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Alkyl and Hydride Bis(trimethylsilyl)Amido Derivatives of the Actinide Elements; Preparation

and Hydrogen-Deuterium Exchange

Stephen J. Simpson, Howard W. Turner, and Richard A. Andersen*

Contribution from Chemistry Department and Materials and Molecular Research Division of Lawrence Berkeley Laboratory, University of California, Berkeley, California 94720

Abstract

The monomeric, hydrocarbon-soluble monohydrides and monodeuterides of the actinide metals (thorium or uranium) of the type HM[N(SiMe₃)₂]₃ have been prepared. Their reaction chemistry, n-BuLi followed by MeBr yields MeM[N(SiMe₃)₂]₃ and borane in tetrahydrofuran yields BH₄M[N(SiMe₃)₂]₃, suggests that the hydrogen atom is hydridic. Pyrolysis of the hydrides yields the novel, four-membered ring metallocycle, $[(Me_3Si)_2N]_2 - MCH_2Si(Me)_2NSiMe_3$ where M is Th or U. These metallocycles are the key intermediates in the hydrogen-deuterium exchange reaction that yields $\{[(CD_3)_3Si]_2N\}_3MD.$

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Metal alkyl and hydride derivatives are important compounds in inorganic and organometallic chemistry.^{1a} These functional groups are of interest in their own right as well as for their utility in organic synthesis.^{1b,C} Though the organometallic and hydride derivatives of the p- and d-block elements are extensive, those of the f-block elements have been rather less extensively explored. The latter compounds are generally of the type Cp_3MR where M is an actinide element or Cp_2MR where M is a lanthanide element.² In contrast to the rather extensive alkyl chemistry, only two hydride derivatives, $(Me_5C_5)_4M_2H_4$ (M = Th or U) are known^{3a,b} though attempts at their preparation have been described.^{3c,d}

The large size and "hard" acid character of the actinide elements affords, as a natural consequence, compounds with high coordination numbers with nitrogen or oxygen ligands. In order to prepare low-coordination number, volatile, and hydrocarbon soluble compounds of these metals, sterically bulky ligands are The bis(trimethylsilyl)amido ligand, (Me₃Si)₂N, is required. ideally suited for this task.⁴ The steric bulk of this amide is shown, by way of example, by the observation that the uranium (III) species, $[(Me_3Si)_2N]_3U$, is monomeric and three coordinate.⁵ Further, the tetravalent methyl and tetrahydroborate derivatives, $RM[N(SiMe_3)_2]_3$ where R is Me or BH_A and M is Th or U, are also monomeric.⁶ Since the simple methyls and tetrahydroborates can be prepared readily there is no reason why the simple hydride species should not exist.⁷

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Refluxing a tetrahydrofuran solution of four molar equivalents of sodium bis(trimethylsilyl)amide and uranium or thorium tetrachloride yields the hydrides, $HM[N(SiMe_3)_2]_3$, M = Th or U. The hydrides may also be prepared from $ClM[N(SiMe_3)_2]_3$ and one molar equivalent of $NaN(SiMe_3)_2$ in refluxing tetrahydrofuran. Tetrahydrofuran is implicated as the hydrogen atom source since if the latter reaction is conducted in refluxing diethyl ether, benzene, or isooctane, the starting chloro-species, $ClM[N(SiMe_3)_2]_3$ is recovered quantitatively. This inference is substantiated by using perdeuterotetrahydrofuran as reaction solvent. In this solvent the deuteride, $DM[N(SiMe_3)_2]_3 M = Th$ or U, is isolated. The observation that tetrahydrofuran is the source of the hydrogen atom is not unique, as it has been observed previously.⁸

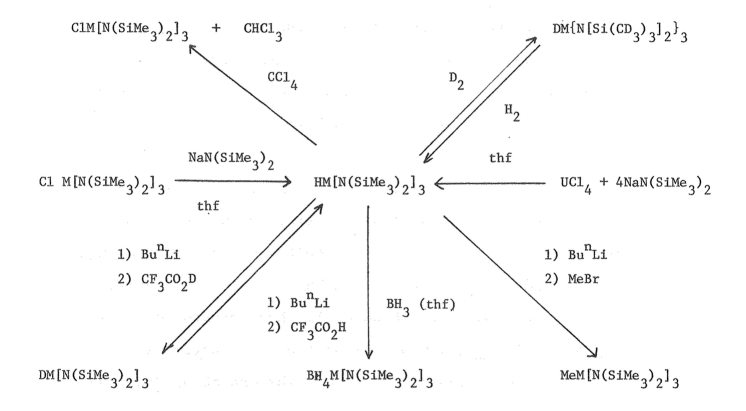
The thorium hydride is colorless and diamagnetic. It is soluble in pentane from which it may be crystallized. The uranium hydride is brown-yellow, paramagnetic ($\mu_B = 2.62$ B.M.), and soluble in pentane. Both hydrides may be sublimed in vacuum at ca. 80-100°C, and afford monomeric molecular ions (M-2) in the mass spectrometer. They are also monomeric in the solid state.⁹ The thorium hydride has a Th-H stretching frequency in the infrared spectrum at 1480 cm⁻¹ and the deuteride absorbs at 1060 cm⁻¹. The uranium-hydrogen stretching frequency occurs at 1430 cm^{-1} and that of the deuteride occurs at 1020 cm^{-1} . The low terminal metal-hydrogen stretching frequency is rather surprising.¹⁰ However, the metal-hydrogen stretching frequency in $(Me_5C_5)_2TiH_2$ and $(Me_5C_5)_2ZrH_2$ appear at 1560 and 1555 cm⁻¹ respectively.^{11,12} Further, the zinc-hydride stretching frequency

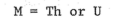
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in the dimeric amidozinc hydride, $(Me_2NCH_2CH_2NMeZnH)_2$, with terminal zinc hydrogen bonds, appears at 1695 cm⁻¹.¹³ In the only other molecular hydrides that have been described for the actinide elements, $(MeC_5)_4M_2H_4$, where M is thorium or uranium, the absorptions associated with the thorium-hydrogen bond occur at 1406, 1361, 1215, and 1114 cm⁻¹.^{3a}

The ¹H nuclear magnetic resonance spectrum of the diamagnetic thorium hydride shows two absorptions at δ 0.63 and δ 0.37, the former being due to the unique hydride and the latter due to the $(Me_3Si)_2N$ ligand. The δ 0.63 absorption is absent in the deuteride. The ¹³C NMR spectrum is a quartet centered at δ 6.80 (¹J_{CH} = 118 Hz). We have been unable to find the hydride signal for the paramagnetic (f²) uranium species though the $(Me_3Si)_2N$ protons and carbons resonate at δ -19.5 and δ -40.8, respectively. By way of comparison the ¹³C NMR spectrum of the uranium (III) species, U[N(SiMe_3)_2]_3 yields a guartet centered at δ -71.9 (J_{CH} = 118 Hz, 28°C, v_{l_2} = 3 Hz). The chemical shift of the Th-H resonance (δ 0.63) in HTh[N(SiMe_3)_2]_3 is not unusual since the hydride resonances in $(Me_5C_5)_2MH_2$ (M = Ti, Zr, or Hf) appear at δ 0.28, ¹¹ 7.46, ¹² and 15.6¹² respectively.

The spectroscopy indicates that the hydrides are indeed authentic mononuclear compounds with actinide-hydrogen bonds. This conclusion is further supported by chemical reactions (Scheme I). The hydrides can be converted quantitatively into the chloro-species $ClM[N(SiMe_3)_2]_3$ and chloroform (detected by ¹H NMR spectroscopy), a well-known reaction of transition metal hydrides.¹⁴ The hydrides may also be converted to the tetra-





Scheme I

hydroborate derivatives by reaction of borane in tetrahydrofuran.¹⁵ Further, the hydrogen atom can be removed by n-butyllithium.¹⁶ Though we have been unable to isolate the lithium-containing compounds, they may be converted to MeM[N(Si- $Me_3)_2]_3$ or DM[N(SiMe_3)_2]_3 by further reaction with MeBr or CF₃CO₂D, respectively. Thus, the hydrogen atom in these actinide compounds is hydridic as is to be expected for a hydrogen atom bonded to an electropositive metal atom. Indeed, these actinide hydrides might be viewed profitably as a hydrocarbon soluble source of sodium hydride.

Another characteristic reaction of transition metal hydrides is their ability to undergo exchange reactions with deuterium.¹⁷ Exposure of the hydrides to deuterium results not only in exchange of the hydridic hydrogen atom but in exchange of all fifty-five hydrogen atoms in the molecule (Scheme I). Not surprisingly this process is reversible, <u>viz</u>., exposure of the perdeuterometaldeuteride to hydrogen yields the hydride. The extent of deuteration was determined by hydrolyzing the compound (NaOD in D_2O) and analyzing the $[(CD_3)_3Si]_2ND$ by mass spectrometry. In the case of uranium, the extent of deuteration was \geq 97% and that for thorium was \geq 93%.

The mechanism of this exchange process is of obvious interest. Hydrogen for deuterium exchange processes are generally explained by a series of oxidative-addition, reductive-elimination cycles.¹⁷ This type of a mechanism can be invoked for uranium (IV) since the hexavalent oxidation state is well-known; however, thorium (VI) is unknown. The reverse process, a series of reductive-elimination,

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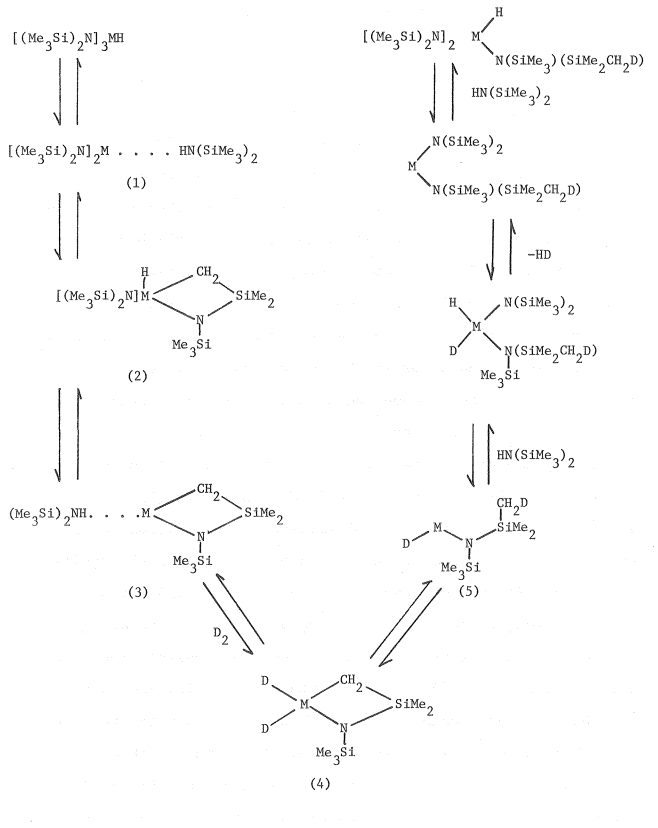
oxidative-addition cycles, is worthy of consideration. Though neither thorium (II) nor uranium (II) compounds have been isolated, they in principle could exist as transition states. This pathway, shown in Scheme II, has its origins in cyclopentadienyl- and pentamethylcyclopentadienylmetal chemistry.¹⁸

The first step is a reductive-elimination of bis(trimethylsilyl)amine giving (1) which undergoes an oxidative-addition of the γ -hydrogen atom giving metallocycle (2).¹⁹ Another reductiveelimination of bis(trimethylsilyl)amine gives (3) which undergoes oxidative-addition of deuterium yielding (4). Insertion of deuterium into the metal-carbon bond of the metallocycle giving (5) will generate a species that has a deuterium in the silylamide ligand. The remaining steps in the cycle will generate monodeutero-[(Me₃Si)₂N]₃MH. Repeating this process will generate the fully deuterated compound.

The key tenant of Scheme II is reductive-elimination of $\operatorname{silylamine}$, $(\operatorname{Me}_3\operatorname{Si})_2\operatorname{NH}$. Thus the hydrides, $\operatorname{HM}[\operatorname{N}(\operatorname{SiMe}_3)_2]_3$, should exchange with perdeutero silylamine, $[(\operatorname{CD}_3)_3\operatorname{Si}]_2\operatorname{ND}$, giving a metal-containing product containing deuterium, since the equilibrium constant for the exchange of the amine with perdeutero-amine is unlikely to be zero. Stirring a solution of either (M = Th or U) undeuterated hydride with perdeuterosilylamine leads to recovery of the hydride compound with no deuterium incorporation. Thus, another mechanism is required to explain the results.

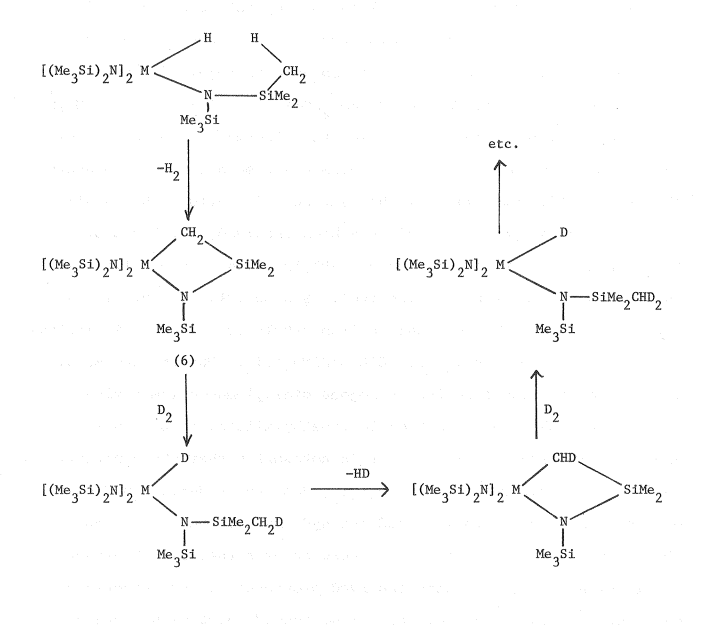
Such a mechanism is shown in Scheme III. This pathway involves the elimination of a γ -hydrogen atom of the silylamide as dihydrogen, yielding the four-membered ring metallocycle (6).

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Scheme II

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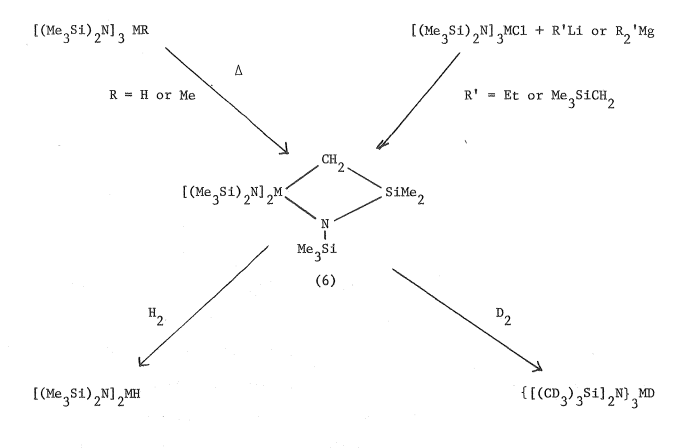
SCHEME III

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This process could occur by a variety of processes that are dis-The key tenant of this mechanistic suggestion is the postulate of the metallocycle (6) as the key intermediate and that the metallocycle will undergo hydrogen for deuterium exchange. Thus, a synthetic route to the metallocycle is required. Some related metallocycles, $Cp_2 TiCH_2 Si(Me)_2 NSiMe_3^{20}$ and $[Et_2N]_3 TaCH(Me)NEt^{21}$

have been prepared. The observation that thermal decomposition of $(R_2N)_3$ TiMe gives methane,²² and presumably a metallocycle, suggests that thermal decomposition of the hydrides will afford the metallocycle (6) and dihydrogen. This has been observed, pyrolysis of the thorium or uranium hydrides or methyls yields the metallocycles and hydrogen or methane, respectively (Scheme IV). The uranium metallocycle may also be obtained from ethyl- or trimethylsilylmethyllithium and the mono-chloride, ClU[N(SiMe₃)₂]₃. The thorium metallocycle can be made from the analogous dialkyl-magnesium deriva-In each case, ethane or tetramethylsilane, respectively, tives. are the only gases liberated. It is noteworthy that the hypothetical ethyl derivative, $EtM[N(SiMe_3)_2]_3$, does not undergo β -hydrogen elimination to give ethylene and the hydride but the ethyl group abstracts a γ -hydrogen atom. The absence of a thermal β -hydrogen elimination process has been observed previously in organoactinide chemistry.²³ The metallocycles (6) do indeed react with hydrogen or deuterium to yield the hydride or the perdeuterodeuteride, respectively. Thus our postulate that the metallocycle is a key intermediate in the hydrogen for deuterium exchange is supported. It is noteworthy that neither the chloro, nor the tetrahydroborate species, $XM[N(SiMe_3)_2]_3$ (X = Cl, BH_4 ; M = Th or U) undergo the

cussed later.



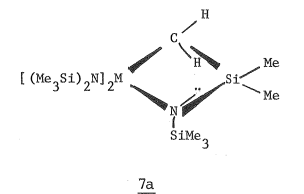
M = Th or U

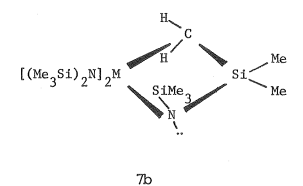


hydrogen-deuterium exchange. Although the methyl derivatives MeM[N(SiMe₃)₂]₃ do not undergo exchange with deuterium at 0°C, they undergo slow exchange at room temperature. This is due to the slow decomposition of the methyls to their respective metallocycles at room temperature, as shown by monitoring the elimination of methane by ¹H NMR spectroscopy. Further, the uranium (III) species, [(Me₃Si)₂N]₃U, does not undergo exchange with deuterium. These observations reinforce the pathway shown in Scheme III.

The conformation of the four-membered ring of the metallocycle is of considerable interest. The ¹³NMR spectrum of the methylene resonance is a triplet centered at δ 68.8 with ${}^{1}J_{_{
m CH}}$ equal to 120 Hz for the thorium compound. The coupling constant is normal for a sp^3 -hydridized carbon atom. The methylene-carbon resonance for the uranium species could not be observed, though the MeSi resonances were observable. The ¹H nuclear magnetic resonance spectrum of the thorium or uranium derivatives is temperature independent from -85° to +100°C, indicative of a planar ring. The apparent planarity could be due to a rapid interconversion $(7a \stackrel{?}{\leftarrow} 7b)$ since the methylene protons and methyl groups of the Me₂Si unit in either 7a or 7b are chemically non-equivalent. If this process was rapid on the NMR timescale, the proton and carbon atoms would appear equivalent. It is unlikely that these protons are chemical shift degenerate in the paramagnetic uranium compound, since the chemical shifts of the other protons differ greatly. On the other hand, the ring might be planar. X-ray crystallography is the only way to answer this question, but we have been unable to obtain suitable single crystals for such an analysis. In this

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regard, the spirocyclic compounds, $[t-BuNSi(Me)_2(t-Bu)N]_2M$, where M is titanium or zircomium, have planar four-membered rings.²⁴ Further, all crystal structures of $[(Me_3Si)(R)N]_XM$ compounds contain planar MN(R)(SiMe₃) units.⁴

Having shown that the metallocycle is a key intermediate in the hydrogen-deuterium exchange process, it is of interest to inquire into the pathway by which it is formed. Relevant experimental observations are, (a) the metallocycle can be prepared from the hydride by pyrolysis (ca. 200°C) of the neat solid; (b) no detectable concentration of the metallocycle is observed by ¹H NMR spectroscopy of the uranium hydride even on heating in toluene-d_g at 90°C for 20 hours; (c) stirring a pentane solution of the hydrides with deuterium at room temperature for ten minutes leads to significant deuterium incorporation, as judged by infrared spectroscopy. Taken together these observations suggest that metallocycle formation is assisted by hydrogen (deuterium). These results are in accord with the findings of Schwartz and Brintzinger²⁵ and they form a unified view of hydrogen activation in metal systems that are unable to undergo oxidativeaddition reactions.

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Experimental Section

All reactions were carried out under argon. Analyses were performed by the microanalytical laboratory of this department. Infrared spectra were recorded as Nujol mulls using a Perkin-Elmer 580 instrument. Mass spectra were obtained on a AEI-MS-12 machine equipped with a direct inlet. Proton and carbon NMR spectra were obtained at 89.56 MHz and 22.50 MHz respectively on a JEOL-FX 90 Q instrument in benzene-d₆ solution. The chemical shifts are expressed in δ -units relative to tetramethylsilane. Solution magnetic susceptabilities were determined by Evans' method using a Varian T-60 spectrometer.²⁶ Gas chromatographic analyses were performed on a Varian Moduline Series 2700 instrument equipped with a thermal conductivity detector. The columns used were 10 ft. x 0.25 in. stainless steel containing either 50-80 mesh Porapak Q or 10% propylene carbonate on 80-100 mesh Chromsorb Q.

<u>Hydridotris[bis(trimethylsilyl)amido]Thorium</u>. (a) <u>From</u> <u>Thorium Tetrachloride</u>. Sodium bis(trimethlylsilyl)amide (17.1 g, 0.0935 mol) in tetrahydrofuran (200 mL) was added to thorium tetrachloride (8.69 g, 0.0232 mol) suspended in tetrahydrofuran (200 mL). The suspension was refluxed for 84 h, allowed to settle, and the supernatant was decanted. The residue was extracted with tetrahydrofuran (2 x 40 mL) and the extracts were combined with the supernatant. The solvent was removed from the extracts under reduced pressure and the sticky mass was extracted with pentane (4 x 50 mL). The extracts were combined, concentrated to <u>ca</u>. 40 mL and cooled (-70°C). The white needles (9.8 g, 60%) were collected and dried under vacuum, mp 145-147°C. <u>Anal</u>. Calcd for $C_{18}H_{55}N_3Si_6Th$:

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C, 30.3; H, 7.71; N, 5.89. Found: C, 30.2; H, 7.83; N, 5.94. The mass spectrum shows a $(M-2)^+$ ion at 711. The Th-H absorption is at 1480 cm⁻¹ in the IR spectrum. ¹H NMR consists of singlets at δ 0.63 and 0.37 due to Th<u>H</u> and Th[N(SiMe_3)₂]₃, respectively. The ¹³C NMR consists of a quartet centered at δ 6.80 (J_{CH} = 118 Hz). (b) <u>From ClTh[N(SiMe_3)₂]₃</u>. Sodium bis(trimethylsilyl)amide (0.21 g, 0.0013 mol) in tetrahydrofuran (30 mL) was added to chlorotris-[bis(trimethylsilyl)amido]thorium (0.88 g, 0.0012 mol) in tetrahydrofuran (20 mL). After stirring at 60°C for 2 hr. the tetrahydrofuran was removed under reduced pressure and the residue was exposed to vacuum at 50°C for 2 hr. Extraction of the dry solid with pentane (2 x 30 mL) followed by filtration and concentration of the filtrate to <u>ca</u>. 6 mL gave white needles (0.75 g, 89%) on cooling (-70°C). The needles were shown to be HTh[N(SiMe₃)₂]₃ by IR and mp.

<u>Hydridotris[bis(trimethylsilyl)amido]Uranium</u>. (a) <u>From</u> <u>Uranium Tetrachloride</u>. Sodium bis(trimethylsilyl)amide (16.4 g, 0.0897 mol) in tetrahydrofuran (200 mL) was added to a solution of uranium tetrachloride (8.44 g, 0.0222 mol) in tetrahydrofuran (200 mL). The brown-green solution was refluxed for 20 hr., cooled to room temperature and the tetrahydrofuran was removed under reduced pressure. The sticky brown mass was exposed to vacuum at 50°C for 4 hr., then extracted with pentane (4 x 100 mL). The combined extracts were filtered, concentrated to <u>ca</u>. 40 mL and cooled (-70°C). The light brown needles (7.8 g, 48%) were collected and dried in vacuum, mp 97-98°C. <u>Anal</u>. Calcd for $C_{18}H_{55}N_{3}Si_{6}U$: C, 30.0; H, 7.64; N, 5.83. Found: C, 29.8; H, 7.74; N, 5.91.

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The mass spectrum contained a $(M-2)^+$ at 717. The U-H stretching frequency was at 1430 cm⁻¹ in the infrared. The solution (benzene) magnetic moment at 31°C was 2.62 B.M. The ¹H NMR spectrum consisted of a singlet at δ -19.5 ($v_{\frac{1}{2}} = 9$ Hz) and the ¹³C NMR spectrum consisted of a quartet centered at δ -40.8 ($J_{CH} = 118$ Hz). The halfwidth of the ¹³C{¹H} resonance was 4 Hz. The uranium hydride was also prepared from ClU[N(SiMe₃)₂]₃ and NaN(SiMe₃)₂ in refluxing tetrahydrofuran in a manner similar to that of HTh[N(SiMe₃)₂]₃ in 75% yield.

Reactions of Hydridotris[bis(trimethylsilyl)amido]Actinides (a) Carbon Tetrachloride. The thorium hydride (0.37 g, with 0.0052 mol) in pentane (40 mL) was treated with carbon tetrachloride (6.0 x 10^{-3} mL, 0.62 mmol). After stirring for 15 min. the volatile material was removed under reduced pressure and the residue was extracted with pentane (2 x 30 mL). The extracts were combined, concentrated to ca. 10 mL and cooled (-20°C). The colorless prisms were collected, dried under vacuum, and shown to be ClTh[N(SiMe₃)₂]₃ by IR and mp. Yield was 0.24 g (62%). In a separate experiment, carbon tetrachloride was added to the hydride in tetrahydrofuran-d_g in a NMR tube. Chloroform was the only organic product observed by NMR. The uranium hydride behaves similarily; the yield of ClU[N(SiMe₃)₂]₃ was 70%. (b) <u>n-Butyllithium and</u> Methylbromide. n-Butyllithium (0.12 mL of a 2.4 M hexane solution, 0.29 mmol) was added to a solution of hydridotris[bis(trimethylsilyl)amido]uranium (0.20 g, 0.00028 mol) in pentane (35 mL) at The dark red solution was stirred for 2 min., then rapidly 35°C. cooled to 0°C. Methylbromide was bubbled through the solution

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for 2 min., and the light brown solution was stirred for 10 min. (0°C). The solution was filtered and the volatile material was removed under reduced pressure. Crystallization of the brown residue from pentane (-20°C) yielded brown-yellow crystals which were collected and dried under vacuum. The crystals (0.16 g, 78%) were identified as $Meu[N(SiMe_3)_2]_3$ by comparison of their IR and mp with those of an authentic specimen. The methylthorium analogue, MeTh[N(SiMe₃)₂]₃, was prepared in 50% yield in an analogous Borane in Tetrahydrofuran. The tetrahydrofuran manner. (c) complex of borane (1 mL of a 1.0 M tetrahydrofuran solution, 1.0 mmol) was added to hydridotris[bis(trimethylsilyl)amido]thorium (0.21 g, 0.29 mmol) in tetrahydrofuran (50 mL) at 0°C. After stirring the colorless solution for 15 h at 0°C, the tetrahydrofuran was evaporated and pentane (50 mL) was added to the sticky residue. Filtration, followed by concentration of the filtrate to ca. 15 mL and cooling (-20°C) yielded colorless prisms (0.12 g, 57%) which were collected and dried under vacuum. The crystals were identified as $BH_4Th[N(SiMe_3)_2]_3$ by IR and mp.⁶ The uranium analogue was prepared similarly in 60% yield.

Deuterotris[bis(trimethylsilyl)amido]Thorium. A solution of hydridotris[bis(trimethylsilyl)amido]thorium (0.20 g, 0.28 mmol) in pentane (40 mL) was maintained at 35°C while a solution of n-butyllithium (0.12 mL of a 2.4 M hexane solution, 0.29 mmol) was added. The pale yellow solution was stirred for 2 min., then treated with monodeutero-trifluoroacetic acid (2.3 x 10^{-3} mL, 0.30 mmol). The volatile material was immediately removed under reduced pressure and the white residue was crystallized as white needles from pentane (0.17 g, 85%), mp 144-146°C. <u>Anal</u>. Calcd for $C_{18}H_{54}DN_{3}Si_{6}Th$: C, 30.3; H, 7.83; N, 5.89. Found: C, 30.1; H, 7.66; N, 5.92. The Th-D stretching frequency in the infrared was observed at 1060 cm⁻¹. The ¹H NMR spectrum consisted of a singlet at δ 0.38. The deuteride was also prepared (85% yield) from ClTh[N(SiMe_3)_2]_3 and NaN(SiMe_3)_2 in refluxing tetrahydrofurand₈.

Deuterotris [bis(trimethylsilyl)amido]uranium. n-Butyllithium (0.16 mL of a 2.4 M hexane solution, 0.38 mmol) was added to a solution of hydridotris[bus(trimethylsilyl)amido]uranium (0.28 g, 0.39 mmol) in pentane (60 mL) maintained at 35°C. The dark red solution was immediately treated with mono-deuterotrifluoroacetic acid (3.0 x 10^{-3} mL, 0.39 mmol). The volatile material was immediately removed under reduced pressure from the browngreen solution. The brown residue was crystallized as light brown needles from pentane (-20°C) in 90% (0.25 g) yield, mp 98-99°C. Anal. Calcd for C₁₈H₅₄DN₃Si₆U: C, 30.0; H, 7.78; N, 5.83. Found: C, 30.1; H, 7.59; N, 5.90. The U-D stretching frequency was observed at 1020 cm^{-1} in the IR. The ¹H NMR spectrum was a singlet at δ -19.4 (28°C) with $v_{\frac{1}{2}} = 10$ Hz. The deuteride was also prepared from $Clu[N(SiMe_3)_2]_3$ and $NaN(SiMe_3)_2$ in refluxing tetrahydrofuran-d $_{8}$ in 90% yield.

 $\frac{\text{Preparation of } [(\text{Me}_{3}\text{Si})_{2}\text{N}]_{2}\text{ThCH}_{2}\text{Si}(\text{Me})_{2}\text{NSiMe}_{3}.$ (a) From $\frac{\text{MeTh}[\text{N}(\text{SiMe}_{3})_{2}]_{3}.$ Methyltris[bis(trimethylsilyl)amido]thorium (1.34 g, 0.00184 mol) contained in a Schlenk tube (175 mL capacity) under one atmosphere of argon was heated to 140°C over 45 min. using an oil bath. The colorless melt was carefully run into a

thin film round the bottom of the Schlenk tube and heated at 150-160°C for 10 min (until sublimation was evident). The melt was then allowed to cool to room temperature and purified either by vacuum sublimation (100°C) in ca. 60% yield or crystallized from pentane (-70°C) in ca. 60% yield. The colorless needles melted at 109-111°C. Anal. Calcd for C₁₈H₅₃N₃Si₆Th: C, 30.3; H, 7.45; N, 5.90. Found: C, 30.0; H, 7.14; N, 5.69. A molecular ion, M^+ , was observed at 711 in the mass spectrometer. The ${}^{\perp}H$ NMR spectrum consisted of singlets at δ 0.37, 0.38, 0.49, and 0.56 in area ratio 36:9:2:6 due to [(Me_3Si)2N]2Th, Me_3SiNTh, CH2Th, and \underline{Me}_2 SiCH₂Th, respectively. The ¹³C NMR spectrum consisted of quartets centered at δ 3.46 (J $_{\rm CH}$ =117 Hz), 4.52 (J $_{\rm CH}$ = 117 Hz), 5.55 (J_{CH} = 118 Hz) due to $[(\underline{Me}_3Si)_2N]_2Th$, \underline{Me}_3SiN , and \underline{Me}_2SiCH_2Th , respectively, and a triplet centered at δ 68.8 (J $_{\rm CH}$ = 120 Hz) due to Me_SiCH_Th. In a separate experiment, the gases formed upon pyrolysis were examined by gas chromatography and shown to be only methane. The metallocycle may be obtained by pyrolysis of $HTh[N(SiMe_3)_2]_3$ in a similar fashion to a final temperature of 180-190°C in 20% yield. (b) From $ClTh[N(SiMe_3)_2]_3$ and Et_2Mg . Diethylmagnesium (0.30 mL of a 0.88 M diethyl ether solution, 0.26 mmol) was added to a solution of chlorotris[bis(trimethylsilyl)amido]thorium (0.38 g, 0.00051 mol) in diethyl ether (100 mL) 0°C. After stirring at 0°C for 24 h the solvent was removed under reduced pressure at 0°C. The pale yellow residue was extracted with pentane (2 x 50 mL), filtered, and the filtrate was concentrated to ca. 5 mL and cooled (-70°C). The colorless blocks were collected and dried under vacuum. Yield was 0.28 g

(77%). The compound was identified by comparison of its IR and mp with an authentic specimen. Gas chromatographic analysis of the vapor above the reaction mixture prior to commencement of the work-up showed that ethane was the only hydrocarbon present. Reaction of the chloro-amide with $(Me_3SiCH_2)_2Mg$ under similar conditions gives the metallocycle and tetramethylsilane.

Preparation of $[(Me_3Si)_2N]_2UCH_2Si(Me)_2NSiMe_3$. (a) From MeU[N(SiMe_3)_2]_3. The uranium metallocycle was prepared in a similar manner as that of the thorium metallocycle just described in ca. 85% yield. The compound was isolated either by vacuum sublimation (110°C) or crystallization from pentane (-70°C) as yellow needles, mp 126-129°C. Anal. Calcd for C₁₈H₅₃N₃Si₆U: C, 30.1; H, 7.39; N, 5.86. Found: C, 29.8; H, 7.14; N, 5.77. The mass spectrum contained a M^+ ion at 717. The magnetic susceptibility in benzene solution (30°C) was 2.71 B.M. The ¹H NMR spectrum (28°C) consists of singlets at δ 2.08, -9.90, -23.3, and -128.6 in area ratio 6:9:36:2 due to \underline{Me}_2SiCH_2U , \underline{Me}_3SiNU , [(\underline{Me}_3Si)₂-N]U and Me₂Si<u>CH</u>₂U, respectively. The ¹³C NMR spectrum (28°C) afforded quartets centered at δ 49.6 (J $_{\rm CH}$ = 118 Hz), 24.5 (J $_{\rm CH}$ = 118 Hz), and -40.8 ($J_{CH} = 118$ Hz) due to $\underline{Me}_2 \text{SiCH}_2 U$, $\underline{Me}_3 \text{SiNU}$, and $[(\underline{Me}_3 \text{Si})_2 N]_2 U$, respectively. The methylene carbon atoms could not be observed. Gas chromatographic analysis of the gases above the melt indicated that methane was the only organic hydrocarbon present. (b) From ClU[N(SiMe_3)_2]_ and EtLi. Ethyllithium (6.0 mL of a 0.47 M toluene solution, 2.8 mmol) was added to a solution of chlorotris[bis-(trimethylsilyl)amido]uranium (2.1 g, 0.0028 mol) in pentane (150 mL) The resulting dark yellow solution was stirred at 0°C for at 0°C.

16 h, then the volatile material was removed under reduced pressure at 0°C. The brown mass was exposed to vacuum for 3 h. Extraction of the dry residue with pentane (2 x 40 mL) followed by filtration and concentration of the filtrate to <u>ca</u>. 20 mL and cooling (-70°C) yielded bright yellow needles. The needles (1.8 g, 90%) were collected and dried under vacuum. The needles were identified by IR and mp. The gases above the reaction mixture were shown to be ethane by gas chromatographic analysis. The analogous reaction using Me_3SiCH_2Li in hexane rather than EtLi yielded the metallocycle in 60% yield and tetramethylsilane as the sole gaseous product.

<u>Preparation of $\{[(CD_3)_3Si]_2N\}_3UD$ </u>. Hydridotris[bis(trimethylsilyl)amido]uranium (0.76 g, 0.00011 mol) in pentane (80 mL) was stirred under deuterium in a thick-walled bottle (100 mL) fitted with a pressure-cap at a total pressure of 15 atm for 48 h. The gas above the solution was vented and the bottle was recharged. After a further 48 h the process was repeated three additional times using reaction times of 24 h. Finally the solution was transferred to a Schlenk tube and the solvent was removed under reduced pressure. Crystallization of the brown residue from pentane (-70°C) gave light brown blocks which were collected and dried under vacuum. Yield was 0.52 g (64%), mp 95-97°C. Anal. Calcd for C₁₈D₅₅N₃Si₆U: C, 27.9; D, 14.2; N, 5.43. Found: C, 27.8; D, 13.8; N, 5.34. The infrared spectrum contained C-D and U-D stretching frequencies at 2210 and 1027 cm^{-1} , respectively. In order to assay the extent of deuteration, the deuterated specimen (0.30 g, 0.00042 mol) was finely ground and exposed to vacuum

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at 50°C. The flask was cooled to 0°C and a solution for 3 h of sodium deuteroxide in deuterium oxide (0.5 mL of a 4M solution, 2.0 mmol) was added. The closed system was stirred for 1 min. at 30°C and the volatile material was distilled into a cooled (-196°C) receiver under vacuum. The distillate was allowed to stand over molecular sieves (4X) for 30 min. and the clear organic product (ca. 0.2 mL) was decanted into a dry capillary tube. The mass spectrum of the liquid was recorded at 15 eV and 70 eV. Analysis of the envelope from m/e 160-180 by standard methods revealed that the amine was \geq 97% deuterated.²⁷ The metallocycle, [(Me_3Si)2N]2UCH2Si(Me)2NSiMe3, was deuterated in an analogous fashion to that of its thorium analogue, described above. The isolated yield was 60%, m.p. 144-147°C. Anal. Calcd for C₁₈D₅₅N₃Si₆Th: C, 28.1; D, 14.3; N, 5.49. Found: C, 28.4; D, 13.7; N, 5.54. The deuterium content of the amine upon deuterolysis was > 91%, statistically distributed.

Other Deuteration Studies. The hydrides do not exchange with benzene-d₆ (refluxing, 24 h) nor with $[(CD_3)_3Si]_2ND$ (pentane solution at room temperature for 24 h). Neither $(Me_3Si)_2NH$, NaN $(SiMe_3)_2$, U[N $(SiMe_3)_2$]₃ nor the derivatives XM[N $(SiMe_3)_2$]₃, where X is Cl or BH₄ and M is Th or U, exchange with deuterium (1-20 atm) at room temperature for periods of 24-36 h. The methyl derivatives, MeM[N $(SiMe_3)_2$]₃ where M is Th or U, do not exchange with deuterium (1-20 atm) in pentane at 0°C over a 36 h period, Acknowledgement This work was supported by the Division of Nuclear Sciences, Office of Basic Energy Sciences, U.S. Department of Energy under Contract W-7405-ENG-48.

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