-state molecular structure of 1 shows the presence of a neutral diuranium(IV) complex where each uranium is bound by three -OSi(O^tBu)₃ ancillary ligands and are bridged by an imido (NH^{2-}) ligand and two THF molecules (Fig. 2). The U-N-U bond angle changes dramatically from linear in A (U-N-U: 170.2(3)°) to bent in 1 (118.3(3)°). The U-N bond distances in 1 are elongated (U1-N1: 2.243(7), U2-N1: 2.219(6) Å) in comparison to A, consistent with a bridging imido (μ-NH).14l

Complex 2 crystalizes in the space group $P\bar{1}$ with one molecule per asymmetric unit. Its solid-state molecular structure shows the presence of an ion pair consisting of one BPh₄ anion and the [U^{IV}(OSi(O^tBu)₃)₃(THF)₂(NH₃)] cation comprising of a terminal UIV-NH3 bound ligand (Fig. 1a). Only two examples of crystallographically characterized U-NH3 complexes have been reported so far: the bis(1,1,1,5,5,5-hexafluoropentane-2,4dionato)NH3-uranyl(v1)19 and a series of trinuclear complexes of formula $[(NH_3)_8U(\mu-N)(NH_3)_3(X)_2U(\mu-N)U(NH_3)_8]Y_n \cdot ZNH_3 (X =$ NH₃, Br⁻, or Cl⁻; n = 6-8; Y = Cl⁻ or Br⁻; Z = 26, 21, or 6).²⁰ Where the U-NH₃ bond distances in these examples, 2.48(6)¹⁹ and 2.605(3)20 respectively, are consistent with the U1-N1 (2.540(4) Å) bond distance in complex 2. Additionally, the U-NH₃ bond distance in 2 is dramatically elongated in comparison to previously reported terminal amido (NH2) complexes, $[U(1,2,4-({}^{t}Bu)_{3}C_{5}H_{2})_{2}(NH_{2})_{2}]$ 2.228(4)Ă), (U-N_{amido}; [U(Tren^{TIPS})(NH₂)] (U- N_{amido} ; Å),21 2.194(5)and $[U(COT^{TIPS2})(Cp^*)(NH_2)]$ $(U-N_{amido}; 2.217(4) Å),^{10}$ further supporting a terminal bound NH3 molecule. The protons of the NH3 ligand can be crystallographically identified, making their assignment unambiguous.

Complex 3 crystalizes in the space group $P\bar{1}$ with 0.5 molecules per asymmetric unit. Its solid-state molecular structure

Table 1 Selected bond lengths (Å) and angles (°) of the previously reported complexes $[Cs(U^{IV}(OSi(O^tBu)_3)_3)_2(\mu-N)]$, A, and $[(U^{IV}(N(SiMe_3)_2)_2(-1)_3]$ THF))₂(μ -NJ)][BPh₄], B, as well as complexes [(U^{IV}(OSi(O^tBu)₃)₃)₂(μ -NH)], 1, [(U^{IV}(OSi(O^tBu)₃)₃(THF))₂(NH₃)][BPh₄], 2, [(U^{IV}(OSi(O^tBu)₃)₃(NH₃))₂(μ -NH)], 1, [(U^{IV}(OSi(O^tBu)₃)₃(THF))₂(NH₃)][BPh₄], 2, [(U^{IV}(OSi(O^tBu)₃)₃(NH₃))₃(NH₃)][BPh₄], 2, [(U^{IV}(OSi(O^tBu)₃)₃(NH₃)](D₃(NH₃))₃(NH₃)₃(NH₃)₃(NH₃)₃(NH₃))₃(NH₃)₃ NH_{2} , 3, $[(U^{IV}(N(SiMe_{3})_{2})(THF)_{3})_{2}(\mu-NH)_{2}][BPh_{4}]_{2}$, 4, and $[(U^{IV}(THF)_{4})_{2}(\mu-N)(\mu-NH_{2})_{2}][BPh_{4}]_{3}$, 5

Complex	A (ref. 14h)	1	2	3	B (ref. 13 <i>k</i>)	4	5
U-N _{nitride} U-N _{imido} U-N _{amido}	_	 NH: 2.243(7); 2.219(6) 		 NH ₃ : 2.608(3)	2.055(3) —	 NH: 2.179(5); 2.198(6) 	2.062(3); 2.018(3) — NH ₂ : 2.449(3); 2.455(3)
U-N-U U-NH-U U-NH ₂ -U	170.2(3) —		_ _ _ _	NH ₂ : 2.451(3); 2.463(2) — — 106.5(1)	168.97(14) — —		107.41(12) — 84.36(9)

can be described as a neutral complex consisting of a dinuclear U^{IV}/U^{IV} core, comprising of two $[U^{IV}(OSi(O^tBu)_3)_3(NH_3)]$ units bridged by two amido (NH_2^{1-}) ligands (Fig. 1b). The U1–N1 and U2–N4 (2.608(3) Å) bond distances are consistent with a terminal NH₃, as found in 2 (U1–N1: 2.540(4) Å). The U1–N2 = 2.451(3) Å and U1–N3 = 2.463(2) Å bond distances are consistent with two bridging amido (NH_2^{1-}) ligands, with delocalized charge on the uranium ions as indicated by the nearly symmetrical U–N_{amido} metrical parameters. The protons of the imido ligands can be crystallographically identified, making their assignment unambiguous.

Next, we investigated protonation of the previously reported dinuclear U^{IV} -nitride complexes bearing more basic $-N(SiMe_3)_2$ ancillary ligands.

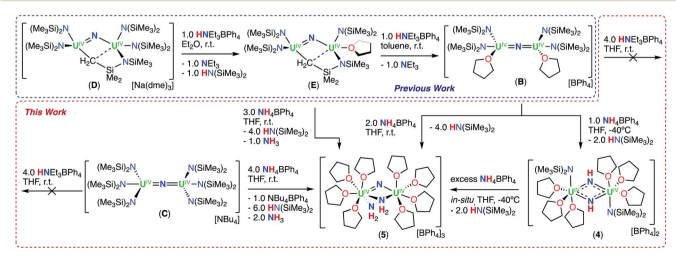
Reactivity of N(SiMe₃)₂-containing complexes

First, we probed the reactivity of complex, **B** with CO₂, CO, and H₂ to compare it with that of complexes **A** and C (Scheme 1). Previous work has shown that **B** has an increased reactivity toward reducing agents in comparison to the all $-N(SiMe_3)_2$, complex C, but the reactivity with CO₂, CO, and H₂ was not explored in the previous report.^{13m} We found that complex **B** reacts with 2.0 equiv. of ¹³CO₂ in d_8 -THF over 4 days, in which **B** is fully consumed and results in a complicated ¹H NMR spectrum (Fig. S11†). Attempts to grow single crystals from the reaction mixture were unsuccessful. However, after the removal of volatiles and hydrolysis with D₂O (pD = 12) of the reaction residue, N¹³CO⁻ was identified by ¹³C NMR spectroscopy (Fig. S12†), similarly to what observed for complex **A**.

Complex **B** reacts with 2.0 equiv. of 13 CO in d_8 -THF over 6 days, in which **B** is fully consumed and results in a complicated 1 H NMR spectrum (Fig. S13†). Attempts to grow crystals from the reaction mixture were unsuccessful. However, after quenching the reaction mixture with D₂O (pD = 12), 13 CN $^-$ was identified by 13 C NMR spectroscopy (Fig. S14†). This indicates that, as previously reported for **A**, which reacts readily with CO to yield [Cs(U(OSi(O^fBu)₃)₃)₂(μ -CN)(μ -O)], 14i the nucleophilic character of the bridging nitride promotes the cleavage and deoxygenation of

Next, we probed the reactivity of cationic bridging nitride (**B**), which was synthesized as reported by our group^{13m} by successive protonolysis of one $-N(SiMe_3)_2$ ligand in the complex **D**, previously reported by Fortier and Hayton, ^{13b} and of the metal-amide methanide bond in **E**, using a total of 2.0 equiv. of HNEt₃BPh₄ (Scheme 3). Addition of excess (4.0 equiv.) HNEt₃BPh₄ to a solution of **B** in d_8 -THF, resulted in unreacted starting materials as seen by ¹H NMR spectroscopy (Fig. S16†). Therefore, we pursued the protonation using the stronger acid, NH₄BPh₄, resulting in different reactivity compared to what was found for the $-OSi(O^tBu)_3$ complex, **A**.

¹H NMR studies showed that addition of 1.0 equiv. of NH₄BPh to a solution of **B** in d_8 -THF resulted in the formation of a new species displaying a resonance at δ 12.6 ppm and unreacted **B** (Fig. S17†). Golden crystals suitable for XRD analysis were obtained from the reaction mixture at -40 °C after 12 hours. The solid-state molecular structure shows the presence of a dinuclear U^{IV}/U^{IV} bridging bis-imido (NH²⁻) complex, [(U^{IV}(N(SiMe₃)₂)(THF)₃)₂(μ-NH)₂][BPh₄]₂ (4) (Fig. 3a and Scheme 3). The product is insoluble in most common solvents including toluene, Et₂O, THF, and *n*-hexanes, but is soluble in pyridine. The ¹H NMR spectrum of the isolated crystals of 4 in d_5 -pyridine, displays a resonance at δ 13.2 ppm, three aromatic BPh₄ phenyl signals, and THF resonances due to the displacement



Scheme 3 Reactivity of complexes B, C, D, and E toward various H⁺ (acid) sources.

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Fig. 3 Molecular structures of (a) $[(U^{IV}(N(SiMe_3)_2)(THF)_3)_2(\mu-NH)_2][BPh_4]_2$, 4, and (b) $[(U^{IV}(THF)_4)_2(\mu-N)(\mu-NH_2)_2][BPh_4]_3$, 5, with thermal ellipsoids drawn at the 50% probability level. Hydrogen atoms and methyl groups on the ancillary ligands, and BPh₄ counterions have been omitted for clarity.

upon coordination of d_5 -pyridine, but were unable to identify resonances for the imido (NH²⁻) groups (Fig. S18†).

Attempts to isolate 4 analytically pure failed most likely due to the cocrystallization of **B**. Recrystallization of a pyridine solution containing 4, by slow diffusion of Et₂O at -40 °C, resulted in single crystals suitable for XRD analysis, identifiable as the pyridine adduct of 4, [(U^{IV}(N(SiMe₃)₂)(pyridine)₃)₂(µ-NH)₂][BPh₄]₂ (4-pyr; Fig. S24†). All metrical parameters are analogous to the THF adduct of 4. Attempts to isolate larger quantities of 4-pyr proved to be unsuccessful.

¹H NMR studies showed that addition of 2.0 equiv. NH₄BPh₄ to **B** in d_8 -THF after 20 minutes, results in the partial disappearance of the signals of B, and in the appearance of the resonance at δ 12.6 ppm assigned to 4. The ¹H NMR spectrum of the reaction mixture measured after 12 hours showed complete disappearance of the signals corresponding to B and to 4, with concomitant appearance of the resonance assigned to HN(SiMe₃)₂ (Fig. S19a and b†). Furthermore, the reaction mixture, after standing 12 hours at -40 °C, afforded gold crystals of $[(U^{IV}(THF)_4)_2(\mu-N)(\mu-NH_2)_2][BPh_4]_3$ (5) in 83% yield (Fig. 3b and Scheme 3). The product is insoluble in most common solvents including toluene, Et₂O, THF, and n-hexanes, but is soluble in pyridine. The ¹H NMR spectrum of complex 5 in d_5 -pyridine displays three aromatic BPh₄ phenyl signals, THF resonances due to the displacement upon coordination of d_5 pyridine, and two amido (NH₂¹⁻) resonances at -468 ppm. Additionally, complex 5 can also be obtained in 89% yield through direct addition of 3.0 equiv. NH₄BPh₄ to complex E. Therefore, isolating complex **B** is not required for the synthesis of complex 5 (Scheme 3).

The presence of two amido $(\mathrm{NH_2}^-)$ bridging groups in 5, can be interpreted in terms of the binding and concomitant N–H cleavage of two NH $_3$ molecules. This involves protonation of the $\mathrm{U^{IV}}$ –N(SiMe $_3$) $_2$ bonds, resulting in the release of four molecules of HN(SiMe $_3$) $_2$. To date, there is only one other example of the facile N–H activation of NH $_3$ by an uranium complex. The $\mathrm{U^{IV}}$

"tucked-in" mixed-sandwich complex, $[U(\eta\text{-COT}^{\text{TIPS2}})(\eta^5:\kappa^1\text{-}C_5\text{Me}_4\text{CH}_2)]$, was reported to promote the ligand assisted NH₃ activation yielding the terminal U^{IV}–NH₂ complex $[U(\text{COT}^{\text{TIPS2}})\text{Cp*(NH}_2)]$. Uranium complexes of primary amides remain rare, ²¹ and complex 5 provides the second example of a bridging amide. ^{13a}

In order to confirm that the two bridging amido (NH₂¹⁻) moieties in 5, are derived from NH₃, we prepared the ¹⁵N labeled analogue 5-^{14/15}N by reacting the isotopically enriched ¹⁵NH₄BPh₄ with **B**. Adding a solution of HCl in Et₂O to 5-^{14/15}N leads to the formation of both ¹⁵NH₄Cl and ¹⁴NH₄Cl (95% yield), indicated by a doublet and triplet respectively in the ¹H NMR spectrum (2 : 1 ratio, ¹⁵NH₄Cl : ¹⁴NH₄Cl).

These results indicate that formation of the bridging nitride bis-amido complex (5) occurs via protonation of two -N(SiMe₃)₂ ligands, binding of the resulting NH3, and subsequent N-H heterolysis of NH3 assisted by the remaining two basic -N(SiMe₃)₂ ligands. We suggest that the reaction proceeds first through the imido (NH²⁻) intermediate (4), which arises from protonation of one U^{IV}-N(SiMe₃)₂ bond and N-H heterolysis of one NH3 molecule with concomitant elimination of two $HN(SiMe_3)_2$ ligands. This is likely to occur *via* a nitride-amido bridged complex, $[(U^{IV}(N(SiMe_3)_2)(THF))_2(\mu-NH_2)(\mu-N)][BPh_4]_2$ (4int) but the higher nucleophilicity of the nitride, compared to B, results in a proton redistribution, yielding complex 4 (Scheme S2†). Further protonation of a U^{IV}-N(SiMe₃)₂ bond in 4, followed by NH3 binding and N-H heterolysis, should afford a bis-imido, mono-amido complex, [(U^{IV}(THF)₄)₂(μ-NH)₂(μ-NH₂)[BPh₄]₃ (5_{int}), but subsequent rearrangement of the protons instead yields complex 5. Such redistribution of protons can be explained in terms of an increased stability of the nitride compared to the imide when two N(SiMe₃)₂ ligands are replaced by two THF and one amido ligands. Previous studies demonstrated the important effect of ancillary ligands on the stability and reactivity of bridging uranium nitrides.13k

Next, we probed the protonation and reactivity of the neutral nitride complex, C. Complex C is analogous to B in that it contains $-N(SiMe_3)_3$ ancillary ligands, but has two additional ligands within the complex. Previous studies from our group showed that C is less reactive towards small molecule activation in comparison to complexes B and A (Scheme 1). This was interpreted, based on DFT studies, in terms of a decreased nucleophilicity of the nitride moiety, due to an increased bond order of the bridging nitride in the $U^{IV}-N(SiMe_3)_3$ complex C compared to the $U^{IV}-OSi(O^tBu)_3$ complex A. Therefore, we investigated if protonation of the amide ligands and NH_3 activation could also occur in complex C.

Similar to the reactivity of **B** with HNEt₃BPh₄, addition of 4.0 equiv. of HNEt₃BPh₄ to a solution of **C** in d_8 -THF resulted in no reaction (Fig. S20†). Alternatively, treatment with 3.0 equiv. NH₄BPh₄ resulted in an immediate color change from brown to golden yellow. Within 5 minutes, golden crystals of 5 suitable for XRD analysis were obtained in 72% yield. Analysis by ¹H NMR spectroscopy and elemental analysis indicated 5 is the only product isolated, showing similar reactivity to complex **B**. This suggests that the previously determined unreactive nature of **C** can be circumvented by use of a strong acid, promoting a series of protonation/protonolysis reactions with subsequent N-H heterolysis of NH₃ to 5.

Structural characterization of N(SiMe₃)₂-containing complexes

The solid-state molecular structures of complexes 4 and 5 were determined by X-ray diffraction studies. The metrical parameters are presented in Table 1, including those previously reported for complex B.

Complex 4 crystalizes in the space group $P\bar{1}$ with 0.5 molecules per asymmetric unit. Its solid-state structure can be described as an ion pair consisting of two BPh₄⁻ anions and a U^{IV}/U^{IV} mixed dication comprising of two [(U^{IV}(N(SiMe₃)₂)(THF)₃)₂] units bridged by two imido (NH²⁻) ligands. The U1-N1 = 2.198(6) Å and U1-N2 = 2.179(5) Å bond distances are similar and fully consistent with bridging imido (NH²⁻) ligands (Fig. 3a).¹³ⁱ All distances are consistent with delocalized charge on the uranium ions as indicated by the nearly symmetrical U-N_{imido} metrical parameters. The protons of the amido ligands can be crystallographically identified, making their assignment unambiguous.

Complex 5 crystalizes in the space group $P2_1/c$ with one molecule per asymmetric unit. The solid-state molecular structure of 5 shows the presence of an ion pair consisting of three BPh₄⁻ anions and a U^{IV}/U^{IV} trication comprising of two $[(U^{IV}(THF)_4)]$ units bridged by one nitrido (N^3^-) and two amido (NH_2^{1-}) ligands, forming a face-sharing geometry (Fig. 3b). The U1–N1 and U2–N1 (2.062(3); 2.018(3) Å) bond distances are consistent with the presence of a bridging nitride $(\mu$ -N). The U-N–U bond angle changes dramatically from linear in **B** (U–N–U: 168.97(14)°) to highly bent in the diamond core geometry of 5 (107.41(12)°), and is analogous to the previously reported complex **D** (Scheme 3). Bent U1–N2 = 2.455(3) Å and U1–N3 = 2.449(3) Å bond distances are consistent with two bridging amido (NH_2^{1-}) ligands, with nearly symmetrical U–N_{amido} metrical

parameters. The protons of the NH_2 ligands can be crystallographically identified, making their assignment unambiguous. The coordination around the U(n) ions is unprecedented for molecular nitride complexes as it contains only solvent molecules and bridging amido ligands, as all ancillary $-N(SiMe_3)_3$ ligands have been removed upon reaction with NH_4BPh_4 .

Conclusion

In summary, we have investigated the reactions of H⁺ and NH₃ with a series of nitride bridged diuranium(IV) complexes differing in the type and number of supporting ligands.

We found that the nitride ligand is easily protonated by 1.0 equiv. of weak acid (HNEt₃BPh₄) in the –OSi(O^fBu)₃ supported complex **A**, yielding an imido bridged complex **1**. Further protonation of the imido moiety to a terminal NH₃ complex, **2**, can be achieved by using a large excess of weak acid and longer reaction time, or by means of a stronger acid. These results indicate that the uranium-nitride bond is easily protonated to afford NH₃. Ammonia binds the U(IV) ion strongly in the resulting mononuclear U–NH₃ complex **3**, but can be displaced by addition of strong acid. Furthermore, the U–OSi(O^fBu)₃ bonds were found to be stable, even in the presence of stronger acids, such as NH₄BPh₄, therefore indicating that –OSi(O^fBu)₃ supporting ligands are well suited for use when acidic conditions are required, such as in the H[†]/e⁻ mediated catalytic conversion of N₂ to NH₃. ^{5j,23}

Conversely, a very different reactivity is observed for both the cationic and anionic nitride bridged complexes, B and C, supported by -N(SiMe₃)₂ ligands. Opposed to complex A, the nitride in both **B** and **C** are more resistant toward protonation by acids. For example, when a weak acid, such as HNEt₃BPh₄, is employed, the nitride in these complexes is unreactive. This is consistent with the lesser nucleophilic character of the bridging nitride in N(SiMe₃)₂-containing complexes compared to the analogous -OSi(OtBu)₃ systems, which was also supported by their reactivity toward small molecules (CO, CO₂ or H₂). Alternatively, addition of the stronger acid, NH₄BPh₄, resulted in the complete loss of -N(SiMe₃)₂ supporting ligands, while the bridging nitride remains intact. Moreover, the basic -N(SiMe₃)₂ ligands promote the N-H heterolytic cleavage of NH₃, yielding a stable bis-NH₂, mono-nitride bridged complex (5), where only ancillary solvent molecules support the metal center. These results demonstrate that basic supporting ligands, such as -N(SiMe₃)₂, present several disadvantages compared to the -OSi(OtBu)3 ligands for usage in the development of catalysts for N₂ conversion to NH₃. Utilizing OSi(O^tBu)₃-containing complexes is not only advantageous for their resistance toward acids, but also for the high reactivity of bound nitrides to yield NH₃. In contrast, N(SiMe₃)₂-supported complexes may be of interest for studies pertaining to the heterolytic bond activation of NH₃.

Author contributions

M. K. carried out the synthetic experiments and analysed the experimental data. R. S. carried out the X-ray single crystal structure analyses. M. M. originated the central idea,

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coordinated the work, and analysed the experimental data. The manuscript was written through contributions of all authors.

Conflicts of interest

There are no conflicts to declare.

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