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## Introduction

Multimetallic cooperativity plays an important role both in the biological and industrial reduction and cleavage of dinitrogen  $(N_2)$  to ammonia  $(NH_3)$ , but the involved mechanisms remain ambiguous.<sup>1,2</sup> Although the binding and reduction of  $N_2$  by molecular metal complexes have been intensively studied, in most cases only a few compounds have been reported that can cleave  $N_2$  without the assistance of supporting ligands, or the addition of external reducing agents.<sup>2-4</sup>

The first example of N<sub>2</sub> cleavage by a metal complex was reported more than 20 years ago when Cummins and coworkers showed that the Mo( $\pi$ ) complex, [Mo<sup>III</sup>(N(<sup>t</sup>Bu)Ar)<sub>3</sub>] (Ar = 3,5-C<sub>6</sub>H<sub>3</sub>(CH<sub>3</sub>)<sub>2</sub>), cleaves N<sub>2</sub> to yield a Mo(vi) nitride *via* an end-on

# Dinitrogen cleavage by a dinuclear uranium(III) complex<sup>†</sup>

Nadir Jori, ២ <sup>a</sup> Megan Keener, ២ <sup>a</sup> Thayalan Rajeshkumar, <sup>b</sup> Rosario Scopelliti, ២ <sup>c</sup> Laurent Maron 🔎 \*<sup>b</sup> and Marinella Mazzanti ២ \*<sup>a</sup>

Understanding the role of multimetallic cooperativity and of alkali ion-binding in the second coordination sphere is important for the design of complexes that can promote dinitrogen  $(N_2)$  cleavage and functionalization. Herein, we compare the reaction products and mechanism of  $N_2$  reduction of the previously reported K<sub>2</sub>-bound dinuclear uranium(III) complex,  $[K_2\{[U^{III}(OSi(O^tBu)_{3})_3]_2(\mu-O)\}]$ , B, with those of the analogous dinuclear  $[K(2.2.2-cryptand)][K{U^{III}(OSi(O^tBu)_3)_3}_2(\mu-O)],$ complexes, uranium(III) 1, and [K(2.2.2cryptand)]<sub>2</sub>[{U<sup>III</sup>(OSi(O<sup>t</sup>Bu)<sub>3</sub>)<sub>3</sub>}<sub>2</sub>( $\mu$ -O)], 2, where one or two K<sup>+</sup> ions have been removed from the second coordination sphere by addition of 2.2.2-cryptand. In this study, we found that the complete removal of the K<sup>+</sup> ions from the inner coordination sphere leads to an enhanced reducing ability, as confirmed by cyclic voltammetry studies, of the resulting complex 2, and yields two new species upon  $N_2$  addition, namely the  $U({}_{III})/U({}_{IV}) complex, [K(2.2.2-cryptand)][{U^{II}(OSi(O^{t}Bu)_{3})_{3}}(\mu-O){U^{IV}(OSi(O^{t}Bu)_{3})_{3}}], 3, and the N_{2} cleavage (M_{2})_{3}(M_{2})_{3})], 3, and the N_{2} cleavage (M_{2})_{3}(M_{2})_{$ product, the bis-nitride, terminal-oxo complex,  $[K(2.2.2-cryptand)]_2[{U^V(OSi(O^tBu)_3)_3}(\mu-N)_2{U^V(OSi(O^tBu)_3)_2}(\kappa-V)_2(\mu-N)_2$ O)]], 4. We propose that the formation of these two products involves a tetranuclear uranium- $N_2$ intermediate that can only form in the absence of coordinated alkali ions, resulting in a six-electron transfer and cleavage of N<sub>2</sub>, demonstrating the possibility of a three-electron transfer from U(III) to N<sub>2</sub>. These results give an insight into the relationship between alkali ion binding modes, multimetallic cooperativity and reactivity, and demonstrate how these parameters can be tuned to cleave and functionalize N<sub>2</sub>.

bridging intermediate.<sup>5,6</sup> An analogous reactivity was reported recently for a Mo(III) siloxide complex,  $[Mo^{III}(OSi(O^tBu)_3)_3]$ .<sup>7</sup> Remarkably to date, these complexes remain the only examples of N<sub>2</sub> cleavage involving mononuclear metal complexes in the absence of reducing agents. A low-valent heteromultimetallic Nb<sup>III</sup><sub>x</sub>/Na<sub>y</sub> complex was also reported that can cleave N<sub>2</sub>, but the inner sphere cation was proposed to be key in the binding and polarization of N<sub>2</sub>.<sup>89</sup>

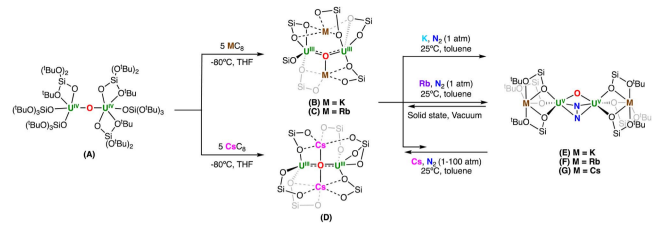
Alkali metals are often employed as external reducing agents in combination with metal complexes, and their key role in the binding, activation, and cleavage of N<sub>2</sub> has been demonstrated by many recent studies.<sup>10–19</sup> In particular, it was proposed that alkali ions are key in promoting the assembly of multimetallic iron complexes that effect the cleavage of N<sub>2</sub>, but the hypothesis was difficult to verify due to the absence of key intermediates.<sup>14</sup> More recently, Holland and coworkers showed that removing the potassium (K<sup>+</sup>) cation from the K-bridged, dinuclear FeNNFe complex, led to a high degree of activation of the N<sub>2</sub> in the resulting mononuclear Fe(N<sub>2</sub>)<sub>2</sub> complex.<sup>20</sup>

Although the number of studies reporting binding and reduction of  $N_2$  by uranium complexes has significantly increased in recent years,<sup>3,18,19,21-35</sup> only a handful of examples of  $N_2$  cleavage to nitrides by uranium have been identified, in which all required the use of an external reducing agent,<sup>18,19,24,33,36</sup> or assistance from a non-innocent ligand,<sup>34</sup>

<sup>&</sup>lt;sup>e</sup>Insititut des Sciences et Ingénierie Chimiques, Ecole Polytechnique Fédérale de Lausanne (EPFL), 1015 Lausanne, Switzerland. E-mail: marinella.mazzanti@epfl.ch <sup>b</sup>Laboratoire de Physique et Chimie des Nano-objets, Institut National des Sciences Appliquées, 31077 Toulouse Cedex 4, France

<sup>&</sup>lt;sup>c</sup>X-Ray Diffraction and Surface Analytics Platform, Institut des Sciences et Ingénierie Chimiques, École Polytechnique Fédérale de Lausanne (EPFL), CH-1015 Lausanne, Switzerland

<sup>&</sup>lt;sup>†</sup> Electronic supplementary information (ESI) available: Experimental details, NMR spectra, XRD, electrochemistry, EPR, IR, UV/vis/near IR, data, computational details. CCDC 277300-2277304. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d3sc05253b



Scheme 1 Previous works on dinitrogen ( $N_2$ ) reduction by multimetallic U(III)-alkali metal ion complexes bearing tris(*tert*-butoxy)siloxide ligands.<sup>19,30</sup>

although very small amounts of a N<sub>2</sub> cleavage side product were identified in one case in the reaction of a putative U(m) with N<sub>2</sub>.<sup>35</sup> We previously reported the reactivity of N<sub>2</sub> with a series of dinuclear U(m) oxo-bridged complexes supported by (tris*-tert*-butoxy)siloxide ligands which contained different alkali metal ions,  $[M_2{[U^{III}(OSi(O'Bu)_3)_3]_2(\mu-O)}]$  (M = K, Rb, Cs) **B–D** (Scheme 1).<sup>19,30</sup> We demonstrated that N<sub>2</sub> binding is less favored with larger cations by steric factors, but that in all the N<sub>2</sub> complexes, a four-electron reduction of N<sub>2</sub> occurs. Full cleavage of N<sub>2</sub> could only be effected by further reduction of the bound N<sub>2</sub><sup>4–</sup> moiety with an external reducing agent, leading to dinuclear and tetranuclear uranium nitride complexes.<sup>18,19</sup>

However, the role of the alkali metal ions in the  $N_2$  cleavage by these multimetallic systems remained ambiguous, and in particular, we were interested in understanding if the presence of the alkali metal ions is indeed essential for  $N_2$  cleavage, or if U(m) ions can cooperatively cleave  $N_2$ .

Herein, we demonstrate that removal of the alkali metal ion from the second coordination sphere has an unexpected outcome on the reactivity of the diuranium(m) complexes with N<sub>2</sub>. Removal of one alkali metal ion forming complex, [K(2.2.2-cryptand)][K {U<sup>III</sup>(OSi(O<sup>f</sup>Bu)<sub>3</sub>)<sub>3</sub>}<sub>2</sub>(µ-O)], **1**, and further reactivity with N<sub>2</sub>, resulted in the formation a highly activated (N<sub>2</sub>)<sup>4-</sup> species. In contrast, both the reaction of the full alkali metal ion-sequestered complex, [K(2.2.2-cryptand)]<sub>2</sub>[{U<sup>III</sup>(OSi(O<sup>f</sup>Bu)<sub>3</sub>)<sub>3</sub>}<sub>2</sub>(µ-O)], **2**, with N<sub>2</sub>, and removal of the K<sup>+</sup> cation from the isolated (N<sub>2</sub>)<sup>4-</sup> complex, **E**, resulted in the immediate cleavage of N<sub>2</sub>. Overall, this study provides the first example of direct stoichiometric N<sub>2</sub> cleavage by an isolated uranium(m) compound without the assistance of the supporting ligand or external alkali metal reducing agents.

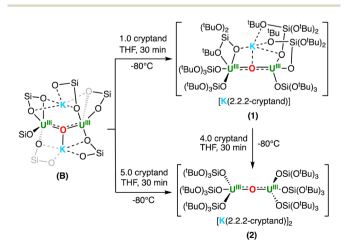
## Results and discussion

#### Synthesis of anionic diuranium(III) complexes

To investigate how the presence of  $K^+$  cations in the second coordination sphere may affect the reaction of dinuclear U(III) complexes with N<sub>2</sub>, we first pursued the synthesis of  $K^+$  cation-sequestered complexes.

The addition of 1 equiv. of 2.2.2-cryptand to a d<sub>8</sub>-THF solution of **B** at -80 °C resulted in the consumption of **B**, with the appearance of a new resonance at -0.27 ppm, and signals corresponding to  $[K(2.2.2\text{-cryptand})]^+$ , as evidenced by <sup>1</sup>H NMR spectroscopy (Fig. S1b†). Single crystals suitable for XRD analysis of  $[K(2.2.2\text{-cryptand})][K{U^{III}(OSi(O'Bu)_3)_3}_2(\mu\text{-}O)]$ , complex 1, were obtained from layering a concentrated THF solution with *n*-hexanes at -40 °C in 72% yield (Scheme 2).

Similarly, addition of 1 equiv. of 2.2.2-cryptand to 1 in  $d_8$ -THF at -80 °C, led to the disappearance of 1, with the appearance of a single new resonance at 1.03 ppm and signals corresponding to  $[K(2.2.2\text{-cryptand})]^+$ , as observed by <sup>1</sup>H NMR spectroscopy (Fig. S1c†). Dark red single crystals suitable for Xrav diffraction studies of  $[K(2.2.2-cryptand)]_{2}[{-}$  $U^{III}(OSi(O^tBu)_3)_3\}_2(\mu-O)]$ , complex 2, were isolated by layering a concentrated THF solution with *n*-hexanes at -40 °C in 76% yield (Scheme 2). It is important to note that the complete removal of the second K<sup>+</sup> cation proved more difficult, and the isolation of analytically pure 2 in high yields required an excess (5 equiv.) of 2.2.2-cryptand.



Scheme 2 Sequestration of the K<sup>+</sup> cations from complex B.

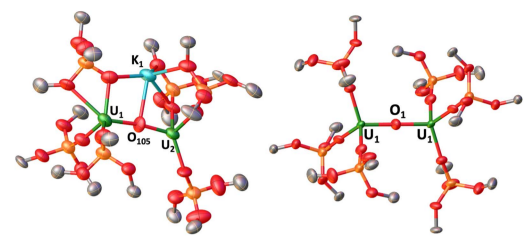


Fig. 1 Solid-state molecular structures of the anion of (left) 1 and (right) 2 with 50% probability ellipsoids. Color code: uranium (green), potassium (light blue), oxygen (red), carbon (grey), silicon (orange). Hydrogen atoms and methyl groups on the  $-OSi(O^tBu)_3$  ligands were omitted for clarity.

Complexes **1** and **2** are insoluble in toluene and showed low thermal stability in THF solutions at 25 °C, resulting in complete decomposition after 6 hours. Removal of one or two  $K^+$  cations in complexes **1** and **2** results in a decreased stability when compared to the previously reported  $K^+$  cation-bound complex, **B**, which is stable in THF at 25 °C for 12 hours.

The solid-state molecular structure of complex 1 (left, Fig. 1) shows an anionic complex with two U(III) ions bridged by an oxo ligand. The inner-sphere  $K^+$  cation (2.851(15) Å) is coordinated to the bridging oxo moiety and by four oxygen atoms of three siloxide ligands, at a shorter distance than that reported for B (2.913(4) Å), suggesting a stronger K1-O<sub>oxo</sub> interaction.<sup>30</sup> The presence of a stronger K-Ooxo bonding is consistent with the fact that an excess of 2.2.2-cryptand is required to fully remove the  $K^+$  cations to yield complex 2. In 2, the second  $K^+$  cation is coordinated by 2.2.2-cryptand as a  $[K(2.2.2-cryptand)]^+$  counterion. The values of the U<sup>III</sup>–O–U<sup>III</sup> bond distances (U1–O105: 2.123(13) and U2–O105: 2.155(13) Å) in 1 are slightly longer than those found in the U<sup>IV</sup>-O-U<sup>IV</sup> complex A (2.085(1), 2.137 (1) Å),<sup>30</sup> and compare well with those found in the  $U^{\rm III}\text{-}O\text{-}U^{\rm III}$ complexes, **B-D** (2.100(5)-2.178(3) Å).<sup>19,30</sup> The U1-O105-U2 core is slightly bent (172.7(8)°), and is consistent with the previously reported K<sup>+</sup> cation-bound U<sup>III</sup>–O–U<sup>III</sup> complex, **B**  $(167.4(2)^{\circ})$ .<sup>30</sup>

The solid-state molecular structure of 2 (right, Fig. 1) shows an anionic complex with two U(III) ions bridged by an oxo ligand, with two outer-sphere  $[K(2.2.2\text{-cryptand})]^+$  counterions. The values of the U<sup>III</sup>-O-U<sup>III</sup> bond distances (U1-O1 and U2-O1:

2.1061(3) Å) are identical, and longer than those found for the  $U^{IV}-O-U^{IV}$  complex **A** (2.085(1), 2.1376(13) Å), but shorter than those observed in 1 (2.123(13) Å, 2.155(13) Å), while comparing well with the previously reported  $U^{III}-O-U^{III}$  complexes, **B**-**D** (2.100(5)–2.178(3) Å). Additionally, the  $U^{III}-O-U^{III}$  bond angle in 2 is linear (180°; Table 1).

#### **Electrochemical studies**

To determine and compare the reducing ability of complexes, 2 and 1, with those of the previously reported complexes **B–D**, cyclic voltammetry studies were carried out under argon with  $[Bu_4N][BPh_4]$  (0.1 M in THF) as the supporting electrolyte (Fig. 2).

As previously observed for complexes **B**–**D** (ref. 19) distinctive irreversible oxidation events at  $E_{pa} = -1.89$  V and -2.25 V are observed in the cyclic voltammograms of **1** and **2**, respectively, which are assigned to the U(m)/U(w) couple (Fig. 2). The corresponding reduction events at  $E_{pc} = -3.23$  V and -3.48 V, respectively, are only observed after initial oxidation, in which these redox events can be attributed to the U(w)/U(m) couple. The reduction potentials for **1** and **2** are more negative compared to the previously reported complex **B** (Table 2), with  $E_{pc}$  values comparable to that observed for **D**. This suggests that partial and full sequestration of the K<sup>+</sup> cations by use of 2.2.2cryptand results in a higher reducing ability for both **1** and **2**, and is comparable to what was previously observed when the K<sup>+</sup> cation is replaced by a weaker Lewis acid, such as Cs<sup>+</sup>.

Table 1 Average	Average values of selected bond lengths (Å) and angles (°) in the complexes					
Complex	U–U	U1-O <sub>oxo</sub>	U2-O <sub>oxo</sub>	U1-O-U2	M1-O <sub>oxo</sub>	M2–O <sub>oxo</sub>
<b>A</b> <sup><i>a</i></sup> <b>B</b> <sup><i>a</i></sup> <b>1</b> <b>2</b> <b>D</b> <sup><i>a</i></sup>	$\begin{array}{c} 4.2128(9) \\ 4.2619(10) \\ 4.2697(12) \\ 4.2123(8) \\ 4.247(1) \end{array}$	2.0852(13) 2.178(3) 2.123(12) 2.1061(3) 2.137(7)	$\begin{array}{c} 2.1376(13)\\ 2.120(3)\\ 2.155(13)\\ 2.1061(3)\\ 2.126(7)\end{array}$	172.19(8) 167(4) 172.7(8) 180 177.9(4)		

<sup>*a*</sup> Values from ref. 19 and 30.

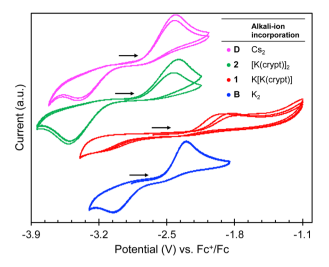


Fig. 2 [-3.9 V; -1.1 V] region of cyclic voltammogram of 3.0 mM THF solutions of complexes D (pink),<sup>19</sup> 2 (green), 1 (red), and B (blue)<sup>19</sup> recorded in 0.1 M [NBu<sub>4</sub>][BPh<sub>4</sub>] under Ar (scan rate = 100 mV s<sup>-1</sup>; referenced against [Fe(C<sub>5</sub>H<sub>5</sub>)<sub>2</sub>]<sup>+/0</sup>).

Table 2 Reduction potentials assigned to the U(iv)/U(iii) couples measured for 1 and 2 compared to values reported for B and D

Complex	$E_{\rm pc}$ (V)	$E_{\mathrm{pa}}\left(\mathrm{V}\right)$	$\Delta E$ (V)
$\mathbf{B}^{a}$	-3.07	-2.30	0.77
1	-3.23	-1.89	1.34
2	-3.48	-2.25	1.23
$\mathbf{D}^{a}$	-3.4	-2.41	0.99
<sup><i>a</i></sup> From ref. 19.	0.1	2.11	0.55

The higher oxidation potential observed for 1 ( $E_{pa} = -1.89 \text{ V}$ ) compared to complexes, 2 ( $E_{pa} = -2.25 \text{ V}$ ) and B ( $E_{pa} = -2.30 \text{ V}$ ), could be attributed to the stronger K–O<sub>oxo</sub> interaction, which renders the removal of the K<sup>+</sup> cation from the inner coordination sphere more difficult, and therefore the overall process more irreversible.

#### Dinitrogen binding and cleavage

Considering the high reducing ability of complexes 1 and 2, we set out to investigate how removal of one or two  $K^+$  cations from the inner coordination sphere of the complex could affect the reactivity with N<sub>2</sub>, when compared to the previously reported **B**–**D** complexes.

First, exposing a dark red solution of 2 in THF to N<sub>2</sub> at -40 °C, resulted in an immediate color change to dark orange. Analysis by <sup>1</sup>H NMR spectroscopy of the reaction mixture at -40 °C revealed the consumption of 2, and the formation of a <sup>1</sup>H NMR silent species, with the appearance of the signal corresponding to [K(2.2.2-cryptand)][OSi(O<sup>t</sup>Bu)<sub>3</sub>] (formation of 0.5 equiv. determined by quantitative <sup>1</sup>H NMR spectroscopy) (Fig. S11†). Instead, analysis of the same reaction mixture at -80 °C by <sup>1</sup>H NMR spectroscopy revealed refined resonances at  $\delta = 35.48$ , 6.61, and -14.81 ppm (Fig. S12†), corresponding to the U(m)/U(rv) anion, [{U<sup>III</sup>(OSi(O<sup>t</sup>Bu)<sub>3</sub>)<sub>3</sub>]( $\mu$ -O}{U<sup>IV</sup>(OSi(O<sup>t</sup>Bu)<sub>3</sub>)<sub>3</sub>]<sup>-</sup>. The assignment of the

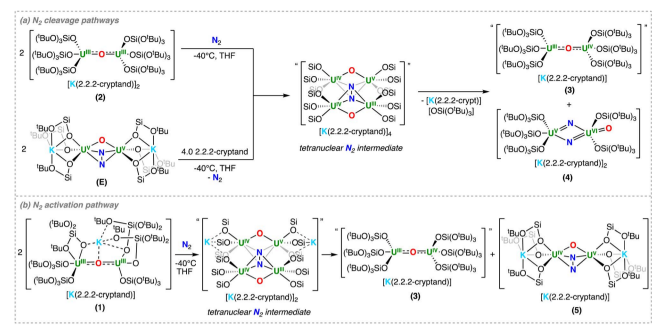
putative, K-sequestered U(m)/U(n) species, [K(2.2.2-cryptand)] [{U<sup>III</sup>(OSi(O'Bu)<sub>3</sub>)<sub>3</sub>}( $\mu$ -O){U<sup>IV</sup>(OSi(O'Bu)<sub>3</sub>)<sub>3</sub>}], **3**, was confirmed by the independent synthesis of the U(m)/U(n) complex, [K {U<sup>III</sup>(OSi(O'Bu)<sub>3</sub>)<sub>3</sub>}( $\mu$ -O){U<sup>IV</sup>(OSi(O'Bu)<sub>3</sub>)<sub>3</sub>], **3-K**, and further addition of 2.2.2-cryptand (Scheme S1†). Notably, the <sup>1</sup>H NMR spectrum of complex **3-K** in the presence of 1 equiv. 2.2.2-cryptand in d<sub>8</sub>-THF at -80 °C, displayed identical resonances (Fig. S10†) as observed in the reaction mixture obtained after addition of N<sub>2</sub> to **2**.

From the reaction mixture obtained after exposing 2 to N<sub>2</sub>, golden crystals suitable for XRD analysis of the bridging bisnitride, terminal-oxo complex,  $[K(2.2.2\text{-cryptand})]_2[{-U^V(OSi(O^tBu)_3)_3}(\mu-N)_2{U^{VI}(OSi(O^tBu)_3)_2(\kappa-O)}]$ , 4, were isolated from a mixture of *n*-hexanes and THF (10:1) at -40 °C in 67% yield (yield is provided considering the reaction stoichiometry in Scheme 3). The <sup>1</sup>H NMR spectrum of 4 in d<sub>8</sub>-THF is silent at -40 °C and -80 °C, and the complex is insoluble in non-polar solvents.

X-Band EPR studies of the *in situ* reaction mixture obtained after addition of N<sub>2</sub> to 2 (20 mM frozen THF/Et<sub>2</sub>O solution), showed two independent sets of signals at 6 K, in which the first is very intense and was fit to an axial set of *g*-values ( $g_1 = 2.35$ ;  $g_2 = 2.05$ ;  $g_3 = 2.05$ ), whereas the second signal is at higher field (g = 1.13 and g = 1.09), and much less intense (Fig. S49†). The first signal was attributed to the U(m)/U(nv) complex, **3**, whereas the second signal could be assigned to the U(v)/U(v1) bis-nitride, terminal-oxo complex, **4**. These assignments were confirmed by independent measurement of the EPR spectra for the putative U(m)/U(nv) complex, **3**, and of isolated **4**, both in frozen THF : Et<sub>2</sub>O solutions (Fig. S52–S58†) and are consistent with *g*values reported for U(m) complexes<sup>37,38</sup> and U(v) terminal oxo complexes.<sup>39–41</sup>

The solid-state molecular structure of 4 (Fig. 3) shows the presence of a dinuclear complex, in which there are two molecules per asymmetric unit, where the two uranium ions are bridged by two nitrides, and are overall bound by five –  $OSi(O^tBu)_3$  ligands, indicating the loss of one ligand.

The overall charge of complex 4 is consistent with the presence of  $U(v)/U(v_1)$  centers. The two U ions are pentacoordinated and display a distorted square pyramidal geometry, and are bridged by two nitride ligands with a short U…U distance of 3.3672(5) Å. The U1 ion is coordinated by three siloxide ligands and the two nitrides, while U2 is coordinated by two siloxide ligands, two nitrides, and a terminal oxo moiety. The U2N2O core is planar, with a N1-N2 separation of 2.543(11) Å, ruling out the presence of a bond between the two nitrogen atoms. The bridging U-N bond distances are asymmetric, featuring a combination of short (U1-N2: 1.950(7) Å, U2-N1: 1.892(8) Å) and elongated (U1-N1: 2.315(8) Å, U2-N2: 2.251(7) Å) bond distances. This is consistent with the presence of U=N multiple bonds and singly bound U-N, respectively, as observed in previously reported  $U(v_I)/U(v_I)$  and  $U(v_I)/U(v)$  bis-nitride bridged complexes.35,42 The bond valence sum analysis and the computational data (vide infra), suggest that U1 and U2 are formally +5 and +6, respectively; however, a delocalized valence cannot be fully ruled out. Overall, the solid-state molecular structure of 4 displays a unique nitride-



Scheme 3 (a) Dinitrogen cleavage by 2 and upon addition of 2.2.2-cryptand to E. (b) Dinitrogen activation by 1.

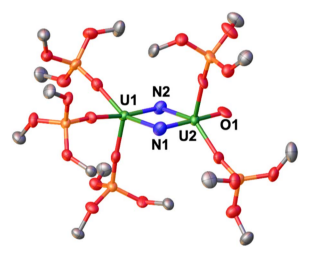


Fig. 3 Solid-state molecular structure of the anionic moiety  $[{U^V(OSi(O^tBu)_3)_3}(\mu-N)_2{U^{VI}(OSi(O^tBu)_3)_2(\kappa-O)}]^{2-}, 4, (50\% probability ellipsoids). Color code: uranium (green), oxygen (red), carbon (grey), nitrogen (blue), and silicon (orange). Hydrogen atoms, solvent molecules and methyl groups the <math>-OSi(O^tBu)_3$  were omitted for clarity.

substituted analogue of the uranyl(VI) ion, which is only the third example of such O=U=N motif. The distances observed in the trans oxo-nitrido moiety [O=U=N] found in 4 (U2-O1: 1.845(6) Å; U2-N1 = 1.892(8) Å) are significantly longer than those found in the trans sodium-capped oxonitrido U(vI) complex reported by Hayton and coworkers (U-O<sub>oxo</sub> = 1.797(7) Å; U-N<sub>nitride</sub> = 1.818(9) Å),<sup>43</sup> and are comparable to those found in the analogous cesium-capped [O=U=N] motif in the previously reported complex, [Cs<sub>3</sub>{-U(OSi(O<sup>t</sup>Bu)<sub>3</sub>)<sub>3</sub>)( $\mu$ -N)<sub>2</sub>{U(OSi(O<sup>t</sup>Bu)<sub>3</sub>)<sub>2</sub>( $\kappa$ -O)}][CsOSi(O<sup>t</sup>Bu)<sub>3</sub>], (U-O<sub>oxo</sub>: 1.856(4) Å; U2-N2<sub>nitride</sub> = 1.85(1) Å).<sup>19</sup>

The molecular structure of **4** is similar to that of the bisnitride, terminal-oxo U(v)/U(v) complex,  $[Cs_3{U(OSi(O^tBu)_3)_3}(-\mu-N)_2{U(OSi(O^tBu)_3)_2(\kappa-O)}][CsOSi(O^tBu)_3]$ . However, it should be noted that in order to promote full cleavage of N<sub>2</sub> from **D**, the addition of 2 equiv. of an alkali metal reducing agent (CsC<sub>8</sub>) under N<sub>2</sub> atmosphere were required, despite the similar redox potentials measured for **2** and **D**. In contrast, the diuranium(m) complex, **2**, is able to effect the direct, stoichiometric cleavage of N<sub>2</sub> without further addition of reducing agent or assistance from supporting ligand, which is unprecedented in f elements chemistry. Additionally, it is remarkable that one uranium ion is able to transfer three electrons to N<sub>2</sub>, yielding a U(v1) ion, as these multielectron transformations still remain rare for uranium.

The formation of the  $U(v_1)/U(v)$  bridging bis-nitride, terminal-oxo complex, **4**, requires the binding and reduction of N<sub>2</sub> by two molecules of complex **2**, most likely through a proposed tetranuclear intermediate, which is supported by the isolation of **4** and the formation of **3** as observed by <sup>1</sup>H NMR spectroscopy (Scheme 3a).

In our previous works, the reaction of the alkali metal-bound diuranium(m) complexes, **B-D** were shown to form the diuranium(v)–N<sub>2</sub><sup>4–</sup> **E-G** complexes (Scheme 1), in which each had different binding constants; however, no direct N<sub>2</sub> cleavage was observed. Considering that the measured reduction potentials for 1 and 2 by cyclic voltammetry studies were greater than for **B**, but very similar to that of **D**, the observed differences in reactivity can be interpreted in terms of unfavorable steric interactions for the alkali ion-bound complexes, which most likely prevent the formation of the tetranuclear intermediate. To further verify this hypothesis, we next studied how removal of the K<sup>+</sup> cation from complex **E** could affect the bound hydrazido (N<sub>2</sub><sup>4–</sup>) moiety.

The addition of 2 equiv. of 2.2.2-cryptand to complex **E** in d<sub>8</sub>-THF at -40 °C, resulted in an immediate color change from dark brown to dark orange. Analysis by <sup>1</sup>H NMR spectroscopy indicated the formation of [K(2.2.2-cryptand)][OSi(O<sup>t</sup>Bu)<sub>3</sub>] and 3 (Fig. S15†), similar to the reaction mixture of 2 and N<sub>2</sub>. Golden crystals of 4 were isolated in 50% yield (per 1.0 equiv. of **E**) by leaving a concentrated hexane : toluene (10 : 1) solution at -40 ° C over the course of two days.

Overall, these results suggest that removal of the coordinated K<sup>+</sup> cations from complex **E**, results in an important structural change which promotes further reactivity of the bound N<sub>2</sub>. The cleavage of N<sub>2</sub>, and the formation of the putative U<sup>III</sup>–O–U<sup>IV</sup> complex **3**, requires two dimers to fully reduce one molecule of N<sub>2</sub>. We propose that due to steric factors, removal of the K<sup>+</sup> cations results in a weaker binding of N<sub>2</sub>, where an equilibrium between the U(III)/U(III) complex, **2**, and the U(v)/U(v)–N<sub>2</sub><sup>4–</sup> complex, exists (*vide infra*), as previously observed for **D**. However, the removal of the bound alkali metal cations allows two dimeric complexes to interact and form the proposed tetranuclear intermediate, which can then effect the six-electron transfer and subsequent cleavage of one molecule of N<sub>2</sub>, yielding the U(vI)/U(v) bis-nitride, terminal-oxo complex **4**.

We next investigated the reaction of 1 with  $N_2$  to assess if the tetranuclear intermediate may be accessible when one potassium is still bound in the second coordination sphere.

Exposing a dark red solution of **1** in THF at -40 °C to N<sub>2</sub>, resulted in an immediate color change to dark brown-orange. Analysis of the reaction mixture by <sup>1</sup>H NMR spectroscopy in d<sub>8</sub>-THF at -80 °C, showed the complete consumption of **1** and the formation of the putative U(m)/U(n) complex, **3**, suggesting a similar reaction pathway (Fig. S20†). However, the formation of [K(2.2.2-cryptand)][OSi(O<sup>t</sup>Bu)<sub>3</sub>], which had been observed during N<sub>2</sub> cleavage to form complex **4**, is not observed in this reaction mixture (Scheme 3a and b).

Indeed, the X-band EPR spectrum of the reaction mixture for 1 and N<sub>2</sub> in a THF : Et<sub>2</sub>O (1 : 1) frozen glass solution at 6 K shows the presence of the U(m)/U(w) complex, **3**. However, there were no additional signals suggestive of a U(v) species, indicating that **4** is most likely not formed in this reaction (Fig. S59†).

Attempts to isolate the N-containing species from this reaction mixture proved unsuccessful. However, through an alternative route, single crystals of a U(v)/U(v) hydrazido (N<sub>2</sub><sup>4</sup>) complex, [K(2.2.2-cryptand)][K<sub>2</sub>{U<sup>V</sup>(OSi(O<sup>t</sup>Bu)<sub>3</sub>)<sub>3</sub>} {U<sup>IV</sup>(OSi(O<sup>t</sup>Bu)<sub>3</sub>)<sub>3</sub>}(\mu-O)(\mu-N<sub>2</sub>)], 5 (Fig. S39†), could be isolated upon addition of 1 equiv. of 2.2.2-cryptand to E in toluene at -40 °C. It is important to note that complex 5 could only be obtained once, as attempts to isolate analytically pure material were unsuccessful due to the low solubility of the reactants and products in toluene. Isolation of the U(v)/U(v) N<sub>2</sub><sup>4-</sup> complex, 5, provides further support for the proposed tetranuclear intermediate (Scheme 3b).

Overall, the reaction of **1** with  $N_2$  also involves four uranium centers; however,  $N_2$  cleavage is not observed due the presence of  $K^+$  cations in the inner coordination sphere, decreasing the reducing ability of the U ions, preventing the transfer of one additional electron to reduce  $N_2$ . In contrast, the reaction of

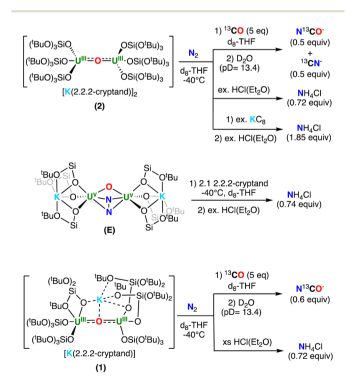
complexes **B–D** with  $N_2$  only involves two uranium centers due to steric hindrance.

#### Reactivity of the N-containing complexes

To further investigate the N-containing species formed upon addition of  $N_2$  to complexes 2 and 1, as well as the addition of 2.2.2-cryptand to E, we next pursued their reactivity with acids  $(H^+)$  and CO.

The addition of excess HCl(Et<sub>2</sub>O) to the residue of the reaction mixture obtained from 2 and N<sub>2</sub>, yielded NH<sub>4</sub>Cl in 72% yield (per 1 equiv. of 2). To further determine whether the NH<sub>4</sub>Cl was unambiguously derived from N2, the reactivity with labelled <sup>15</sup>N<sub>2</sub> was investigated, leading to the formation of isotopically enriched <sup>15</sup>NH<sub>4</sub>Cl as evidenced by <sup>1</sup>H NMR spectroscopy (Fig. S24<sup>†</sup>). Similarly, addition of excess HCl(Et<sub>2</sub>O) to the residue of the reaction mixture obtained from E and 2.1 equiv. of 2.2.2-cryptand in THF, also yielded NH4Cl in 74% yield (per 1 equiv. of E). Overall, these results are consistent with the formation of the bis-nitride, terminal-oxo complex, 4, as the primary N-containing species (Scheme 4). Finally, the addition of excess HCl(Et<sub>2</sub>O) to isolated 4 resulted in the formation of NH<sub>4</sub>Cl in 92% yield (1.84 equiv., 100% conversion corresponding to 2 equiv. of NH<sub>4</sub>Cl), consistent with the presence of the two nitride ligands (Scheme 4).

We also investigated the reactivity with  ${}^{13}$ CO. The addition of 5 equiv. of  ${}^{13}$ CO to the reaction mixture obtained after addition of N<sub>2</sub> to 2, led to an immediate color change from dark orange to brown. The products of the reaction could not be isolated; however, quenching the reaction mixture with basic (pD = 13.4) D<sub>2</sub>O, revealed the presence of 0.5 equiv. of N<sup>13</sup>CO<sup>-</sup> and 0.5



Scheme 4 Reactivity of (top) 2, (middle) E and 2.2.2-cryptand, and (bottom) 1, after addition of N<sub>2</sub> with H<sup>+</sup> and  $^{13}$ CO.

equiv. of  ${}^{13}CN^-$  (per 1 equiv. of 2), as evidenced by quantitative  ${}^{13}C$  NMR spectroscopy (Fig. S29†). The amount of NCO<sup>-</sup>/CN<sup>-</sup> is consistent with the formation of 0.5 equiv. of the bis-nitride, terminal-oxo complex, 4 (top, Scheme 4). The formation of NCO<sup>-</sup>/CN<sup>-</sup> has been previously observed in the reactivity of uranium nitride complexes with CO.<sup>18,19,42,44-46</sup> In particular, the previously reported dinuclear U(v) bis-nitride complex, [K<sub>2</sub>-{U<sup>V</sup>(OSi(O<sup>t</sup>Bu)<sub>3</sub>)<sub>3</sub>}<sub>2</sub>( $\mu$ -N)<sub>2</sub>], showed similar reactivity with CO yielding a 1:1 ratio of N<sup>13</sup>CO<sup>-</sup>: <sup>13</sup>CN<sup>-</sup>.

Alternatively, the addition of 5 equiv. of <sup>13</sup>CO to the reaction mixture obtained after addition of N<sub>2</sub> to 1, led to a series of color changes over 48 hours. The products of the reaction could not be isolated, but quenching the reaction mixture with basic (pD = 13.4)  $D_2O_2$ , revealed the presence of 0.5 equiv. of  $N^{13}CO^{-}$  (per 1 equiv. of 1) by quantitative <sup>13</sup>C NMR spectroscopy (Fig. S32<sup>†</sup> and Scheme 4). This reactivity is most consistent with the formation of 0.5 equiv. of the  $N_2^{4-}$  complex, 5, as the formation of NCO<sup>-</sup> with concomitant release of N<sub>2</sub> was previously observed for the reaction of CO and the U(v)/U(v)hydrazido-amide complex,  $[K_2{U^V(OSi(O^tBu)_3)_3}_2(\mu-NH)(\mu-$ N2)].29 Interestingly, addition of excess HCl(Et2O) to the residue obtained from the reaction mixture of 1 with N<sub>2</sub> yielded NH<sub>4</sub>Cl in 74% yield (per 1 equiv. of 1). Notably, the formation of NH<sub>4</sub>Cl was not previously observed upon addition of HCl to the U(v)/U(v)  $N_2^{4-}$  complexes, E-G,<sup>19,30</sup> suggesting that the  $N_2^{4-}$  in the U(rv)/U(v) complex, 5, is more activated.

#### **Computational studies**

To gain insight into the  $N_2$  reduction pathways promoted by complexes, **1** and **2**, DFT calculations (B3PW91) were performed, including solvent and dispersion forces. Formation of the isolated  $U(v)/U(v_1)$  bis-nitride, terminal-oxo complex, **4**,

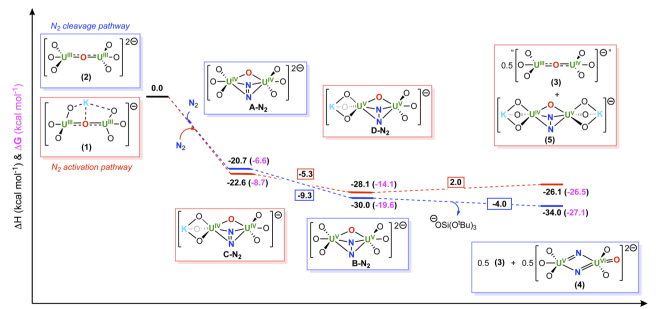
from the reaction of N<sub>2</sub> and **2**, was shown to be thermodynamically favorable by  $-34.0 \text{ kcal mol}^{-1} (-26.5 \text{ kcal mol}^{-1} \text{ in}$ Gibbs free energy) (Scheme 5). Unfortunately, the calculations of the proposed tetranuclear intermediate were intractable; however, the calculations demonstrate that complete reduction of N<sub>2</sub> may involve two intermediate species, namely a U(rv)/U(rv) N<sub>2</sub><sup>2-</sup> (**A-N**<sub>2</sub>) and a U(v)/U(v) N<sub>2</sub><sup>4-</sup> complex (**B-N**<sub>2</sub>), in which both intermediate species involve a step-wise, twoelectron reduction to N<sub>2</sub>.

Interestingly, the first two-electron reduction of  $N_2$ , forming intermediate **A-N**<sub>2</sub>, is computed to be exothermic by  $-20.7 \text{ kcal mol}^{-1}$ , (-6.6 kcal mol $^{-1}$  in Gibbs free energy), as well as the second two-electron reduction to yield the  $N_2^{4-}$  intermediate, **B-N**<sub>2</sub>, is exothermic by -9.3 kcal mol $^{-1}$ (-7.5 kcal mol $^{-1}$  in Gibbs Free energy). The final N–N bond cleavage step, which involves the proposed tetranuclear uranium intermediate, is also exothermic by -4.0 kcal mol $^{-1}$ (-12.4 kcal mol $^{-1}$  in Gibbs free energy).

Next, the oxidation states of the complexes, 2 and 4, as well as the intermediates,  $A-N_2$  and  $B-N_2$ , were verified by computing the different spin states.

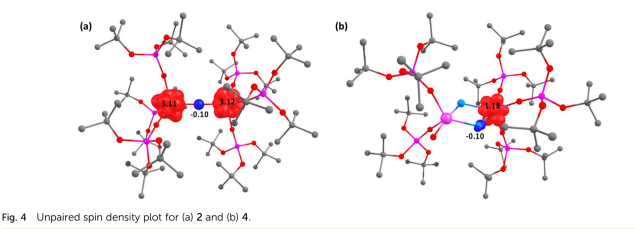
For complex 2, the septet (s = 3), quintet (s = 2), and triplet (s = 1) spin states were considered. As expected for a U(m)/U(m) system, the septet (s = 3; six unpaired electrons) was found to be the ground state, with the other spin states higher than 14.7 kcal mol<sup>-1</sup> in energy. The unpaired spin localization clearly shows that the unpaired electrons are fully localized at the uranium centers (Fig. 4).

The spin states for the N<sub>2</sub>-bound intermediate species, namely, **A-N<sub>2</sub>** and **B-N<sub>2</sub>**, were also computed considering the quintet (s = 2), triplet (s = 1), and singlet (s = 1) ground states. Interestingly, the quintet (s = 2) is 3.1 kcal mol<sup>-1</sup> higher in energy than the triplet (s = 1), while the singlet (s = 0) is higher



**Reaction Coordinate** 

Scheme 5 Computed enthalpy and Gibbs free energy ( $\Delta H$  (black) and  $\Delta G$  (pink) in kcal mol<sup>-1</sup>) profiles for the formation of (blue) 4 and (red) 5.



than 44.0 kcal mol<sup>-1</sup> in energy. Scrutinizing the unpaired spin density on the U centers for the two lower spin states (quintet and triplet), clearly shows that the quintet (s = 2) corresponds to the U(rv)/U(rv) N<sub>2</sub><sup>2-</sup> (**A-N**<sub>2</sub>), whereas the triplet (s = 1) corresponds to the U(v)/U(v) N<sub>2</sub><sup>4-</sup> (**B-N**<sub>2</sub>) intermediate. Interestingly, in the quintet (s = 2) spin-state, the SOMO-2 and the SOMO-3 (SOMO = singly occupied molecular orbital), indicates occupation of two degenerate N–N  $\pi^*$  ligand orbitals, suggesting that the N<sub>2</sub> has undergone a two-electron reduction to a N<sub>2</sub><sup>2-</sup> moiety, yielding formal U(rv)/U(rv) 5f<sup>2</sup> centers. Therefore, some unpaired spin density is observed on the N<sub>2</sub> moiety with a ferromagnetic coupling with the unpaired spins at the uranium.

For the bis-nitride, terminal-oxo complex, 4, a doublet (s = 1/2) spin state was calculated, which is consistent with a mixedvalent 5f<sup>1</sup> U(v)/U(v1) complex. The unpaired spin density is located on only one uranium, suggesting that this is most likely the U(v) 5f<sup>1</sup> ion, whereas the second uranium ion does not display any unpaired spin density, suggesting this is a U(v1) 5f<sup>0</sup> ion (Fig. 4), which is in line with the observed bond valence sum analysis. Interestingly, the two bridging nitride moieties are negatively charged (-0.8, -0.9), indicating a nucleophilic character.

Similar analysis was performed for the reduction of N<sub>2</sub> by complex 1, in which calculations for the proposed tetranuclear intermediate were also intractable. In this system, the two stepwise, two-electron reductions of N<sub>2</sub>, forming the intermediate,  $U(v)/U(v) N_2^{2-}$  (C-N<sub>2</sub>) and  $U(v)/U(v) N_2^{4-}$  (D-N<sub>2</sub>) species, were found to be thermodynamically favorable by -22.6 and -5.3 kcal mol<sup>-1</sup> (-8.7 and -10.9 kcal mol<sup>-1</sup> in Gibbs free energy, respectively) respectively. Whereas, subsequent formation of the bis-nitride, terminal-oxo complex, 4, was endothermically unfavorable by 34.6 kcal  $mol^{-1}$ . Therefore, the overall six-electron reduction and cleavage of N2 is endothermic by 6.7 kcal mol<sup>-1</sup>. In contrast, the formation of the  $U(rv)/U(v) N_2^{4-}$  complex, 5, is slightly endothermic by 2.0 kcal  $mol^{-1}$  (exergonic by 7.5 kcal  $mol^{-1}$ ) leading to an overall thermodynamically favorable reaction pathway  $(-26.1 \text{ kcal mol}^{-1} \text{ in enthalpy}, -27.1 \text{ kcal mol}^{-1} \text{ in Gibbs free})$ energy). Overall, these results are consistent with the experimental findings, suggesting that generation of the U(IV)/U(V)  ${N_2}^{4-}$  species, 5, is more favorable when partial  $K^+$  cation sequestration has occurred, whereas  $N_2$  cleavage to the U(v)/U(v1) bis-nitride, terminal-oxo complex, 4, is more favorable when all  $K^+$  cations have been removed from the inner coordination sphere.

## Conclusions

Herein, we have compared the mechanism and resulting products of N<sub>2</sub> reduction by previously reported K<sub>2</sub>-bound dinuclear uranium(III) complexes with those of the analogous uranium(III) dinuclear complexes, 1 and 2, where one or two K<sup>+</sup> ions have been removed from the inner coordination sphere by addition of 2.2.2-cryptand. The complete sequestration of the K<sup>+</sup> cations resulted in an enhanced reducing ability of complex 2, leading to the formation of two products upon N<sub>2</sub> addition, namely, the U(III)/U(IV) complex, 3, and the  $U(v)/U(v_1)$  bis-nitride complex, 4. The formation of these two products requires four uranium centers to be involved in N2 cleavage, and is proposed to occur via a tetranuclear  $N_2$ intermediate that can only form in the absence of coordinated alkali metal ions, resulting in a six-electron N<sub>2</sub> cleavage. Removal of only one K<sup>+</sup> cation and subsequent reactivity with  $N_2$  led to the formation of the U(III)/U(IV) complex, 3, and to the formation of a U(v)/U(rv)  $N_2^{4-}$  complex, 5, where the  $N_2^{4-}$ is more activated than in the analogous  $U(v)/U(v) N_2^{4-}$ , E-G complexes, as indicated by its high reactivity with electrophiles, such as acid (H<sup>+</sup>) and CO. Most notably, we had previously shown that N2 evolution occurred upon addition of a strong acid (HCl) to the complexes E and F, whereas in this study, we found HCl addition to complex 5 resulted in NH<sub>4</sub>Cl formation. Additionally, computational studies indicate that  $N_2$  cleavage by 2, with concomitant formation of the bisnitride complex, 4, is thermodynamically favored. Both the U(v)/U(v) bis-nitride 4, and the  $U(v)/U(v) N_2^{4-}$ , 5, complexes react with CO in ambient conditions leading to CN<sup>-</sup> and NCO<sup>-</sup> or solely NCO<sup>-</sup>, respectively. Overall, these results provide an important insight into the relationship between alkali ion-binding modes and the multimetallic cooperativity and reactivity within a unique uranium system that cleaves and functionalizes N2, demonstrating the possibility of a three-electron transfer from U(III) to  $N_2$ .

## Data availability

The data that support the findings of this study are openly available in the Zenodo repository at https://doi.org/10.5281/zenodo.10044157.

## Author contributions

N. J. designed and carried most experiments and analyzed the data; M. K. identified the conditions for the isolation of the key complex 4. M. M. designed and supervised the project; T. R. and L. M. carried out the computational study; R. S. measured and analyzed the X-Ray data; N. J., M. K., and M. M. wrote the manuscript with contributions of all authors, and all authors have given approval for the final version of the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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