

Pnictogen-Hydride Activation by (silox)₃Ta (silox = ^tBu₃SiO); Attempts to Circumvent the Constraints of Orbital Symmetry in N₂ Activation

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Activation of N₂ by (silox)₃Ta (**1**, silox = ^tBu₃SiO) to afford (silox)₃Ta=N–N=Ta(silox)₃ (**1**₂-N₂) does not occur despite $\Delta G^{\circ}_{\text{calcd}} = -55.6$ kcal/mol because of constraints of orbital symmetry, prompting efforts at an independent synthesis that included a study of REH₂ activation (E = N, P, As). Oxidative addition of REH₂ to **1** afforded (silox)₃HTaEHR (**2**-NHR, R = H, Me, ⁿBu, C₆H₄-*p*-X (X = H, Me, NMe₂); **2**-PHR, R = H, Ph; **2**-AsHR, R = H, Ph), which underwent 1,2-H₂-elimination to form (silox)₃Ta=NR (**1**=NR; R = H, Me, ⁿBu, C₆H₄-*p*-X (X = H (X-ray), Me, NMe₂, CF₃)), (silox)₃Ta=PR (**1**=PR; R = H, Ph), and (silox)₃Ta=AsR (**1**=AsR; R = H, Ph). Kinetics revealed NH bond-breaking as critical, and As > N > P rates for (silox)₃HTaEHP (**2**-EHP) were attributed to (1) $\Delta G^{\circ}_{\text{calc}}(\text{N}) < \Delta G^{\circ}_{\text{calc}}(\text{P}) \sim \Delta G^{\circ}_{\text{calc}}(\text{As})$; (2) similar fractional reaction coordinates (RCs), but with RC shorter for N < P ~ As; and (3) stronger TaE bonds for N > P ~ As. Calculations of the pnictidenes aided interpretation of UV–vis spectra. Addition of H₂NNH₂ or H₂N–N(^oNC₂H₃Me) to **1** afforded **1**=NH, obviating these routes to **1**₂-N₂, and formation of (silox)₃MeTaNHNH₂ (**4**-NHNH₂) and (silox)₃MeTaNH(^oNCHMeCH₂) (**4**-NH(azir)) occurred upon exposure to (silox)₃Ta=CH₂ (**1**=CH₂). Thermolyses of **4**-NHNH₂ and **4**-NH(azir) yielded [(silox)₂TaMe](μ -N _{α} HN _{β})(μ -N _{γ} HN _{δ} H)-[Ta(silox)₂] (**5**) and [(silox)₃MeTa](μ - η^2 -N, N: η^1 -C-NHNHCH₂CH₂CH₂)[Ta(κ -O, C-OSi^tBu₂CMe₂CH₂)(silox)₂] (**7**, X-ray), respectively. (silox)₃Ta=CPPH₃ (**1**=CPPH₃, X-ray) was a byproduct from Ph₃PCH₂ treatment of **1** to give **1**=CH₂. Addition of Na(silox) to [(THF)₂Cl₃Ta]₂(μ -N₂) led to [(silox)₂ClTa](μ -N₂) (**8**-Cl), and via subsequent methylation, [(silox)₂MeTa]₂(μ -N₂) (**8**-Me); both dimers were thermally stable. Orbital symmetry requirements for N₂ capture by **1** and pertinent calculations are given.

Introduction

Since its discovery, (silox)₃Ta (**1**, silox = ^tBu₃SiO) has produced a remarkable array of products derived from small molecule activation because of its proclivity toward oxidative addition.¹ For example, 1/2 CO is cleaved to afford 1/2 (silox)₃-TaO and 1/4 (silox)₃TaCCTa(silox)₃,² and *p*-CF₃-C₆H₄-NH₂ undergoes a rare C–N bond scission to provide (silox)₃-Ta(NH₂)(C₆H₄-*p*-CF₃).³ Other activations include C–O^{3,4} and E–H (E = N, P, As) bonds,⁵ and the reversible addition of a CH bond at the *para*-position of 2,6-lutidine.⁶ By varying

the siloxide ligand, it has been shown that ^tBu₃SiO (silox) is unique in its ability to sterically protect the Ta(III) center of **1**,⁴ thereby permitting the investigation of these activations. Interestingly, one substrate that is expected to react with **1**, dinitrogen, is noticeably absent.

Schrock's seminal studies of N₂ activation have their origins in the synthesis of L_{*n*}X₃Ta=N–N=TaX₃L_{*n*}^{7–9} and related "dinitrogen" complexes, which were typically generated via the reduction of X_{*n*}M in the presence of L and N₂.^{10–14} The

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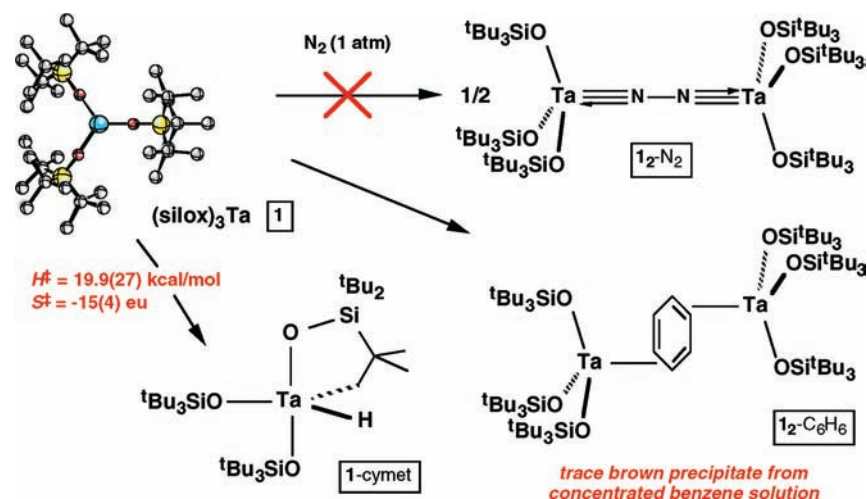
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Scheme 1



species are most aptly described as diimido complexes, as structural parameters clearly reveal the TaN multiple bonds ($\text{BO} \sim 3$) and N–N single bond. Given the amount of reduction of N₂ implied by the structures, it is not surprising that the compounds do not relinquish the bound “dinitrogen”, even under harsh conditions. However, it is surprising that (silox)₃Ta (**1**, silox = tBu₃SiO), does not appear to bind dinitrogen to form (silox)₃Ta(N₂)_x or reduce it to (silox)₃Ta=N–N=Ta(silox)₃ (**I**₂-N₂). As Scheme 1 shows, in the presence of N₂, **1** cyclometallates to afford (silox)₂HTa(κ -O,C-OSi-tBu₂CMe₂CH₂) (**1-cymet**),^{6,15} and if sufficiently concentrated in benzene solution, a small amount is converted to the bridging benzene complex, [(silox)₃Ta]₂(μ -C₆H₆) (**I**₂-C₆H₆).^{6,16} Since the thermodynamics, at least according to Schrock’s preceding studies, certainly suggests that N₂ activation by **1** should be viable, it is likely that orbital symmetry constraints^{17,18} play a crucial role in the lack of observed N₂ chemistry.^{19–24} Even the cyclometalation process, which has not been shown to exhibit reversible behavior, is likely to be constrained by orbital symmetry. Herein are described attempts to circumvent these constraints and synthesize **I**₂-N₂ via alternate methods. In concert with these efforts, a significant study of REH₂ (E = N, P, As) oxidative addition and

1,2-H₂-elimination from (silox)₃HTaEHR is also presented.⁵ Computational studies support the contention that **I**₂-N₂ is considerably more stable than **1** and N₂.

Results

Pnictogen-Hydride Reactivity. 1. Tantalum Pnictogenide-Hydrides.^{25,26} Routes to the desired dinitrogen complex can be envisaged through the oxidative addition of N–H bonds to (silox)₃Ta (**1**), and Scheme 2 shows that there is ample evidence for this approach, not only for nitrogen, but for P–H and As–H bonds as well.^{5,27–31} Treatment of **1** with primary amines^{3,5,25,26} or ammonia^{25,32} in hydrocarbon solvent (or C₆D₆ in the case of NMR tube experiments) led to the formation of (silox)₃HTaNHR (**2-NHR**; R = Me, tBu, C₆H₄-p-X (X = H, Me, NMe₂)) and (silox)₃HTaNH₂ (**2-NH₂**) in near quantitative yield (¹H NMR), although only the latter was isolated because of the subsequent 1,2-H₂-elimination to form the imido derivatives. An alternative synthesis involving the addition of RNH₂ to (silox)₃TaH₂ (**2-H**) provided **2-NHR** (R = C₆H₄-p-X (X = H, NMe₂, CF₃)) accompanied by the evolution of dihydrogen. IR spectra of **2-NH₂** and **2-NHPh** revealed two and one $\nu(\text{NH})$ stretches, respectively, and $\nu(\text{TaH})$ stretches at 1786 cm⁻¹ for the former and 1810 cm⁻¹ for the latter, consistent with terminal tantalum hydrides.³³ The amide-hydrides appeared colorless, and their NMR spectra are provided in Table I. Diagnostic, broad, amide-hydrogen resonances ranged from δ 4.56 for **2-NH₂** to δ 7.41 for **2-NH(C₆H₄-NMe₂)**, and these were accompanied by

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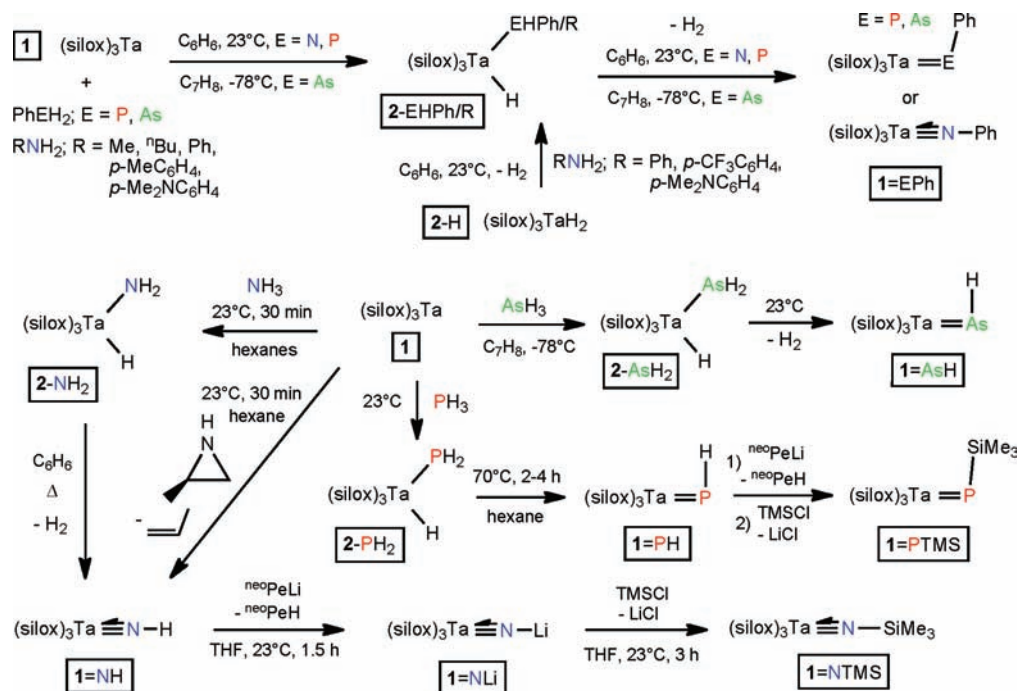
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Scheme 2



typical downfield hydride shifts ranging from δ 19.96 (2-NHMe) to δ 21.76 (2-NH(C₆H₄-CF₃)). Their structures are likely to be trigonal bipyramidal (tbp) with equatorial amide and hydride ligands or square pyramidal with basal Ta-H and Ta-NHR groups as observed for the C-N bond oxidative addition product, (silox)₃-(H₂N)TaC₆H₄-*p*-CF₃, which was previously crystallographically characterized.³

The related exposure of (silox)₃Ta (1) to PhPH₂ led to the formation of bright yellow (silox)₃HTaPPh (2-PHPh), which was crystallized from pentane in only 39% yield because of its high solubility. The parent (silox)₃HTaPH₂ (2-PH₂) complex, a pale yellow species, was isolated in 54% yield, but it was accompanied by an insoluble brick red material, which may be the double activation product (silox)₃HTa-PH-TaH(silox)₃, based on the near insolubility of the size-related, crystallographically characterized (silox)₃Nb=PH-Nb(silox)₃ species.³⁴ Phosphide 2-PH₂ manifests $\nu(\text{PH})$ bands at 2275 and 2285 cm⁻¹ in its IR spectrum, while 2-PHPh exhibits only one at 2320 cm⁻¹, and these were accompanied by tantalum hydride stretches at 1775 and 1790 cm⁻¹, respectively. The phenyl phosphide exhibited a ddt at δ 55.5 in its ³¹P NMR spectrum, with ¹J_{PH} = 243 Hz, ²J_{PH} = 56 Hz, and a coupling to the *o*-Ph hydrogens of ³J_{PH} = 9 Hz. The ³¹P NMR signal for 2-PH₂ was very different, resonating at -142.39 ppm, with ¹J_{PH} = 177 Hz and ²J_{PH} = 3 Hz. The difference prompted a molecular weight study of 2-PH₂, and this was fully consistent with a monomer: *M_r* (found) 858(68); calcd 861. It is plausible that the two phosphide complexes possess substantially different Ta-P-R angles because of disparate steric factors, and that their chemical shifts reflect these features in addition to the electronic effects of H versus Ph. In addition, the

dramatically lower ²J_{PH} for 2-PH₂ (3 vs 56 Hz) and a related, smaller ³J_{HH} = 2 (vs 9 Hz for 2-PHPh) value suggests that the geometries of the phosphide hydrides may also be different at the metal.

Preparation of the arsenide complexes proved to be trickier, as each species rapidly lost dihydrogen to form the corresponding arsinidenes. Treatment of (silox)₃Ta (1) with PhAsH₂ at -78 °C in toluene-*d*₈ provided (silox)₃HTaAsHPh (2-AsHPh), but the complex was stable at low temperature only if 1,4-cyclohexadiene, a radical trap, was present. Given the danger accorded the use of toxic AsH₃, it was generated via hydrolysis of Zn₃As₂ with HCl, and used immediately after passing through a cold trap. In this fashion, exposure of 1 to AsH₃ in toluene-*d*₈ at -78 °C, in the presence of 1,4-cyclohexadiene, afforded (silox)₃HTaAsH₂ (2-AsH₂). Characterization of the two arsenide hydrides was limited to ¹H NMR spectra at -78 °C, and the hydride resonances at δ 24.47 (d, 2-AsHPh) and δ 24.98 (t, 2-AsH₂) that were observed were the furthest downfield for this genre of compounds. The hydrides were coupled to AsH protons at δ 5.53 (d, ³J_{HH} = 8 Hz) in 2-AsHPh and at δ 3.13 (d, ³J_{HH} = 5 Hz) in 2-AsH₂.

2. Tantalum Pnictide Derivatives.³⁵⁻⁴⁶ The pnictide-hydrides addressed in the previous section all underwent

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Table 1. NMR Spectral Assignments^a for (silox)₃M Pnictogen Derivatives and Selected IR Data

compound	¹ H NMR (δ (J(Hz)))		¹³ C{ ¹ H} NMR (δ (J(Hz)))		³¹ P/ ¹⁵ N NMR (δ (J(Hz))) and IR (cm ⁻¹)
	^t Bu	H/ER	CM _{e3} ,C(CH ₃) ₃	ER/R	
(silox) ₃ HTaNH ₂ (2-NH ₂)	1.29	4.56 20.04	23.83, 30.83		-229.37 (td, 72 ^b , 5 ^c) 1786, ν(TaH) 3363, ν(NH) 3453, ν(NH)
(silox) ₃ HTaNHMe (2-NHMe)	1.31	3.34 (d,7) 5.42 (q,7) 19.96			
(silox) ₃ HTaNH ⁿ Bu (2-NH ⁿ Bu) ^d	1.32	0.85 (t, CH ₃) 3.85 (dd, NCH ₂) 5.66 (t, NH) 19.99			
(silox) ₃ HTaNHPh (2-NHPh) ^e	1.26	6.76 (m, <i>p</i> -H) 7.15–7.26 (m, <i>o</i> -, <i>m</i> -H) 7.29 (NH) 21.47			1810, ν(TaH) 3350, ν(NH)
(silox) ₃ HTaNH(C ₆ H ₄ - <i>p</i> -Me) ^e (2-NHC ₆ H ₄ - <i>p</i> -Me)	1.28	2.17 (<i>p</i> -Me) 6.99 (m, <i>o</i> -, <i>m</i> -H) 7.33 (NH) 21.38			
(silox) ₃ HTaNH(C ₆ H ₄ - <i>p</i> -NMe ₂) ^e (2-NHC ₆ H ₄ - <i>p</i> -NMe ₂)	1.30	2.55 (NMe ₂) 6.66 (d, <i>m</i> -H) 7.03 (d, <i>o</i> -H) 7.41 (NH) 21.13			
(silox) ₃ HTaNH(C ₆ H ₄ - <i>p</i> -CF ₃) ^e (2-NHC ₆ H ₄ - <i>p</i> -CF ₃)	1.23	6.91 (d, <i>m</i> -H) 7.04 (NH) 7.43 (d, <i>o</i> -H) 21.76			
(silox) ₃ Ta(NH ₂) ₂ (3-(NH ₂) ₂)	1.30	4.14			-241.62 (t, 71 ^b)
(silox) ₃ HTaPH ₂ (2-PH ₂)	1.28	2.40 (dd, 177 ^f , 2) 21.72 (dt, 3 ^g , 2)	23.75, 30.65		-142.39 (td, 177 ^f , 3 ^g) 1775, ν(TaH) 2275, ν(PH) 2285, ν(PH) 55.5 (ddt, 243 ^f , 56 ^g , 9 ^h)
(silox) ₃ HTaPPh (2-PHPh) ^{e,i}	1.26	6.88 (t, <i>p</i> -H) 7.21 (dd, 243 ^f , 9) 7.72 (t, <i>o</i> -H, 9) 21.62 (dd, 56 ^g , 9)	23.58, 30.66	126.31 127.53 127.61 132.51 (8 ^j)	1790, ν(TaH) 2320, ν(PH)
(silox) ₃ HTaAsH ₂ (2-AsH ₂)	1.29	3.13 (d,5) 24.98 (t,5)			
(silox) ₃ HTaAsHPh (2-AsHPh) ^e	1.26	5.53 (d, 8) 6.96 (m, <i>p</i> -H) 7.84 (d, <i>o</i> -H) 24.47 (d,8)			
(silox) ₃ Ta=NH (1=NH)	1.29	6.01 (br) ^k	23.78, 30.65		-60.60 (¹ J _{NH} = 76) 3470, ν(NH)
(silox) ₃ (H ₃ N)Ta=NH (1=NH(NH ₃))	1.28	0.80 (br, NH ₃) 5.67 (br, NH)			
(silox) ₃ Ta=NLi (1=NLi) ^l	1.20		24.40, 31.43		
(silox) ₃ Ta=NCH ₃ (1=NMe)	1.29	3.89	23.42, 30.54	47.88	
(silox) ₃ Ta=N ⁿ Bu (1=N ⁿ Bu) ^m	1.29	0.92 (t, CH ₃) 1.60 (m, CH ₂) 4.23 (t, NCH ₂)			
(silox) ₃ Ta=NSiMe ₃ (1=NTMS)	1.28	0.28	23.59, 30.66	3.79	
(silox) ₃ Ta=NPh (1=NPh) ^e	1.27	6.75 (m, <i>p</i> -H) 7.16–7.25 (<i>o</i> -, <i>p</i> -H)	23.36, 30.55	121.82 125.35 128.31 129.41	
(silox) ₃ Ta=N(C ₆ H ₄ - <i>p</i> -Me) ^e (1=N(C ₆ H ₄ - <i>p</i> -Me))	1.28	2.14 (Me) 6.99 (d, <i>m</i> -H) 7.08 (d, <i>o</i> -H)			
(silox) ₃ Ta=N(C ₆ H ₄ - <i>p</i> -NMe ₂) ^e (1=N(C ₆ H ₄ - <i>p</i> -NMe ₂))	1.31	2.50 (NMe ₂) 6.62 (d, <i>m</i> -H) 7.13 (d, <i>o</i> -H)	23.41, 30.62	40.92 113.04 125.85 146.31 150.07	
(silox) ₃ Ta=N(C ₆ H ₄ - <i>p</i> -CF ₃) ^e	1.24	7.01 (d, <i>m</i> -H)	23.35, 30.50	123.05 (² J _{CF} = 32)	

Table 1. Continued

compound	¹ H NMR (δ (J(Hz)))		¹³ C{ ¹ H} NMR (δ (J(Hz)))		³¹ P/ ¹⁵ N NMR (δ (J(Hz))) and IR (cm ⁻¹)
	^t Bu	H/ER	CM ₃ ,C(CH ₃) ₃	ER/R	
(1 =N(C ₆ H ₄ - <i>p</i> -CF ₃))		7.38 (d, <i>o</i> -H)		125.21 125.45 (¹ J _{CF} = 271) 125.78 (³ J _{CF} = 4) 160.33 185.02 (135)	
(silox) ₃ Ta=CH ₂ (1 =CH ₂)	1.28	6.66	23.75, 30.59		
(silox) ₃ Ta=C=PPh ₃ (1 =C=PPh ₃)	1.31	7.71 (m, <i>m,p</i> -H) 7.78 (m, <i>o</i> -H)	25.21, 31.31	128.75 (<i>p</i>) 131.15 (<i>m</i>) 133.94 (<i>o</i>) 136.37 (d, 88) 175.01 (d, 53)	-4.26
(silox) ₃ Ta=PH (1 =PH)	1.30	3.57 (d, 82 ^f)			276.33 (d, 82 ^f) 2150, ν(PH)
(silox) ₃ Ta=PPh (1 =PPh) ^e	1.28	6.76 (<i>m</i> -H ^h) 7.20 (<i>o</i> -H) 7.67 (<i>p</i> -H)	23.75, 30.61	125.79 127.67 128.29 137.27 (10 ^g)	334.6 (t, 6 ^h)
(silox) ₃ Ta=PSiMe ₃ (1 =PTMS)	1.34	0.47 (d, 6 ^k)			196.07
(silox) ₃ Ta=AsH (1 =AsH)	1.30	2.04			1950, ν(AsH)
(silox) ₃ Ta=AsPh (1 =AsPh) ^e	1.29	6.81 (<i>m</i> -H) 7.21 (<i>o</i> -H) 7.86 (<i>p</i> -H)		125.83 127.93 139.61 142.59	
(silox) ₃ Ta(AsH ₂) ₂ (3 -(AsH ₂) ₂)	1.23	2.33			2180, ν(AsH) 2190, ν(AsH)
(silox) ₃ MeTaNHNH ₂ (4 -NHNH ₂)	1.26	1.03 (Me) 3.63 (NH ₂) 4.11 (NH)	23.78, 30.43	33.79	64.80 (NH ₂ , 78) ⁿ 148.47 (NH, 78) ⁿ
(silox) ₃ MeTaNH(^c NCH ₂ CMeH) (4 -NH(azir))		1.06 (d, 6, CH ₃) 1.24 (d, 8, CHMe) 1.44 (dd, 8, 4, CHH) 1.54 (d, 4, CHH)			81.99 (^c N) ⁿ 244.46 (NH, 72) ^{n,o}
[(silox) ₂ TaMe][(silox) ₂ Ta] (μ -N _α HN _β)(μ -N _γ HN _δ H) (5)	1.24 1.26 1.28 1.30 1.26	1.21 (Me) 3.00 (N _γ H) 5.42 (N _α H) 6.38 (N _δ H)	24.01, 30.83 24.10, 30.88 24.15, 31.04 24.49, 31.11 23.76, 30.39	38.31	79.50 (N _γ H, 72) ^{n,o} 131.27 (N _α H, 72) ^{n,o} 179.70 (N _δ H, 75) ^{n,o} 298.23 (N _β , 8) ^{n,p}
[(silox) ₂ ClTa] ₂ (μ -N ₂) (8 -Cl)	1.26		23.76, 30.39		
[(silox) ₂ MeTa] ₂ (μ -N ₂) (8 -Me)	1.24	1.24	23.79, 30.44	33.87	

^a Benzene-*d*₆ unless otherwise noted. ^b ¹J_{15NH}. ^c ³J_{15NH}. ^d β-CH₂ and γ-CH₂ obscured. ^e All couplings between *o*-, *m*-, and *p*-Ph positions are 8(1) Hz. ^f ¹J_{PH}. ^g ²J_{PH}. ^h ³J_{PH}. ⁱ *m*-H obscured. ^j ¹J_{PC}. ^k At 80 °C, a 1:1:1 triplet due to ¹J_{14NH} ~ 50 Hz was observed. ^l THF-*d*₈. ^m CH₂ obscured. ⁿ Obtained via HMBCAD correlation spectroscopy. ^o ¹J_{15NH}. ^p ²J_{15NH}.

1,2-H₂-elimination to form the corresponding imido (nitrene) derivatives (silox)₃Ta=NR (**1**=NR; R = H, Me, ⁿBu, C₆H₄-*p*-X (X = H, Me, NMe₂, CF₃)), phosphinidenes (silox)₃Ta=PR (**1**=PR; R = H, Ph) and arsinidenes (silox)₃Ta=AsR (**1**=AsR; R = H, Ph). An alternate preparation of **1**=NH from 2-methylaziridine proved to be convenient. While the imido derivatives were colorless, (silox)₃Ta=PH (**1**=PH) was isolated as pale orange crystals (61%), and red-violet crystals of (silox)₃Ta=PPh (**1**=PPh)⁵ were obtained in 62% yield. Parent arsinidene (silox)₃Ta=AsH (**1**=AsH) was isolated as dark orange crystals, but proved to be contaminated by

a compound tentatively identified as (silox)₃Ta(AsH₂)₂ (**3**-(AsH₂)₂; ν(AsH) = 2180, 2190 cm⁻¹), which comprised ~10% of the material. A red insoluble material also formed, and this is likely to be a dinuclear substance, as speculated for the related phosphorus chemistry above.³⁴ It is likely that **3**-(AsH₂)₂ forms from reaction of excess AsH₃ in solution with (silox)₃HTaAsH₂ (**2**-AsH₂) via the loss of H₂, or via the 1,2-AsH-addition to the Ta=As bond of **1**=AsH. Phenylarsinidene⁴⁶ (silox)₃-Ta=AsPh (**1**=AsPh)⁵ was obtained as green crystals in 85% yield from hexane.

¹H and ¹³C{¹H} NMR spectra of the pnictidenes listed in Table 1 reveal standard silox resonances and others corresponding to the R groups. The parent imide (**1**=NH), phosphinidene (**1**=PH), and arsinidene (**1**=AsH) possess varied pnictogen-hydrogen ¹H NMR spectral shifts. The imide proton is at δ 6.01, and upon warming to 80 °C, manifests a 1:1:1 triplet due to ¹⁴N-coupling (¹J_{N14H} ~ 50 Hz). The PH resonance in **1**=PH is at δ 3.57

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with $^1J_{\text{PH}} = 82$ Hz, and the arsinidene proton is observed at δ 2.04. The observation of large ^{14}N -coupling is indicative of the cylindrical symmetry expected for the triply bonded imido group. In contrast, the $^1J_{\text{PH}}$ for $\mathbf{1}=\text{PH}$ is substantially less than that found for the phosphide-hydride complexes, and is taken as an indication of further diminished s-character in the phosphinidene-hydrogen bond. Structural studies of $(\text{silox})_3\text{TaX}$ species have historically been difficult to obtain or interpret because of disorder and twinning issues common to the 3-fold core, but it is likely that the $\text{Ta}=\text{EH}$ angle ($\text{E} = \text{P}, \text{As}$) in $\mathbf{1}=\text{EH}$ is very close to 90° .

The addition of excess NH_3 to benzene solutions of $\mathbf{1}=\text{NH}$ caused the formation of a white solid, and ^1H NMR spectral analysis of the reaction mixture implicated the formation of $(\text{silox})_3\text{Ta}(\text{NH}_2)_2(\mathbf{3}-(\text{NH}_2)_2)$. $^{15}\text{NH}_3$ was used to obtain ^{15}N NMR data for $\mathbf{3}-(^{15}\text{NH}_2)_2$, whose nitrogens were observed as a triplet ($^1J_{15\text{NH}} = 71$ Hz) at δ -241.62 . This assignment was based on the ^{15}N NMR spectrum of $\mathbf{2}-(^{15}\text{NH}_2)$, obtained from $^{15}\text{NH}_3$ and $(\text{silox})_3\text{-Ta}(\mathbf{1})$, which resonated as a triplet of doublets at δ -229.37 ($^1J_{15\text{NH}} = 72$ Hz; $^3J_{15\text{NH}} = 5$ Hz). Attempts to generate equilibrium information pertaining to $\mathbf{1}=\text{NH} + \text{NH}_3 \rightleftharpoons \mathbf{3}-(\text{NH}_2)_2$ were fraught with complications that were attributed to complex equilibria,^{47,48} since different van't Hoff plots were obtained at different concentrations of NH_3 in benzene.²⁶ It was suspected that aggregation of NH_3 in solution (1, 3, and 10 equiv) was at the origin of the unusual behavior, and no further attempts were made. Another complication concerned the formation of $(\text{silox})_3\text{-Ta}(\text{H}_3\text{N})\text{Ta}=\text{NH}(\mathbf{1}=\text{NH}(\text{NH}_3))$, whose imide resonance was shifted to a broad signal at δ 5.67 in its ^1H NMR spectrum accompanied by a broad NH_3 resonance at δ 0.80. It is suspected that the original white solid is a combination of $\mathbf{3}-(\text{NH}_2)_2$ and $\mathbf{1}=\text{NH}(\text{NH}_3)$, but attempts to isolate either invariably led to ammonia loss. The ^{15}N NMR resonance at δ -60.60 (d, $^1J_{15\text{NH}} = 76$ Hz) observed in solution is attributed to $\mathbf{1}=(^{15}\text{NH})$, but it is conceivable it is slightly shifted because of equilibria with the ammonia adduct. A switch to THF- d_8 revealed four distinct species, but the spectral signature for the fourth species, presumably $\mathbf{1}=\text{NH}(\text{NH}_3)$, continued to be questioned for lack of a signal that could be attributed with confidence to the NH_3 group.

Alternative syntheses of Me_3Si -capped imido and phosphinidene derivatives are also given in Scheme 2. Deprotonation of $(\text{silox})_3\text{Ta}=\text{NH}(\mathbf{1}=\text{NH})$ with neopentylolithium afforded the lithio-imide, $(\text{silox})_3\text{Ta}=\text{NLi}(\mathbf{1}=\text{NLi})$, in 78% yield as colorless crystals from tetrahydrofuran (THF). Subsequent silylation with TMSCl afforded the imide, $(\text{silox})_3\text{Ta}=\text{NSiMe}_3(\mathbf{1}=\text{NTMS}, 50\%)$, again as colorless crystals. The corresponding route was also employed in the preparation of $(\text{silox})_3\text{Ta}=\text{PTMS}(\mathbf{1}=\text{PTMS})$, except that the phosphide anion^{34,49} was generated in situ and silylated to produce a small amount of the phosphinidene for the sake of ^{31}P NMR spectroscopy.

Table 2. UV–vis Absorptions for $(\text{silox})_3\text{Ta}=\text{EPh}$ ($\mathbf{1}=\text{EPh}$; $\text{E} = \text{N}, \text{P}, \text{As}$)

$\mathbf{1}=\text{NPh}$		$\mathbf{1}=\text{PPh}$		$\mathbf{1}=\text{AsPh}$	
λ (nm)	ϵ ($\text{M}^{-1} \text{cm}^{-1}$)	λ (nm)	ϵ ($\text{M}^{-1} \text{cm}^{-1}$)	λ (nm)	ϵ ($\text{M}^{-1} \text{cm}^{-1}$)
215 (sh)	17,000	230 (sh)	18,800	226	21,300
246	13,300	244 (sh)	15,600	249	18,900
280 (sh)	9,700	275	15,900	270	14,800
285	9,750	300 (sh)	5,160	312	2,100
294 (sh)	9,300	361	3,880	385	3,950
		434 (sh)	650	450 (sh)	910
		553	310	602	250

studies. Phosphinidene ^{31}P chemical shifts for $\mathbf{1}=\text{PH}$, $\mathbf{1}=\text{PPh}$, and $\mathbf{1}=\text{PTMS}$ were recorded at δ 276.33, 334.6, and 196.07, respectively. While most of the pnictidene complexes were isolated and subjected to elemental analysis ($\mathbf{1}=\text{ER}$, $\text{E} = \text{N}, \text{R} = \text{H}, \text{Me}, \text{SiMe}_3, \text{C}_6\text{H}_4\text{-}p\text{-X}$ ($\text{X} = \text{H}, \text{CF}_3$); $\text{E} = \text{P}, \text{R} = \text{H}, \text{Ph}$; $\text{ER} = \text{AsPh}$), the remaining complexes were generated in situ for subsequent studies.

3. UV–vis Spectra of Pnictidenes and Calculations.

UV–vis spectra of $(\text{silox})_3\text{Ta}=\text{EPh}$ ($\mathbf{1}=\text{EPh}$, $\text{E} = \text{N}, \text{P}, \text{As}$) were taken for comparison and their UV–vis absorptions are listed in Table 2. The higher energy absorption in each case is taken as an IL (intraligand) band because of comparisons with $\text{NaOSi}^t\text{Bu}_3$,⁶ and the mélange of bands in the 240–300 nm region are likely to be ligand-to-metal charge transfer (LMCT) bands of siloxide and pnictide character. In the cases of $\mathbf{1}=\text{PPh}$ and $\mathbf{1}=\text{AsPh}$, low energy bands of relatively lower intensity are seen, whereas no features are observed below ~ 330 nm for $\mathbf{1}=\text{NPh}$. Calculations reveal the highest occupied molecular orbitals (HOMOs) of $\mathbf{1}=\text{PPh}$ and $\mathbf{1}=\text{AsPh}$ to be essentially the $\text{Ta}=\text{E}$ π -bonding orbitals (Figure 1). The weak 553 nm ($\epsilon = 310 \text{ M}^{-1} \text{cm}^{-1}$) absorption of $\mathbf{1}=\text{PPh}$ is roughly 9–10,000 cm^{-1} lower⁵⁰ than the 361 nm absorption of considerably greater intensity ($\epsilon = 3,880 \text{ M}^{-1} \text{cm}^{-1}$), suggesting that these are the triplet and singlet bands corresponding to $\text{TaP}(\pi^b) \rightarrow \text{Ta}(d_{xz}(\pi^{nb}))$ LMCT. The modest intensities speak to the minimal overlap of the roughly orthogonal $d\pi_{yz}^b$ and d_{xz} orbitals. The shoulder at 300 nm and the low intensity band at 434 nm could be the $\text{TaP}(\pi^b) \rightarrow \text{Ta}(d_{yz}(\pi^*))$ LMCT and its accompanying triplet absorption, but computationally a $\text{P}(3s/3p_z)^{nb} \rightarrow \text{Ta}(d_{xz}(\pi^{nb}))$ LMCT would also be a reasonable fit from both energy and intensity standpoints. Strong parallels are observed in the low energy features of $\mathbf{1}=\text{AsPh}$, such as a weak 602 nm band ($\epsilon = 250 \text{ M}^{-1} \text{cm}^{-1}$) that is ~ 9 –10,000 cm^{-1} below a 385 nm band of modest intensity ($\epsilon = 3,950 \text{ M}^{-1} \text{cm}^{-1}$), which are consistent with triplet and singlet absorptions pertaining to the related to $\text{TaAs}(\pi^b) \rightarrow \text{Ta}(d_{xz}(\pi^{nb}))$ LMCT. The assignment of the remaining low energy transitions as $\text{TaP}(\pi^b) \rightarrow \text{Ta}(d_{yz}(\pi^{nb}))$ or $\text{As}(4s/4p_z)^{nb} \rightarrow \text{Ta}(d_{xz}(\pi^{nb}))$ singlet or triplet LMCT bands also parallels the phosphinidene case. Related interpretations of similar chalcogenide features have been made.^{51,52}

The surprising feature to the calculations of $(\text{silox})_3\text{-Ta}=\text{EPh}$ ($\mathbf{1}=\text{EPh}$, $\text{E} = \text{N}, \text{P}, \text{As}$) shown in Figure 1 are

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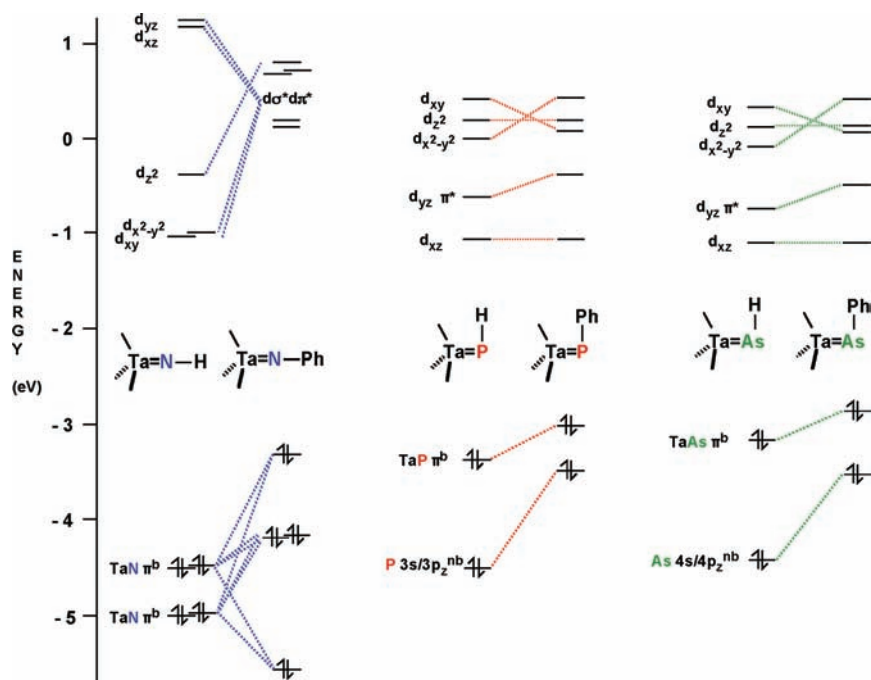


Figure 1. Truncated molecular orbital diagrams featuring pertinent Ta=E occupied levels and the ligand fields for (silox)₃Ta=ER (**1**=ER; E = N, P, As; R = H, Ph). In all cases, the orbital parentage has been tentatively interpreted from orbital pictures, but the low symmetry (*C_s*) cases were difficult, and the *d_{xz}/d_{yz}* set mixed with the *d_{x²-y²}/d_{xy}}* set in **1**=NPh to such an extent that no assignment could be made. Note that the two π -bonds in **1**=NH are spread among four orbitals because of σ/π -mixing, and are further split in **1**=NPh because of mixing with the phenyl orbitals.

the HOMOs of the phosphinidene and arsenidene, which are the Ta=E π -bonding orbitals. The “lone pairs” or “inert pairs” on P and As are expected to be relatively low in energy, and the π -bonds in the system are so weak that the Ta=E bonding orbitals are not driven below the inert pairs. Clearly the large 3s/3p and 4s/4p energy gaps for elemental phosphorus and arsenic, the origin of the so-called inert pair effect,^{53–56} are revealed relative to the π -interaction energies.

4. Pnictidene Structural Studies and Calculations.

Crystallographic studies of (silox)₃Ta=PPh (**1**=PPh, *d*(TaP) = 2.317(4) Å, \angle Ta=P–C = 110.2(4)°) and (silox)₃Ta=AsPh (**1**=AsPh, *d*(TaAs) = 2.428(2) Å, \angle Ta=As–C = 107.2(4)°) have been reported.⁵ A molecular view of (silox)₃Ta=NPh (**1**=NPh) and selected geometric parameters are given in Figure 2, and crystallographic details are given in Table 3. In contrast to the Ta–E–C angles of the phosphinidene and arsenidene, the TaNPh linkage is essentially linear, and the tantalum–nitrogen distance is standard for a triple bond (*d*(TaN) = 1.767(10) Å, \angle Ta=N–C = 172.4(8)°).⁵⁷ Utilization of the 2s orbital of N in the imide linkage is therefore apparent, constituting persuasive evidence for sp hybridization. High level quantum mechanical calculations of (silox)₃Ta=EPH (**1**'=EPH, primes indicated calculated values) are consistent with the reported structures: **1**'=NPh, *d*(TaN) = 1.81 Å, \angle Ta=N–C = 176.0°; **1**'=PPh, *d*(TaP) = 2.36 Å, \angle Ta=P–C = 112.7°; **1**'=AsPh, *d*(TaAs) = 2.47 Å, \angle Ta=As–C = 109.4°. Related calculations of (silox)₃Ta=EH

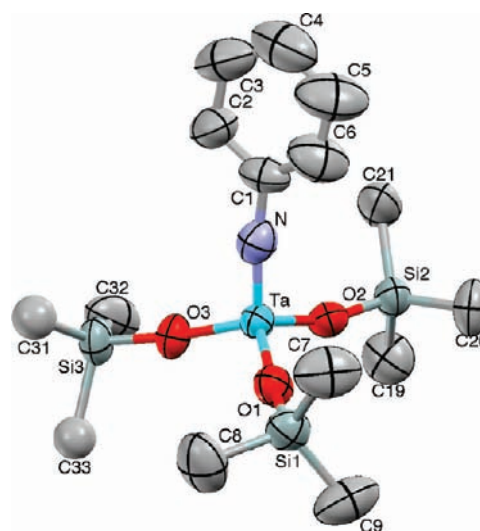


Figure 2. Molecular view of (silox)₃Ta=NPh (**1**=NPh) with the peripheral Me groups removed for clarity. Selected distances (Å) and angles (deg): TaN, 1.767(10); TaO1, 1.894(7); TaO2, 1.891(7); TaO3, 1.893(6); SiO(ave), 1.663(11); NC1, 1.425(15); TaNC1, 1.724(8); NTaO1, 1.080(4); NTaO2, 1.059(5); NTaO3, 1.096(3); O1TaO2, 1.114(3); O1TaO3, 1.110(3); O2TaO3, 1.108(3); TaO1Si1, 1.692(5); TaO2Si2, 1.647(5); TaO3Si3, 1.729(5).

(**1**'=EH) show the Ta=E–H angles to be close to 90° for E = P and As, therefore steric factors probably cause the Ta=E–C angles for **1**'=PPh and **1**'=AsPh to deviate from 90°: **1**'=NH, *d*(TaN) = 1.79 Å, \angle Ta=N–H = 179.4°; **1**'=PH, *d*(TaP) = 2.36 Å, \angle Ta=P–H = 87.6°; **1**'=AsH, *d*(TaAs) = 2.46 Å, \angle Ta=As–H = 87.0°.

5. 1,2-H₂-Elimination from (silox)₃HTaEHR. An assessment of the mechanism of 1,2-H₂-elimination from (silox)₃HTaEHR (**2**-EHR) was made via an evaluation of the rates as monitored by ¹H NMR spectroscopy in C₆D₆

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Table 3. Selected Crystallographic and Refinement Data for (silox)₃Ta=NPh (**1**=NPh), (silox)₃Ta=CPPh₃ (**1**=CPPh₃), [(silox)₃MeTa](μ-η²-N,N:η¹-C-NHNHCH₂CH₂-CH₂)[Ta(κ-O,C-OSi^tBu₂CMe₂CH₂(silox))₂] (**7**), and [(silox)₂ClTa](μ-N₂) (**8**-Cl)

	1 =NPh	1 =CPPh ₃	7	8 -Cl
formula	C ₄₂ H ₈₆ NO ₃ Si ₃ Ta	C ₅₅ H ₉₆ O ₃ PSi ₃ Ta	C ₇₆ H ₁₇₂ N ₂ O ₆ Si ₆ Ta ₂	C ₄₈ H ₁₀₈ N ₂ O ₄ Cl ₂ Si ₄ Ta
formula wt	918.34	1101.51	1740.60	1322.52
space group	<i>Pna</i> 2 ₁	<i>P</i> 2 ₁ / <i>n</i>	<i>P</i> $\bar{1}$	<i>P</i> 2 ₁ / <i>c</i>
<i>Z</i>	4	4	2	4
<i>a</i> , Å	18.114(4)	11.8368(2)	12.7517(6)	14.6904(6)
<i>b</i> , Å	13.436(3)	22.2055(4)	17.5127(9)	27.0349(10)
<i>c</i> , Å	20.650(4)	22.1197(4)	23.2899(11)	17.4888(7)
α, deg	90	90	103.755(2)	90
β, deg	90	90.1200(10)	90.992(2)	113.637(2)
γ, deg	90	90	110.627(2)	90
<i>V</i> , Å ³	5025.8(19)	5813.97(18)	4698.7(4)	6363.0(4)
ρ _{calc} , g·cm ⁻³	1.214	1.258	1.230	1.381
μ, mm ⁻¹	2.291	2.018	2.446	3.633
temp, K	293(2)	173(2)	173(2)	173(2)
λ (Å)	0.71073	0.71073	0.71073	0.71073
<i>R</i> indices [<i>I</i> > 2σ(<i>I</i>)] ^{a,b}	<i>R</i> ₁ = 0.0415 <i>wR</i> ₂ = 0.0976	<i>R</i> ₁ = 0.0404 <i>wR</i> ₂ = 0.0739	<i>R</i> ₁ = 0.0611 <i>wR</i> ₂ = 0.1429	<i>R</i> ₁ = 0.0566 <i>wR</i> ₂ = 0.1309
<i>R</i> indices (all data) ^{a,b}	<i>R</i> ₁ = 0.0733 <i>wR</i> ₂ = 0.1186	<i>R</i> ₁ = 0.0674 <i>wR</i> ₂ = 0.0824	<i>R</i> ₁ = 0.0912 <i>wR</i> ₂ = 0.1549	<i>R</i> ₁ = 0.0980 <i>wR</i> ₂ = 0.1561
GOF ^c	1.080	1.028	1.059	1.014

^a $R_1 = \sum ||F_o| - |F_c|| / \sum |F_o|$. ^b $wR_2 = [\sum w(|F_o| - |F_c|)^2 / \sum wF_o^2]^{1/2}$. ^c GOF (all data) = $[\sum w(|F_o| - |F_c|)^2 / (n - p)]^{1/2}$, *n* = number of independent reflections, *p* = number of parameters.

Table 4. Rate Constants^{a,b} for 1,2-H₂-Elimination from (silox)₃HTaEHR (**2**-EHR), with and without Added H₂ER and Radical Trap C₆H₈ Obtained by ¹H NMR Spectroscopic Monitoring

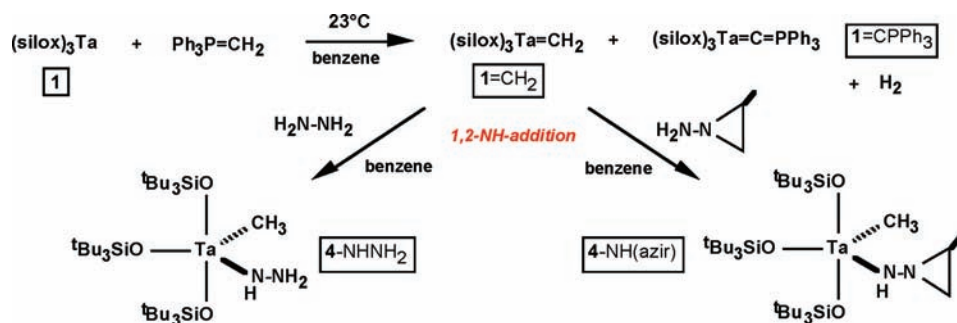
cmpd	[HTaEHR]	[H ₂ ER]	[C ₆ H ₈]	T(°C)	<i>k</i> (×10 ⁵ s ⁻¹)
(silox) ₃ HTaNH ₂ (2 -NH ₂) ^{c,d}	0.050			136.2(4)	166(5)
	0.056			119.0(4)	42.2(7)
	0.053			109.6(4)	18.8(4)
	0.053			99.1(4)	7.3(1)
	0.053			90.0(0)	2.73(2)
(silox) ₃ HTaNHMe (2 -NHMe)	0.029			24.8(4)	2.49(6)
	0.029	0.363		24.8(4)	29.0(10)
	0.038 ^e			24.8(4)	2.60(4)
(silox) ₃ HTaNPh (2 -NHPPh) ^f	0.060			24.8(4)	3.8(2)
	0.062	0.373		24.8(4)	3.8(2)
(silox) ₃ HTaNH(C ₆ H ₄ - <i>p</i> -Me) (2 -NHC ₆ H ₄ - <i>p</i> -Me)	0.058 ^g			24.8(4)	5.2(1)
(silox) ₃ HTaNH(C ₆ H ₄ - <i>p</i> -NMe ₂) (2 -NHC ₆ H ₄ - <i>p</i> -NMe ₂)	0.033 ^h			24.8(4)	22(1)
	0.053			24.8(4)	23.2(6)
(silox) ₃ DTaNH(C ₆ H ₄ - <i>p</i> -NMe ₂) (2D -NHC ₆ H ₄ - <i>p</i> -NMe ₂)	0.065 ^{h,i}			24.8(4)	20.0(2)
(silox) ₃ HTaNH(C ₆ H ₄ - <i>p</i> -CF ₃) (2 -NHC ₆ H ₄ - <i>p</i> -CF ₃)	0.062			24.8(4)	1.14(2)
(silox) ₃ DTaNH(C ₆ H ₄ - <i>p</i> -CF ₃) (2D -NHC ₆ H ₄ - <i>p</i> -CF ₃) ^j	0.062			24.8(4)	1.12(8)
(silox) ₃ HTaPH ₂ (2 -PH ₂) ^{k,l}	0.034			24.8(4)	1.06(1)
	0.056			24.8(4)	1.04(7)
	0.060		0.060	24.8(4)	1.07(2)
(silox) ₃ HTaPPh (2 -PPh) ^m	0.029 ^m			24.8(4)	0.26(2)
	0.064 ^j			24.8(4)	0.272(6)
	0.037 ^j	0.109		24.8(4)	0.28(3)
	0.054 ^m		0.054	24.8(4)	0.275(4)
(silox) ₃ HTaAsHPh (2 -AsHPh) ⁿ	0.053			-76(1)	140(10)
	0.037		0.037	-9.5(10)	33(1)
	0.040		0.398	-9.5(10)	33(1)

^a Determined from unweighted, non-linear, least-squares fitting of the exponential form of the rate expression (averaged where simultaneous runs were obtained); concentrations are in M. ^b **2**-NHR generated from **1** and RNH₂ unless otherwise indicated; monitored via disappearance of TaH. ^c **2**-NH₂ isolated prior to runs. ^d Δ*H*[‡] = 25.3(8) kcal/mol; Δ*S*[‡] = -10(2) eu. ^e Monitored via loss of NH(CH₃). ^f Generated from **2**-H and RNH₂. ^g Monitored by disappearance of ArCH₃. ^h Monitored via the appearance of imido N(CH₃)₂. ⁱ Generated from (silox)₃TaD₂ (**2D**-D) and H₂NC₆H₄-*p*-NMe₂; run under ~1 atm D₂. ^j Generated from (silox)₃TaD₂ (**2D**-D) and H₂NC₆H₄-*p*-CF₃; monitored by disappearance of NH. ^k Isolated prior to run. ^l Monitored via appearance of PH. ^m Monitored via disappearance of TaH. ⁿ **2**-EHR generated from **1** and H₂ER.

or C₇D₈. Table 4 lists rate constants obtained (typically an average of two or three runs) under various conditions. Rates were measured by generating **2**-EHR in situ when isolation of the pure compound was not possible. The eliminations were found to be first order in metal-hydride, and while the addition of H₂NMe was found to accelerate the elimination rate in the case of **2**-NHMe, the addition of PhNH₂ and PhPH₂ to **2**-NHPPh and **2**-PPh, respec-

tively, failed to induce a change in the observed rates. It may be that only small bases have the ability to catalytically assist in the H-transfer, perhaps via the formation of (silox)₃Ta(EHR)₂ and H₂, and subsequent elimination of H₂ER. EH₃ could not be added to **2**-EH₂, since **3**-(EH₂)₂ was found to form for E = N, As. Trace amounts of EH₂R were common in NMR tube experiments where hydrolysis of Ta-EHR bonds can typically occur during

Scheme 3



NH_3 . It appears that 1,2- H_2 -elimination from putative $(\text{silox})_3\text{HTaNH}(\text{NH}_2)$ (2-NHNH_2) is simply uncompetitive with respect to 1,2- HN -elimination, although $\text{N}-\text{N}$ oxidative addition followed by NH_3 loss cannot be excluded. A related approach, utilizing the addition of N -amino-2-methylaziridine,⁶⁶ was considered in the hope that propene loss and formation of $(\text{silox})_3\text{Ta}=\text{N}-\text{NH}_2$ ($1=\text{NNH}_2$) would permit a stepwise synthesis of 1_2-N_2 . Unfortunately, $(\text{silox})_3\text{Ta}=\text{NH}$ ($1=\text{NH}$) was the product, consistent with 2-methylaziridine loss analogous to that delineated for hydrazine (eq 2); byproducts (e.g., $(\text{silox})_3\text{Ta}(\eta\text{-C}_2\text{H}_3\text{Me})$) as in Scheme 2) were consistent with the reaction of 2-methylaziridine (shown as the initial product in eq 2) with 1.

2. Synthesis of $(\text{silox})_3\text{Ta}=\text{CH}_2$. In an attempt to obviate the presumed deleterious, swift 1,2- HN -elimination reaction, routes to $(\text{silox})_3\text{RTaNHNH}_2$ and $(\text{silox})_3\text{-RTaNH}(2\text{-Me-aziridine})$ were sought in the hope that 1,2- RH -elimination to give the desired imides would be controllable. As Scheme 3 illustrates, successful implementation of this route required the preparation of $(\text{silox})_3\text{Ta}=\text{CH}_2$ ($1=\text{CH}_2$),²⁶ which was accomplished via methylene transfer to $(\text{silox})_3\text{Ta}$ (**1**) from $\text{Ph}_3\text{P}=\text{CH}_2$,⁶⁷ related attempts with Me_3PCH_2 ,⁶⁸ which would yield the easily removable PMe_3 as a byproduct, afforded complex mixtures. As is typical for reactions with the Wittig reagent, removal of byproduct Ph_3P was difficult, but a 29% yield of $1=\text{CH}_2$ was obtained after multiple crystallizations from hexane. During the course of optimizing the synthesis, a curious byproduct was obtained that was devoid of hydrogens other than those of phenyl or silox. The compound, which could be isolated in 20% yield, possessed a single $^{31}\text{P}\{^1\text{H}\}$ NMR spectral shift at $\delta -4.26$ and a unique $^{13}\text{C}\{^1\text{H}\}$ NMR spectral resonance at $\delta 175.01$ that appeared as a doublet with $J_{\text{PC}} = 53$ Hz, consistent with the formulation $(\text{silox})_3\text{Ta}=\text{C}=\text{PPh}_3$ ($1=\text{CPPh}_3$). Its structure was confirmed by X-ray crystallography.

3. Structure of $(\text{silox})_3\text{Ta}=\text{C}=\text{PPh}_3$. Selected crystallographic details of $(\text{silox})_3\text{Ta}=\text{C}=\text{PPh}_3$ ($1=\text{CPPh}_3$) are given in Table 3, and Figure 4 illustrates its near C_3v core configuration, with the silox groups staggered relative to the phosphine phenyl substituents on the periphery. The $\text{Ta}-\text{C}-\text{P}$ linkage is virtually linear ($177.83(17)^\circ$), and the $d(\text{TaC})$ is quite short at $1.862(2)$ Å, signifying substantial

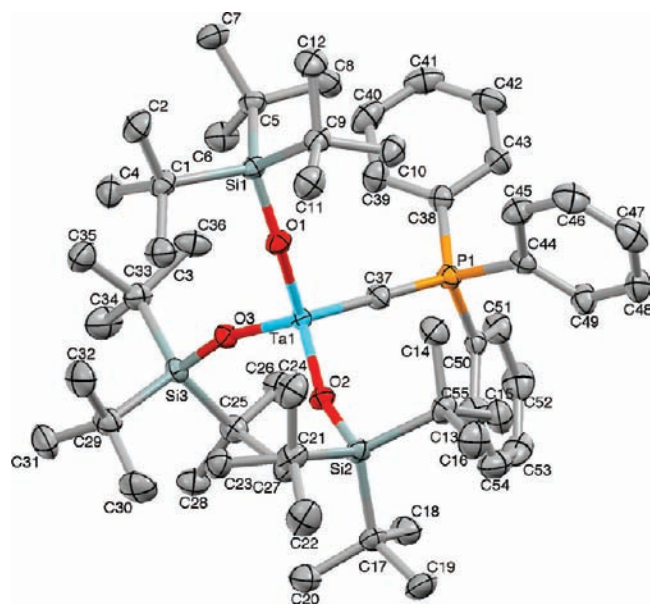


Figure 4. Molecular view of $(\text{silox})_3\text{Ta}=\text{C}=\text{PPh}_3$ ($1=\text{CPPh}_3$). Selected distances (Å) and angles (deg): $\text{Ta}-\text{C}37$, 1.862(2); $\text{Ta}-\text{O}1$, 1.9122(16); $\text{Ta}-\text{O}2$, 1.9072(15); $\text{Ta}-\text{O}3$, 1.9139; $\text{P}-\text{C}37$, 1.667(2); $\text{P}-\text{C}^{\text{Ph}}(\text{ave})$, 1.816(3); $\text{Si}-\text{O}(\text{ave})$, 1.655(5); $\text{Ta}-\text{C}37-\text{P}$, 177.83(17); $\text{O}1-\text{Ta}-\text{C}37$, 109.24(9); $\text{O}2-\text{Ta}-\text{C}37$, 108.66(9); $\text{O}3-\text{Ta}-\text{C}37$, 108.27(9); $\text{O}1-\text{Ta}-\text{O}2$, 110.45(7); $\text{O}1-\text{Ta}-\text{O}3$, 108.81(7); $\text{O}2-\text{Ta}-\text{O}3$, 111.36(7); $\text{C}37-\text{P}-\text{C}^{\text{Ph}}(\text{ave})$, 113.7(7); $\text{C}^{\text{Ph}}-\text{P}-\text{C}^{\text{Ph}}(\text{ave})$, 105.0(7); $\text{Ta}-\text{O}-\text{Si}(\text{ave})$, 163.6(19).

triple bond character.⁵⁹ The accompanying PC distance of $1.667(2)$ Å is slightly shorter than expected for a $\text{P}-\text{C}(\text{sp})$ single bond (1.78 Å), consistent with considerable double bond character; hence, the unit can be construed as having contributions from both $\text{Ta}=\text{C}=\text{P}$ and $\text{Ta}^{(-)}\equiv\text{C}-\text{P}^{(+)}$ resonance structures. Similar constructs have been useful in rationalizing phosphacarbene/ylne groups in related molecules.^{69–72} The remainder of the core has standard $(\text{silox})_3\text{Ta}^{\text{V}}$ parameters that are given in the caption to Figure 3.

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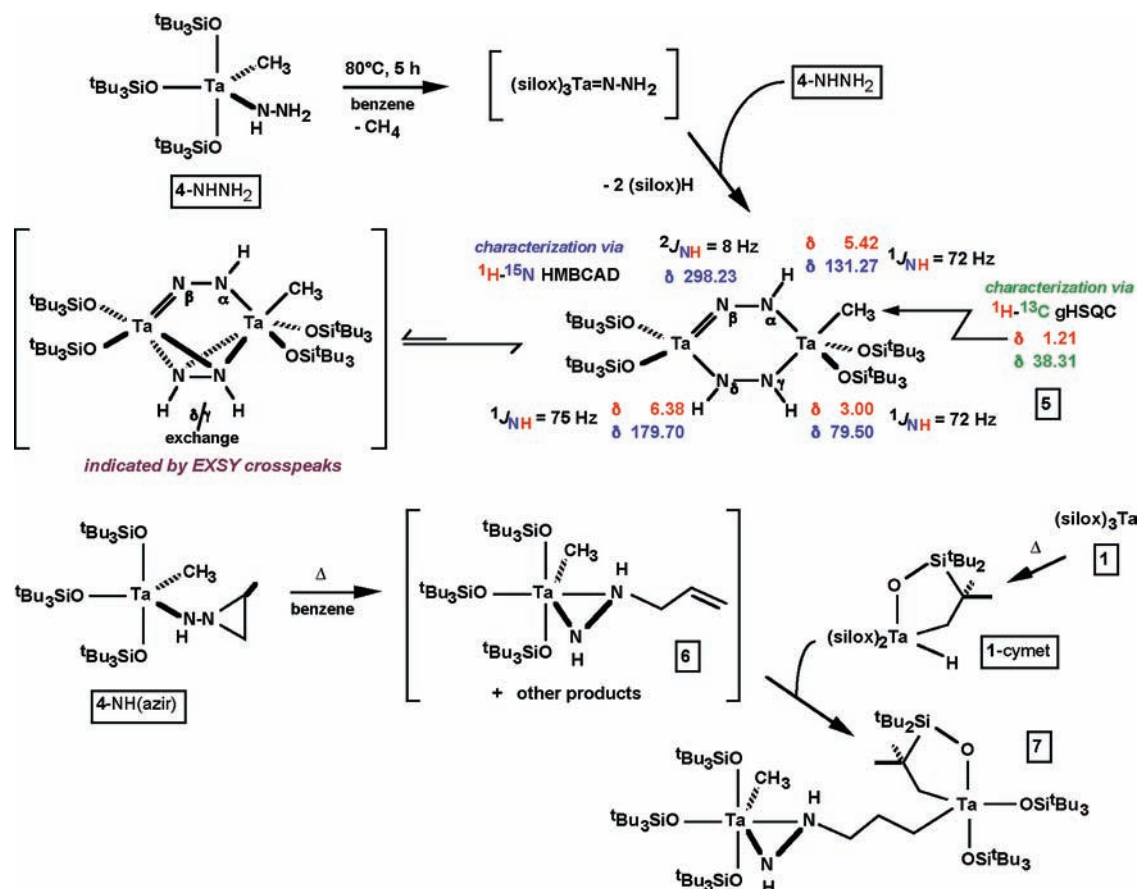
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Scheme 4



4. 1,2-RH-Elimination Attempts. As Scheme 3 reveals, the addition of hydrazine to $(\text{silox})_3\text{Ta}=\text{CH}_2$ ($\mathbf{1}=\text{CH}_2$) afforded $(\text{silox})_3\text{MeTaNHNH}_2$ ($\mathbf{4-NHNH}_2$) in 70% yield. Although $\mathbf{4-NHNH}_2$ was modestly thermally sensitive, NMR spectral characterization was readily accomplished using $^1\text{H}-^{15}\text{N}$ HSQCAD correlation spectroscopy.⁷³ In addition to the silox and Me proton resonances, different nitrogens at δ 64.80 and δ 148.47 ($J_{\text{NH}} = 78\text{ Hz}$) were observed to correspond with proton signals at δ 3.63 and δ 4.11, and these were assigned to the NH_2 and NH groups, respectively. Similarly, treatment of $\mathbf{1}=\text{CH}_2$ with amino-2-methylaziridine provided $(\text{silox})_3\text{MeTaNH}(-^c\text{NCHMeCH}_2)$ ($\mathbf{4-NH(azir)}$) in 60% yield as a waxy solid. The aziridine nitrogen was indirectly found at δ 81.99 (^{15}N), and the accompanying NH was located at δ 244.46 with $J_{\text{NH}} = 72\text{ Hz}$. Subsequent thermalyses of the methyl-hydrazides generated derivatives that proved to be more complicated than those expected from simple 1,2-MeH-eliminations.

Thermal degradation of $(\text{silox})_3\text{MeTaNHNH}_2$ ($\mathbf{4-NH-NH}_2$) led to the loss of MeH , but evidence of $(\text{silox})\text{H}$ was also obtained, and NMR spectroscopic data were more consistent with a dinuclear species with four inequivalent silox groups. Attempts to rid the solutions of byproduct $(\text{silox})\text{H}$ invariably led to further degradation, thus it is conceivable that the silanol is weakly hydrogen-bonded to the product, especially when solutions are concentrated. Successful implementation of $^1\text{H}-^{15}\text{N}$ HMBCAD

NMR correlation spectroscopy enabled the connectivity of the N-containing bridges to be ascertained, and an elusive Me group, partially masked by the silox resonances, was brought to light by $^1\text{H}-^{13}\text{C}$ gHSQC NMR correlation spectroscopy.⁷⁴ The product is thought to be $[(\text{silox})_2\text{TaMe}(\mu\text{-N}_\alpha\text{HN}_\beta)(\mu\text{-N}_\gamma\text{HN}_\delta)]$ ($\mathbf{5}$), which is portrayed in Scheme 4. Attempts to crystallize $\mathbf{5}$ in the presence of $(\text{silox})\text{H}$ led to waxy solids of reasonable purity, but these were not amenable to X-ray diffraction analysis. As such, the hapticity and orientation of bridges and silox groups are undetermined. Nonetheless, the NMR spectroscopy clearly shows a diamido $\mu\text{-NH-NH}$ bridge accompanied by an imido-amide $\mu\text{-NH-N}$ unit. The shift of one amide (N_γH , δ 79.50) is substantially lower than the remaining two (N_αH , δ 131.27; N_δH , δ 179.70), and may be indicative of a strong σ -donor (i.e., Me) opposite it in the complex. The imido nitrogen is observed more downfield at δ 298.23. EXSY correlation spectroscopy revealed exchange between N_γH and N_δH , thus it is possible that this bridge exists as an $\eta^2, \eta^{2/1}\text{-NH}$, NH unit^{12,75,76} or this configuration is at least energetically feasible to permit rapid exchange, as Scheme 4 shows.

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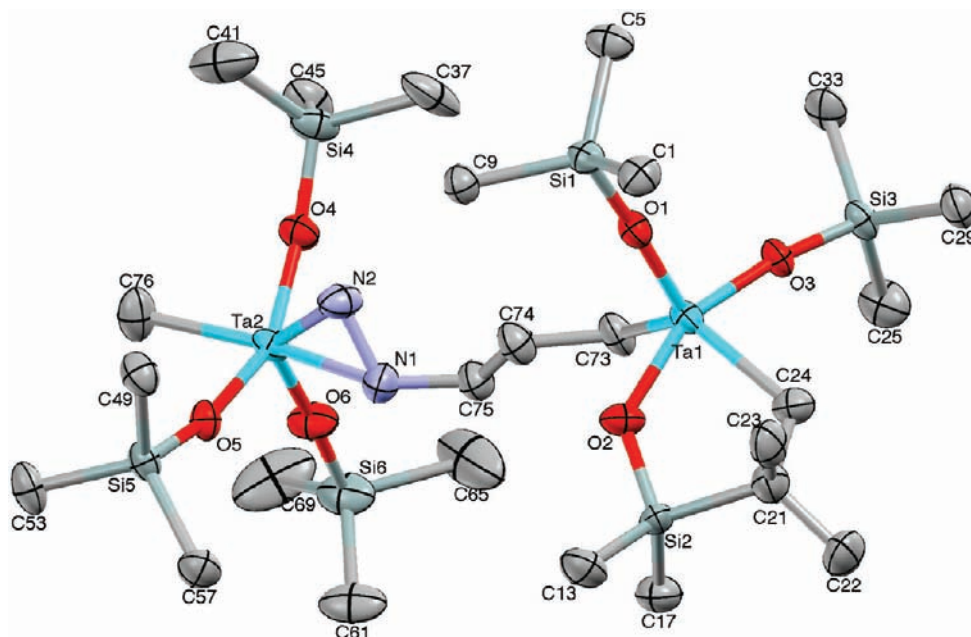


Figure 5. Molecular view of $[(\text{silox})_3\text{MeTa}](\mu\text{-}\eta^2\text{-}N,N:\eta^1\text{-}C\text{-NHNHCH}_2\text{CH}_2\text{CH}_2)[\text{Ta}(\kappa\text{-O,C-OSi}^b\text{Bu}_2\text{CMe}_2\text{CH}_2)(\text{silox})_2]$ (**7**) with the methyls of the ^tBu groups removed for clarity. Selected distances (Å) and angles (deg): Ta1–O1, 1.862(8); Ta1–O2, 1.932(5); Ta1–O3, 1.905(5); Ta1–C24, 2.217(9); Ta1–C73, 2.133(8); C73–C74, 1.523(11); C74–C75, 1.481(12); N1–C75, 1.441(11); N1–N2, 1.384(11); Ta2–N1, 2.283(7); Ta2–N2, 2.008(7); Ta2–C76, 2.211(9); Ta2–O4, 1.915(5); Ta2–O5, 1.951(5); Ta2–O6, 1.896(6); Si–O(ave), 1.662(11); O1–Ta1–O2, 100.4(2); O1–Ta1–O3, 104.8(2); O2–Ta1–O3, 88.3(3); O1–Ta1–C24, 123.8(3); O1–Ta1–C73, 111.0(3); O2–Ta1–C24, 77.0(3); O2–Ta1–C73, 86.0(3); O3–Ta1–C24, 86.1(3); O3–Ta1–C73, 88.3(3); O4–Ta2–N1, 108.7(3); O4–Ta2–N2, 85.0(3); O4–Ta2–O5, 152.7(2); O4–Ta2–O6, 100.4(2); O4–Ta2–C76, 83.1(3); O5–Ta2–N1, 79.4(2); O5–Ta2–N2, 87.4(3); O5–Ta2–O6, 105.3(2); O5–Ta2–C76, 82.8(3); O6–Ta2–N1, 93.1(3); O6–Ta2–N2, 126.0(3); O6–Ta2–C76, 101.3(3); N1–Ta2–N2, 36.9(3); N1–Ta2–C76, 159.6(3); N2–Ta2–C76, 132.6(3); Ta1–C73–C74, 118.2(6); C73–C74–C75, 114.5(8); C74–C75–N1, 115.8(8); N2–N1–C75, 114.7(7); N1–N2–Ta2, 82.4(4); Ta2–N1–C75, 136.8(6); Ta2–N1–N2, 60.7(4); Ta1–O2–Si2, 131.5(4); O2–Si2–C21, 96.4(4); Si2–C21–C24, 102.7(6); Ta1–C24–C21, 119.7(6).

The failure to generate $(\text{silox})_3\text{Ta}=\text{NNH}_2$ as a stable entity^{77,78} prompted a turn toward the aziridine chemistry, and $(\text{silox})_3\text{MeTaNH}(\text{-}^c\text{NCHMeCH}_2)$ (**4-NH**(azir)) was treated with 2 equiv of $(\text{silox})_3\text{Ta}$ (**1**) with the hope of producing $(\text{silox})_3\text{MeTaNH}=\text{N}=\text{Ta}(\text{silox})_3$ and $(\text{silox})_3\text{Ta}(\eta\text{-C}_3\text{H}_6)$. Unfortunately, no deoxygenation of the aziridine was noted at 23 °C, and thermolysis **4-NH**(azir) under varied conditions could not be accomplished without significant cyclometalation of **1** to give $(\text{silox})_2\text{HTa}(\kappa\text{-O,C-OSi}^b\text{Bu}_2\text{CMe}_2\text{CH}_2)$ (**1-cymet**).^{6,15} During the thermolysis, several products were noted, including those containing olefinic residues, consistent with ring-opening of the aziridine⁷⁹ to $(\text{silox})_3\text{MeTaNHNNH-CH}_2\text{CH}=\text{CH}_2$ (**6**). Yellow crystals that proved to be nearly insoluble in common hydrocarbons were observed to form during this process, and X-ray crystallographic studies revealed this material to be $[(\text{silox})_3\text{MeTa}](\mu\text{-}\eta^2\text{-}N,N:\eta^1\text{-}C\text{-NHNHCH}_2\text{CH}_2\text{CH}_2)[\text{Ta}(\kappa\text{-O,C-OSi}^b\text{Bu}_2\text{CMe}_2\text{CH}_2)(\text{silox})_2]$ (**7**). As Scheme 4 illustrates, **7** is best thought of arising from insertion of the olefin of **6** into the Ta–H bond of **1-cymet**.¹⁵

The treatment of $(\text{silox})_3\text{Ta}=\text{CH}_2$ (**1=CH}_2**) with diimines was also considered in view of Schrock's preparation of $[(\text{THF})_2\text{Cl}_3\text{Ta}]_2(\mu\text{-N}_2)$.⁹ Unfortunately, stable diimines such as $\text{PhCH}=\text{N}=\text{N}=\text{CHPh}$ failed to react with the methylene derivative, presumably because of deleterious steric interactions in the putative 4-membered ring intermediate.

6. Structure of $[(\text{silox})_3\text{MeTa}](\mu\text{-}\eta^2\text{-}N,N:\eta^1\text{-}C\text{-NHNHCH}_2\text{CH}_2\text{CH}_2)[\text{Ta}(\kappa\text{-O,C-OSi}^b\text{Bu}_2\text{CMe}_2\text{CH}_2)(\text{silox})_2]$ (7**).** With solubility precluding a definitive NMR analysis of $[(\text{silox})_3\text{MeTa}](\mu\text{-}\eta^2\text{-}N,N:\eta^1\text{-}C\text{-NHNHCH}_2\text{CH}_2\text{CH}_2)[\text{Ta}(\kappa\text{-O,C-OSi}^b\text{Bu}_2\text{CMe}_2\text{CH}_2)(\text{silox})_2]$ (**7**), an X-ray crystallographic study (Table 3) was conducted on the aforementioned yellow crystals. Despite one badly disordered set of silox ^tBu groups (on Si6), the study yielded a reasonable model that illustrates the ring-opened aziridine product as illustrated by Figure 5. The cyclometalated center is fairly well described as a pseudo-tbp geometry with the two less-electronegative alkyl substituents in their expected equatorial positions.^{3,80} The chain linkage, Ta1–C73, is 2.133(8) Å, and the cyclometalated $d(\text{Ta1}-\text{C24})$ is 2.217(9) Å, and both are separated by 124.5(4)°. The remaining pseudo-equatorial position is occupied by O1, which is 1.862(5) Å from the tantalum, and 111.0(3)° and 123.8(3)° from C73 and C24, respectively. Siloxide oxygens O2 and O3 are 1.932(5) and 1.905(5) Å from the tantalum, respectively, which are slightly longer than O1 and consistent with their pseudoaxial positions. The bite angle of the cyclometalated silox is 77.0(3)°, and its oxygen, O2, is 100.4(2)° and 86.0(3)° from O1 and C73, respectively.

One can similarly assess the hydrazide-methyl core, taking TaO4 (1.915(5) Å) and TaO5 (1.951(5) Å) as pseudoaxial siloxides ($\angle\text{O4-Ta2-O5} = 152.7(2)^\circ$), and TaO6 (1.896(6) Å) as a pseudo-equatorial siloxide, 100.4(2)° and 105.3(2)° from O4 and O5, respectively.

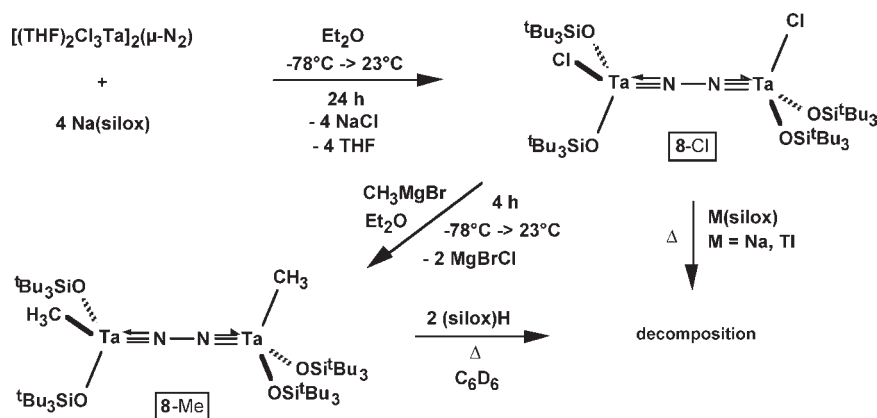
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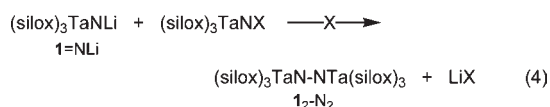
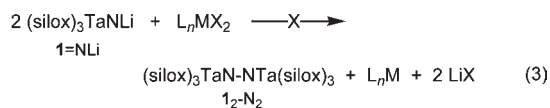
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Scheme 5



This assessment treats the NH-NHR ligand as occupying one site. The methyl is pseudo-equatorial (2.211(9) Å) along with the amide of the hydrazide ($d(\text{Ta}_2\text{-N}_2) = 2.008(7)$ Å; $d(\text{N}_1\text{-N}_2) = 1.384(11)$ Å), and the trans influence of the donor nitrogen, N2 (2.283(7), $\angle\text{C76-Ta}_2\text{-N}_1 = 159.6(3)^\circ$), helps slightly weaken the methyl interaction. The remaining core distances and angles are listed in the Figure 5 caption. Distances and angles pertaining to the bridge are consistent with a $(\text{CH}_2)\text{-NHNH}$ linkage derived from ring-opening of the aziridine and trapping of the subsequent olefin by the hydride of the 1-cymet, just as Scheme 4 portrays.

7. Attempted Oxidative Coupling of $(\text{silox})_3\text{TaNLi}$. Conceptually, oxidation of the nitride anion, $(\text{silox})_3\text{-TaNLi}$ ($\mathbf{1}=\text{NLi}$), can lead to the desired N_2 -complex, but one electron oxidants invariably generated $(\text{silox})_3\text{-Ta}=\text{NH}$ ($\mathbf{1}=\text{NH}$), presumably via H-atom abstraction by an incipient $(\text{silox})_3\text{TaN}\cdot$ species. A more probable strategy would rely on N-N oxidative coupling induced by reductive elimination from $\text{L}_n\text{M}(\text{NTa}(\text{silox})_3)_2$, as eq 3 implicates. Utilization of $(\text{Ph}_3\text{P})_2\text{PtCl}_2$, $(\text{Et}_3\text{P})_2\text{PdCl}_2$, $(\text{DME})\text{NiCl}_2$, and other related complexes failed to induce N-N bond formation, and no evidence for M-N intermediates was found. Only for Cp_2TiCl_2 was NMR spectroscopic evidence for $\text{Cp}_2\text{CITi-N}=\text{Ta}(\text{silox})_3$ obtained, and its thermolysis yielded $\mathbf{1}=\text{NH}$ and Cp_2TiCl . Attempts to oxidatively add the NH bond of $\mathbf{1}=\text{NH}$ to $(\text{Ph}_3\text{P})_4\text{Pd}$ and trap the product hydride with olefin also failed to provide any evidence of $(\text{Ph}_3\text{P})_n\text{PdH}(\text{N}=\text{Ta}(\text{silox})_3)$.



A potentially simpler route would require conversion of the anionic nitride into an electrophile of the type $(\text{silox})_3\text{Ta}=\text{NX}$ ($\mathbf{1}=\text{NX}$), which would then be susceptible

to attack by nitride $(\text{silox})_3\text{TaNLi}$ ($\mathbf{1}=\text{NLi}$), as indicated by eq 4.⁸¹ Unfortunately, attempts to halogenate the nitride anion with Br_2 , I_2 , or NBS failed to elicit any tractable material under a variety of conditions. Since $(\text{silox})_3\text{Ta}=\text{NH}$ was a common product in virtually all of the attempts, it appears that Ie^- reactivity is at the core of the problem.

Compromise, $[(\text{silox})_2\text{CITa}](\mu\text{-N}_2)$. **1. Synthesis of $[(\text{silox})_2\text{CITa}](\mu\text{-N}_2)$.** Although a number of variants on the preceding methods are plausible, Schrock's diimide complexes, in particular $[(\text{THF})_2\text{Cl}_3\text{Ta}]_2(\mu\text{-N}_2)$,⁹ were considered as starting materials for metathesis reactions to attain some measure of success in a reasonable period of time. As Scheme 5 illustrates, treatment of the chlorodiimide derivative with 4 equiv of $\text{Na}(\text{silox})$ afforded $[(\text{silox})_2\text{CITa}](\mu\text{-N}_2)$ (**8-Cl**) in 56% yield. Further metathesis with $\text{Na}(\text{silox})$ or $\text{Tl}(\text{silox})$ on **8-Cl** failed to provide the desired $[(\text{silox})_3\text{Ta}]_2(\mu\text{-N}_2)$ ($\mathbf{1}_2\text{-N}_2$) complex, and thermolysis ultimately led to degradation. Prior work⁴ suggested that smaller siloxides/alkoxides would not support a 3-coordinate tantalum center. The chloride was converted to the Me derivative, $[(\text{silox})_2\text{MeTa}](\mu\text{-N}_2)$ (**8-Me**, 70%) with MeMgBr , but subsequent attempts at silanolysis reactions with $(\text{silox})\text{H}$ did not generate MeH , and heating again led to decomposition. NMR spectra of **8-Cl** and **8-Me** revealed one type of silox, consistent with either C_{2h} symmetry and/or free rotation about the bridge.

2. Structure of $[(\text{silox})_2\text{CITa}](\mu\text{-N}_2)$. Table 3 lists pertinent crystallographic information pertaining to the structure of $[(\text{silox})_2\text{CITa}]_2(\mu\text{-N}_2)$ (**8-Cl**), and Figure 4 provides two views of the molecule. Unfortunately, while the data confirm the C_2 structure, the electron density between the two heavy tantalum atoms would not yield a reliable solution; using an isotropic model for the bridging nitrogens, TaN distances of 1.780(7) and 1.739(8) Å, and a $d(\text{NN})$ of 1.355(11) Å were found, but these should not be construed as accurately portraying the bridge. Nonetheless, the conformation of **8-Cl** was confirmed, including a pseudo-eclipsed placement of the core substituents relative to the adjacent tantalum, as previously seen in $[(\text{silox})_2\text{XM}]_2$ systems.⁸² This arrangement essentially minimizes steric interactions across the bridge by

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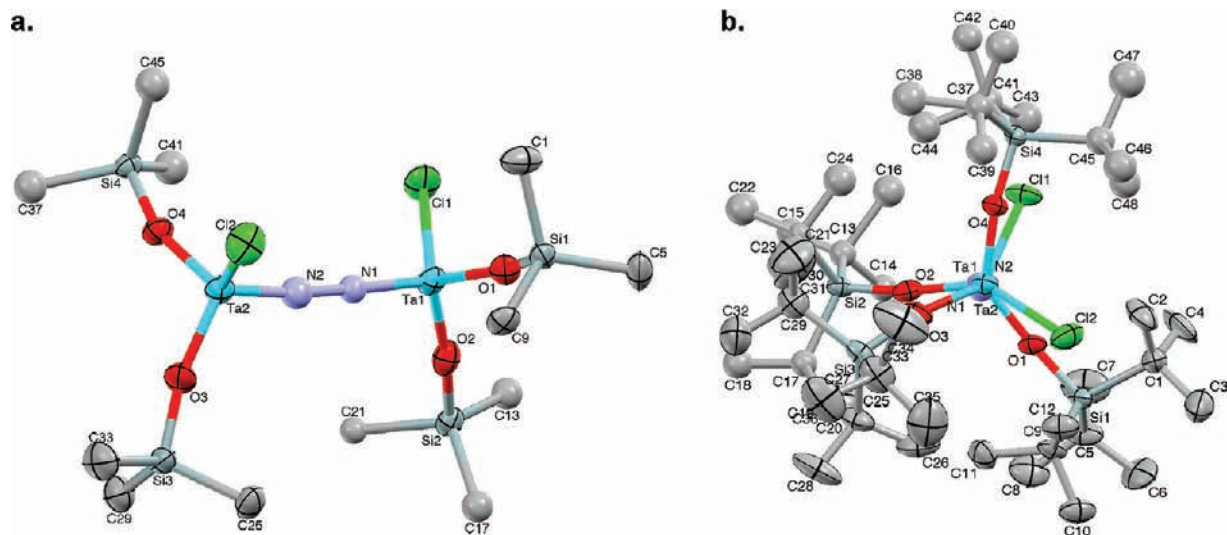


Figure 6. (a) Molecular view of $[(\text{silox})_2\text{ClTa}]_2(\mu\text{-N}_2)$ (**8-Cl**) with peripheral Me groups removed for clarity; note the N atoms refined isotropically; see text for comments pertaining to the TaNNTa bridge. (b) End on view of **8-Cl** revealing typical C_2 symmetry of dimeric species. Selected distances (Å) and angles (deg): Ta1–O1, 1.869(6); Ta1–O2, 1.871(7); Ta2–O3, 1.868(7); Ta2–O4, 1.854(6); Ta1–Cl1, 2.312(2); Ta2–Cl2, 2.307(3); Si–O(ave), 1.652(5); N1–Ta1–O1, 109.2(3); N1–Ta1–O2, 111.4(3); N1–Ta1–Cl1, 107.5(3); O1–Ta1–O2, 110.7(3); O1–Ta1–Cl1, 109.5(2); O2–Ta1–Cl1, 108.5(2); N2–Ta2–O3, 111.6(4); N2–Ta2–O4, 108.5(3); N2–Ta2–Cl2, 107.9(3); O3–Ta2–O4, 109.6(3); O3–Ta2–Cl2, 109.8(2); O4–Ta2–Cl2, 109.5(2).

trading two minimal silox/Cl interactions for one significant silox/silox interaction. Only one type of silox group is observed in NMR spectra of **8-Cl** or $[(\text{silox})_2\text{-MeTa}]_2(\mu\text{-N}_2)$ (**8-Me**), presumably because of rapid rotation about the TaNNTa linkage. Distances and angles of the pseudo-tetrahedral cores ($109.5(13)^\circ$) are listed in Figure 6.

Discussion

Pnictogen Chemistry. 1. H₂ER Oxidative Addition to (silox)₃Ta. A previous study that employed substituents on $\text{H}_2\text{N}(\text{C}_6\text{H}_4\text{-X})$ (X, k_{rel} : 3-CF₃, 0.30(7); 3-F, 0.63(12); 4-F, 0.89(17); H, 1; 4-Ph, 0.93(17); 4-Me, 1.2(2); 4-OMe, 1.9(2); 4-NMe₂, 2.5(2)) examined the relative rates (aniline at $k_{\text{rel}} = 1$) of NH oxidative addition via competition experiments.³ As the basicity of the aniline increased ($\rho = -0.69$, $R = 0.93$), the relative rate of oxidative addition increased, consistent with nucleophilic attack by the amine at a vacant d_{xz}/d_{yz} orbital preceding oxidative addition. The correlation was modest, and separate Hammett parameters were indicative of slightly greater resonance than inductive contributions. While complementary studies of REH₂ oxidative addition to (silox)₃Ta (**1**) were not examined, they qualitatively correlated with basicity of the substrate modified by steric factors.

2. 1,2-Dihydrogen-Elimination from (silox)HTaNHR. The 1,2-dihydrogen elimination data listed in Table 4 represent a limited, but reasonably interpretable set with regard to (silox)₃HTaNHR (**2-NHR**). The activation parameters ($\Delta H^\ddagger = 25.3(8)$ and $\Delta S^\ddagger = -10(2)$ eu) for H₂ elimination from (silox)₃HTaNH₂ (**2-NH₂**) are fully consistent with previous 1,2-RH-eliminations of RH (silox)_{*n*}(^tBu₃SiNH)_{3-*n*}MR species, whose ΔS^\ddagger values were typically -12 to -13 eu.^{59–61} Since H does not need to reorganize prior to elimination, in contrast to R, the slightly less negative activation entropy observed is expected. Loss of RH/D from (silox)₂(^tBu₃SiNH/D)TiR results in KIEs ranging from ~ 7 – 15 at 24.8 °C,⁵⁹ and

while the related experiments were not conducted in this study, the absence of a significant isotope effect in 1,2-HD-elimination from (silox)₃DTaNH(C₆H₄-*p*-X) (**2D-NHC₆H₄-*p*-X**; X = CF₃, NMe₂) suggests that the activation is primarily one involving N–H breakage.

The Hammett study of 1,2-H₂-elimination from (silox)₃-HTa(NHC₆H₄-*p*-X) (**2-NHC₆H₄-*p*-X**) in Figure 3, the relative rates of dihydrogen loss from (silox)₃HTa(NHR) (**2-NHR**; R = Ph (3420), Me (2340), and H (1)), and the modest correlation with pK_a 's of the corresponding amines support the contention that positive charge on N develops in the transition state. In **2-NHR**, N→Ta π -bonding also justifies the acidic character of the NH hydrogen, while the electropositive nature of tantalum relative to hydrogen ensures that the TaH bond is Ta(δ^+)–H(δ^-). Two additional factors suggest that N–H bond breaking, viewed as having a (NH) δ^+ component and hence having some proton transfer character, is the major bond activation. First, virtually no isotope effect is observed in the hydride position, as assessed via $k(\text{2-NHC}_6\text{H}_4\text{-}i\text{-X})/k(\text{2D-NHC}_6\text{H}_4\text{-}i\text{-X})$ (X = NMe₂, KIE = 1.10(6); CF₃, KIE = 1.02(10)). Second, a modest inverse correlation of rate versus $D(\text{H-NHR})$ is observed, where the weaker the bond, the faster the 1,2-H₂-elimination.

3. 1,2-H₂-Elimination from (silox)₃HTaEHR. While dihydrogen elimination from the nitrogen-containing species was reasonably explained, and consistent with the widely held view of charge distribution in early metal complexes, inclusion of phosphorus and arsenic muddies the waters. To reiterate, if ΔS^\ddagger for H₂ elimination from **2-AsHPh** is assumed to be -10 eu, the relative 1,2-H₂-elimination rates from (silox)₃HTaEHPH (**2-EHPH**) at 24.8 °C are As (5800) > N (14) > P (1). The aforementioned thermodynamic $D(\text{E-H})$ or $\Delta H_f^\circ(\text{EH}_3)$ correlations predict a standard periodic trend; As should be faster than P, followed by N. Given the relative paucity of information on the actual thermodynamics of such events, high level quantum calculations were conducted.

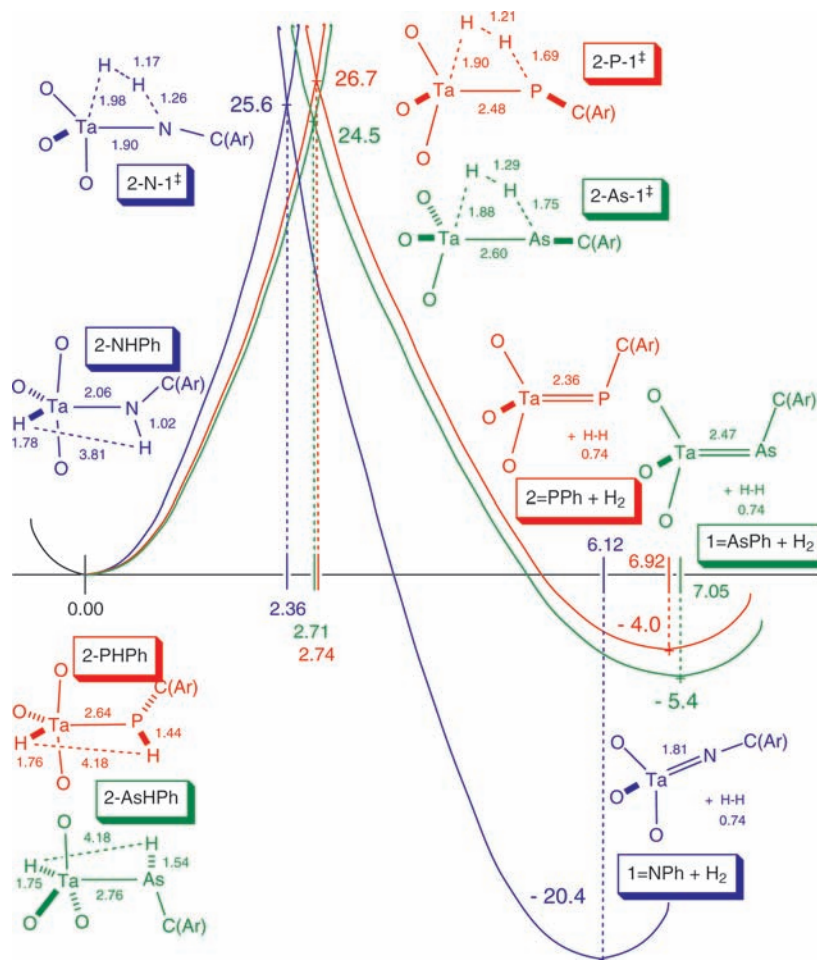


Figure 7. Calculated energies (y -axis, kcal/mol) of 1,2- H_2 -elimination from $(\text{silox})_3\text{HTaEHPH}$ (**2-EHPH**; $\text{E} = \text{N}, \text{P}, \text{As}$) to give $(\text{silox})_3\text{Ta}=\text{EPh}$ (**1=EPh**, $\text{E} = \text{N}, \text{P}, \text{As}$) + H_2 via transition states $[\mathbf{2-E-1}]^\ddagger$ ($\text{E} = \text{N}, \text{P}, \text{As}$). The reaction coordinate (x -axis) is described in Table 5 (RC(x)).

4. Calculations on 1,2- H_2 -Elimination from $(\text{silox})_3\text{-HTaEHR}$. Figure 7 reveals the geometries and relative energies of $(\text{silox})_3\text{HTaEHPH}$ (**2-EHPH**; $\text{E} = \text{N}, \text{P}, \text{As}$), the products $(\text{silox})_3\text{Ta}=\text{EPh}$ (**1=EPh**, $\text{E} = \text{N}, \text{P}, \text{As}$) + H_2 , and intervening transition states $[\mathbf{2-E-1}]^\ddagger$ ($\text{E} = \text{N}, \text{P}, \text{As}$). Remarkably, the calculations (24.8°C : $\Delta G^\ddagger(\text{P})_{\text{cald}} = 26.7$ kcal/mol, $\Delta G^\ddagger(\text{N})_{\text{cald}} = 25.6$ kcal/mol, $\Delta G^\ddagger(\text{As})_{\text{cald}} = 24.5$ kcal/mol) parallel the general experimental trend although the activation free energies are somewhat different (24.8°C : $\Delta G^\ddagger(\text{P}) = 25.0$ kcal/mol, $\Delta G^\ddagger(\text{N}) = 23.5$ kcal/mol, $\Delta G^\ddagger(\text{As}) = 19.9$ kcal/mol). One straightforward explanation for the lack of a periodic trend concerns the thermodynamics for 1,2- H_2 -elimination, which is calculated to be 20.4 kcal/mol exoergic for N, but only 4.0 and 5.4 kcal/mol favorable for P and As, respectively. The greater favorable free energy for elimination of H_2 from the amide-hydride suggests a lower transition state than the P and As cases that have only a modest driving force.

The argument that the more favorable free energy change for N renders a lower transition state also suggests that $[\mathbf{2-N-1}]^\ddagger$ should occur *earlier* in the reaction coordinate. Table 5 provides a view of the reaction coordinate parametrized solely via bond distance changes: (1) as a change in total bond lengths starting from $(\text{silox})_3\text{HTaEHPH}$ (**2-EHPH**; $\text{E} = \text{N}, \text{P}, \text{As}$; $x = 0$), given as RC(x); (2) as a fractional change in a reaction coordinate in bond

length normalized to 1.00 with the pnictide-hydrides **2-EHPH** at 0.00. Somewhat surprisingly, the RC for all of the pnictogens is similar, where the TS is about 40% of the way to the product pnictidene (N, 0.39; P, 0.40; As, 0.38), despite a significant disparity in the sum of bond distance changes: N, 6.12 Å; P, 6.92 Å; As, 7.05 Å. One way to rationalize these features is to accept the premise that the $\text{Ta}=\text{N}$ bond is likely to have a substantially greater force constant than the remaining $\text{Ta}=\text{P}$ and $\text{Ta}=\text{As}$ bonds, thus affording a “steeper sided” free energy surface (Figure 7) for the product **1=NPh** relative to **1=PPh** and **1=AsPh** (likely to be also true for **2-EHPH**). A steeper free energy surface would counteract the expectation of an earlier transition state for N because of the more favorable ΔG° , while allowing the overall reaction coordinate to be more compressed. This can be observed in the changes in the pnictide and pnictidene distances. From **2-EHPH** to $[\mathbf{2-E-1}]^\ddagger$, the $\text{Ta}-\text{E}$ distances all change by -0.16 Å, but this is a far greater percentage change for the amide (7.8% relative to $\sim 6\%$ for P and As), while the percent change in $d(\text{TaE})$ from $[\mathbf{2-E-1}]^\ddagger$ to **1=EPh** are all roughly 5% for all three cases. There is a corresponding greater percent change in the $d(\text{EH})$ for $\text{N} > \text{P} > \text{As}$ that complements the $d(\text{TaE})$ explanation. Essentially, the greater bond strengths in the N species, especially that of the imido product, render the transition state later than it would be based solely on the standard free energy change for

Table 5. Calculated Reaction Coordinate^a Changes from 2-EHPH to [2-E-1][†] to 1=EPH

	<i>d</i> (TaH)	<i>d</i> (TaE)	<i>d</i> (EH)	<i>d</i> (HH)	∑ <i>d</i>	RC(<i>x</i>) ^b	RC ^c
2-NHPh	1.78	2.06	1.02	3.81	8.67	0.00	0.00
Δ	0.20	-0.16	0.24	-2.47	-2.36		
[2-N-1] [†]	1.98	1.90	1.26	1.17	6.31	2.36	0.39
Δ		-0.09		-0.43	-3.76		
1=NPh		1.81		0.74	2.55	6.12	1.00
2-PHPh	1.76	2.64	1.44	4.18	10.02	0.00	0.00
Δ	0.14	-0.16	0.25	-2.97	-2.74		
[2-P-1] [†]	1.90	2.48	1.69	1.21	7.28	2.74	0.40
Δ		-0.12		-0.47	-4.18		
1=PPh		2.36		0.74	3.10	6.92	1.00
2-AsHPh	1.75	2.76	1.54	4.18	10.23	0.00	0.00
Δ	0.13	-0.16	0.21	-2.89	-2.71		
[2-As-1] [†]	1.88	2.60	1.75	1.29	7.52	2.71	0.38
Δ		-0.13		-0.55	-4.34		
1=AsPh		2.47		0.74	3.18	7.05	1.00

^a The reaction coordinate (RC) is roughly considered as the sum of the interatomic distances involved in the elimination; the product RC is simply defined as *d*(TaE) + *d*(HH). ^b The positional coordinate *x* indicates the total distance changes along the reaction coordinate. ^c The RC refers to the fraction of reaction progress relative to the pnictide-hydrides at 0.00.

1,2-H₂-elimination. Since the torsional motions of the atoms in the 1,2-H₂-elimination events are likely to parallel the bond strengths from the standpoint of energies (i.e., easier to bend X–E–H bonds as As(easier) < P < N), this parametrization of the RC is reasonable.

Dinitrogen Binding. 1. Collateral Discoveries. While the indirect synthesis of the hypothetical dinitrogen complex, [(silox)₃Ta]₂(μ-N₂) (**1**₂-N₂), which is likely to be a diimide,^{7–9} has not been realized, several discoveries during the course of these investigations are noteworthy. The synthesis of (silox)₃Ta=C=PPh₃ (**1**=CPPH₃), for example, may permit examination of the reactivity of an incipient “(silox)₃TaC” (**1**-C) via phosphine dissociation. Preliminary indications are that carbonylation yields the known ketylidene (silox)₃Ta=C=C=O (**1**=CCO), suggesting that loss of PPh₃ can occur under the right circumstances. The possibility of Wittig-like reactivity of the phosphalkylidyne fragment is another plausible path of discovery; studies are ongoing.

In the history of silox-based chemistry from these laboratories, the loss of silanol, other than from degradative hydrolysis, had not been cleanly observed until the thermolysis of (silox)₃MeTaNHNH₂ (**4**-NHNH₂) led to [(silox)₂TaMe](μ-N_αHN_β)(μ-N_γHN_δH)[Ta(silox)₂] (**5**). Unfortunately, the association of (silox)H with this complex, presumably via hydrogen bonding as solutions were concentrated, hampered purification of the unusual cyclic ditantalum species. Nonetheless, the loss of (silox)H via amination will be kept in mind as a future synthetic tool.

2. Compromise Diimide, [(silox)₂ClTa]₂(μ-N₂). The structure and stability of [(silox)₂ClTa]₂(μ-N₂) (**8**-Cl) leave little question that the hypothetical [(silox)₃Ta]₂(μ-N₂) (**1**₂-N₂) complex should be stable. Even though the crystallographic model precluded a true determination of *d*(TaN) and *d*(NN), the ditantalum interatomic distance of 4.874 Å is more consistent with two tantalum-imido interactions and an elongated dinitrogen distance than a bridging dinitrogen complex. Again, its stability with regard to dinitrogen loss supports the notion of the title quest.

3. Calculations on Dinitrogen Binding and Activation by (silox)₃Ta. Although [(silox)₂ClTa]₂(μ-N₂) (**8**-Cl)

provides a potent example of the entitled dinitrogen activation, the forward reaction of N₂ + “(silox)₂ClTa” cannot be examined because the *bis*-siloxchloride tantalum monomer is not a stable entity; only (silox)₃Ta (**1**) has been shown to possess the necessary sterics coupled to a unique electronic, 3-coordinate environment to exist in monomer form.⁴ As a consequence, high level quantum calculations were performed on the hypothetical dinitrogen activation illustrated in Scheme 1.

Figure 8 illustrates some of the complexity pertaining to dinitrogen activation by (silox)₃Ta (**1**, ⁿTa), first in binding N₂ to form (silox)₃TaN₂ (**1**-N₂, ⁿTa–N₂), and second to form the diimide (silox)₃Ta=N–N=Ta(silox)₃ (**1**₂-N₂, ⁿTa–NN–ⁿTa) within the constraints of a *linear* reaction coordinate. 3-Coordinate **1** is a singlet (¹Ta, (d_{z²})²) with *d*(TaO) of 1.89 Å and O–Ta–O angles of 120(1)^o that is ~19 kcal/mol below the triplet of nearest energy, ³Ta, whose electronic configuration is (d_{z²})¹(d_{xz} or d_{yz})¹, that is, ³E'' in D_{3h}. Borrowing from previous calculations,⁸⁴ the barrier to convert from ¹Ta to ³Ta is a few kcal/mol higher. Binding N₂ requires intersystem crossing to a triplet surface, as **1**-N₂ exists as a triplet that is 7.2 kcal/mol above the separated reagents, and ~5.5 kcal/mol below its respective singlet. Triplet ³Ta–N₂ possesses a *d*(TaN) of 1.96 Å, an elongated *d*(NN) of 1.18 Å (relative to the calculated 1.108 Å for free N₂), and a slightly pyramidalized core suggesting only modest reorganization energy. The electronic configuration of d_{xz}¹d_{yz}¹ for ³Ta–N₂ (³A₂ in C_{3v}) correlates with neither the ground state (GS) nor the first excited state (ES) of **1**, and suggests that a higher lying triplet (³A₂'', red dashed line) must somehow be accessed for binding. In summary, *all transition states for ⁿTa + N₂ → ⁿTa–N₂ require intersystem crossing events that are forbidden by orbital symmetry.* Some also require spin crossovers, but previous calculations suggest that these are inconsequential, especially within second and third row transition metal systems.^{83–88}

The second step, binding of (silox)₃TaN₂ (**1**-N₂, ³Ta–N₂) to another (silox)₃Ta (**1**, ⁿTa) can be estimated to be roughly the same energy as the initial binding event, but the surface is now a quintet, since both tantalum centers are triplets. One possible way for (silox)₃Ta (**1**) to bind **1**-N₂ is to intersystem cross to the same triplet that may be necessary to bind N₂ in the first step (i.e., the dashed blue surface), incurring a large barrier in the process. Once the binding has occurred to give (silox)₃Ta–N₂–Ta(silox)₃ (**1**₂-N₂, ⁿTa–NN–ⁿTa), electronic reorganization must occur to reach the singlet surface of the GS product. Since the diimide product (*d*(TaN) = 1.84 Å; *d*(NN) = 1.29 Å; ∠O–Ta–O = 109(1)^o) is ~55 kcal/mol lower than 2 **1** + N₂, orbital symmetry constraints may be more readily

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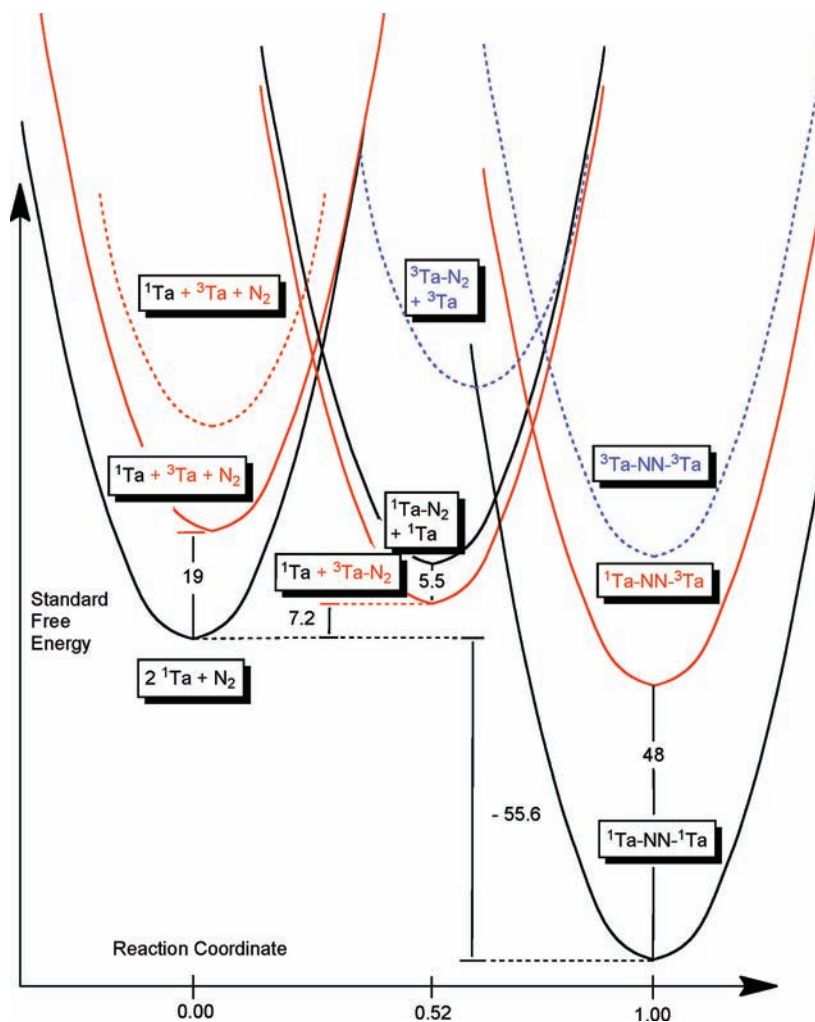


Figure 8. Plausible *linear* reaction coordinate (RC) versus standard free energy diagram for 2 (silox)₃Ta (**1**) + N₂ ((silox)₃Ta = ⁿTa) to hypothetical (silox)₃TaN₂Ta(silox)₃ (**1**₂-N₂, ⁿTa-NN-ⁿTa) via (silox)₃TaN₂ (**1**-N₂, ⁿTa-N₂). Calculated standard free energies are in kcal/mol and the RC is based on the calculated *d*(Ta-N) and *d*(N-N) changes, with all surface representations parabolically equivalent. The energies of species pertaining to dashed surfaces have been estimated.

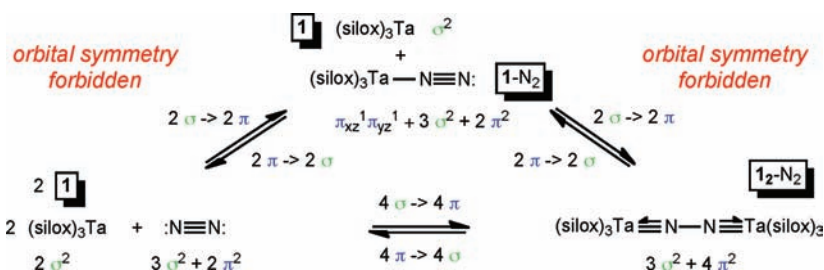


Figure 9. Simple illustration of orbital symmetry constraints in the conversion of 2 (silox)₃Ta (**1**) + N₂ ((silox)₃Ta = ⁿTa) to hypothetical (silox)₃TaN₂Ta(silox)₃ (**1**₂-N₂, ⁿTa-NN-ⁿTa) via (silox)₃TaN₂ (**1**-N₂, ⁿTa-N₂).

overcome, but the reduction of dinitrogen to diimide is also orbital symmetry forbidden from **1**-N₂.

It is clear from Figure 8 that the constraints of orbital symmetry must be overcome in both the initial and the second binding events of N₂. One way to overcome the electronic constraints depicted, that is, lower the intersystem crossing barriers, is to consider non-linear paths that effectively “mix” electron configurations that are intrinsically disparate, namely, those of σ -character with those of π -character. These paths are too varied to individually address without extensive calculations or

additional evidence, but one reaction coordinate pertinent to the reorganization of (silox)₃TaN₂Ta(silox)₃ (**1**₂-N₂, ⁿTa-NN-ⁿTa) has already been vetted.

The cleavage of N₂ by (ArRN)₃Mo to GS singlet nitride products, 2 (ArRN)₃MoN, is orbital symmetry forbidden.^{19–23} In this case, the binding of N₂ to form [(ArRN)₃Mo]₂(μ -N₂) is aided by the *S* = 3/2 states of the Mo(III) precursor, which require little promotional energy (to Mo(III) *S* = 1/2 states) to achieve the *S* = 1 state of the dinitrogen intermediate. Compared to the tantalum case, binding is readily accomplished, and it is

the *cleavage* of N_2 that is orbital symmetry forbidden. Scission eventually occurs because the parentage of only one orbital pair needs to be converted from π - to σ -character, and there is a considerably favorable free energy to aid in the process. In the tantalum case, each step (Figure 9) carries a significant energy penalty to overcome orbital symmetry constraints, either in changing electronic surfaces or in the significant reorganization energy required in lowering the symmetry to mix σ - and π -character.

Experimental Section

General Considerations. All manipulations were performed using either glovebox or high vacuum line techniques. All glassware was oven-dried. THF and ether were distilled under nitrogen from purple sodium benzophenone ketyl and vacuum transferred from the same prior to use. Hydrocarbon solvents were treated in the same manner with the addition of 1–2 mL/L tetraglyme. Benzene- d_6 and toluene- d_8 were dried over sodium, activated 4 Å molecular sieves, vacuum transferred, and stored under nitrogen. THF- d_8 was dried over sodium and vacuum transferred from sodium benzophenone ketyl prior to use. H_2 and D_2 were passed over a column containing activated 4 Å sieves and copper oxide. Primary amines, TMSCl, 1,4-cyclohexadiene, and phenylarsine oxide were purchased from Aldrich and dried over activated 4 Å sieves. NH_3 (Matheson) was dried over sodium and degassed prior to use. PH_3 (Matheson) was used as received. AsH_3 (**Caution! Danger, very toxic!**) was generated in small quantities from dilute mineral acid digestion of Zn_3As_2 (Cerac). Phenyl phosphine was purchased from Aldrich and used without further purification. $(^1Bu_3SiO)_3Ta$ (**1**),² $(^1Bu_3SiO)_3TaH_2$ (**2-H**),³³ and phenyl arsine⁸⁹ were prepared following published procedures.

1H , $^{13}C\{^1H\}$, and ^{31}P NMR spectra were obtained on Varian XL-200 and XL-400, and Inova 400, 500 and 600 MHz spectrometers and chemical shifts are reported relative to benzene- d_6 (1H , δ 7.15; $^{13}C\{^1H\}$, δ 128.00) or external H_3PO_4 (^{31}P , δ 0.00). UV-vis spectra were acquired on a Hitachi U-2000 spectrometer, and IR spectra were recorded on a Mattson FT-IR, Perkin-Elmer 299B grating IR, or PE 377 grating IR. Combustion analyses were performed by Oneida Research Services (Whitesboro, NY), Robertson Microlit Laboratories (Madison, NJ), or Texas Analytical (Houston, TX). Molecular weight determinations via cryoscopy in benzene or vapor phase osmometry were performed on home-built instruments.

Procedures. **1.** $(^1Bu_3SiO)_3HTaNH_2$ (**2-NH₂**). To a 10 mL flask containing 360 mg ($SiOx)_3Ta$ (**1**, 0.435 mmol) was distilled 8 mL of hexanes at $-78^\circ C$. The flask was opened to a calibrated gas bulb containing NH_3 (0.456 mmol, 1.05 equiv) and allowed to warm to $23^\circ C$ with stirring. The initial blue color discharged completely after 30 min. Reduction of the volume to 3 mL, cooling to $-78^\circ C$, and filtration led to the isolation of 217 mg of white crystals (59%). Anal. Calcd. for $C_{36}H_{84}O_3NSi_3Ta$: C, 51.22; H, 10.03; N, 1.66. Found: C, 51.31; H, 10.04; N, 1.41.

2. $(^1Bu_3SiO)_3HTaPH_2$ (**2-PH₂**). To a 100 mL flask containing $(SiOx)_3Ta$ (**1**, 0.341 g, 0.412 mmol) was distilled 50 mL of hexane at 77 K. PH_3 was condensed into the flask from a calibrated bulb (415 Torr in 91 mL, 2.06 mmol, 5 equiv), and the solution was allowed to warm slowly to $23^\circ C$. The initial blue color faded to green, then yellow over a period of 30 min, and a red solid formed. After another 30 min, the excess PH_3 was removed, and the volume of the solution reduced to 10 mL. The solution was filtered, concentrated, and cooled to $-78^\circ C$, to yield pale yellow crystals (0.192 g, 54%). IR (nujol, cm^{-1}) 2285

(m), 2275 (m), 1775 (br, s), 1450 (m), 1375 (m), 1065 (w), 1010 (w), 1000 (w), 970 (w), 930 (w), 880 (br, s), 800 (br, s), 625 (s). M_r found: 858(68); calcd: 861. Combustion analysis was precluded by thermal instability.

3. $(^1Bu_3SiO)_3HTaPPh$ (**2-PPh**). A 25 mL flask was charged with $(SiOx)_3Ta$ (**1**, 0.682 g, 0.824 mmol), and 10 mL of pentane was added at $-78^\circ C$. Phenyl phosphine (90.7 μL , 0.825 mmol) was transferred into a separate flask, and 10 mL pentane was added at $-78^\circ C$. The solutions were warmed and stirred to ensure homogeneity. The solution of **1** was cooled to $-78^\circ C$, and the $PhPH_2$ solution was added via syringe in two portions. This mixture was stirred for 5 min and then warmed to $23^\circ C$ as a color change from blue to yellow-orange was observed (~ 10 min); stirring was continued for another 10 min. The volatiles were removed, and pentane (10 mL) was added. The solution was filtered, concentrated to 5 mL, and cooled to $-78^\circ C$ to afford bright yellow crystals (0.298 g, 39%). IR (nujol, cm^{-1}) 2320 (m), 1790 (br, s), 1580 (m), 1470 (s), 1375 (m), 1005 (w), 960–800 (br, s), 725 (m), 690 (m), 620 (s). Combustion analysis was precluded because of thermal instability.

4. $(^1Bu_3SiO)_3Ta=NH$ (**2=NH**). A 100 mL flask was charged with **1** (0.979 g, 1.18 mmol), attached to a frit assembly, and evacuated. Hexane (50 mL) was added via vacuum transfer. 2-Methylaziridine (360 Torr in 91 mL, 1.79 mmol, 1.5 equiv) was added to the stirred solution at $-78^\circ C$. As the solution was warmed to room temperature, the blue color of the solution quickly faded to pale yellow. After stirring for 30 min the volatiles were removed and fresh hexane was added. The solution was filtered and concentrated to 10 mL. White crystals formed upon cooling to $-78^\circ C$ and were isolated by filtration (0.771 g). A second crop was obtained in a like manner (0.080 g, 85% total). IR (nujol, cm^{-1}) 3470 (m), 1375 (m), 1010 (w), 935 (w), 880 (br, s), 820 (s), 630 (s). Anal. Calcd for $C_{36}H_{82}O_3N-Si_3Ta$: C, 51.34; H, 9.81; N, 1.66. Found: C, 51.38; H, 10.09; N, 1.54.

5. $(^1Bu_3SiO)_3Ta=NCH_3$ (**1=NCH₃**). Into a glass bomb reactor was transferred a solution of $(SiOx)_3Ta$ (**1**, 0.438 g, 0.529 mmol) in hexane (10 mL). The solution was freeze/pump/thaw degassed three times, and $MeNH_2$ was condensed in from a calibrated gas bulb (77 Torr in 125 mL, 0.53 mmol). As the bomb warmed to $23^\circ C$, the blue color faded to pale yellow. After stirring for 12 h, the solution was frozen and a copious quantity of gas was evacuated from the bomb. The solution was again warmed and stirred. This cycle was repeated intermittently over a period of 3 d until no more gas was evolved. The resulting solution was concentrated to 4 mL, cooled to $-78^\circ C$, and filtered to yield a white powder (0.228 g, 50%). IR (nujol, cm^{-1}) 1475 (s), 1375 (m), 1320 (m), 1010 (w), 975 (m), 935 (w), 875 (br, s), 820 (s), 625 (s). Anal. Calcd for $C_{37}H_{84}O_3NSi_3Ta$: C, 51.90; H, 9.89; N, 1.64. Found: C, 51.79; H, 9.79; N, 1.45.

6. $(^1Bu_3SiO)_3Ta=NPh$ (**1=NPh**). To a glass bomb reactor was added a solution of $(SiOx)_3Ta$ (**1**, 0.486 g, 0.587 mmol) dissolved in 6 mL of benzene. Aniline (53.5 μL , 0.587 mmol) was dissolved in benzene (2 mL) and added to the bomb. This mixture was freeze/pump/thaw degassed three times and stirred at $23^\circ C$ for 6 h, during which time a generous quantity of gas evolved. The solution was degassed and warmed to $23^\circ C$ with continued stirring, and the process was repeated after an additional 12 h. The solution was then heated to $100^\circ C$ for 30 min and cooled to $23^\circ C$. After the volatiles were removed, 15 mL of hexane was added, and the solution was degassed, filtered, and the volatiles removed. Ether (5 mL) was added, and the solution was cooled to $-78^\circ C$ to afford white crystals (0.346 g, 64%). IR (nujol, cm^{-1}) 1590 (m), 1475 (s), 1385 (m), 1360 (s), 1010 (w), 985 (w), 950 (s), 875 (br, s), 820 (s), 750 (s), 685 (m), 620 (s). Anal. Calcd for $C_{42}H_{86}O_3NSi_3Ta$: C, 54.93; H, 9.44; N, 1.53. Found: C, 54.92; H, 9.45; N, 1.48.

7. $(^1Bu_3SiO)_3Ta=N(C_6H_4-p-CF_3)$ (**1=NC₆H₄-p-CF₃**). A sample of $(SiOx)_3Ta$ (**1**, 0.208 g, 0.251 mmol) was dissolved in 5 mL

(89) (a) Doak, G. O.; Freedman, L. D. *Organometallic Compounds of Arsenic, Antimony, and Bismuth*; Wiley-Interscience: New York, 1970. (b) Wiberg, K. B. Z. *Naturforsch.* **1957**, *12B*, 127–128.

of hexane and placed in a glass bomb reactor. Dihydrogen (~600 Torr at 77 K) was admitted to the reactor, and the solution was stirred at 23 °C for 20 h to ensure formation of (silox)₃TaH₂ (2-H), and degassed. In a glovebox, 4-aminobenzotrifluoride (31.6 μL, 0.252 mmol) was syringed into the hexane solution, which was degassed. After thermolysis in a wax bath at 90 °C for 10 h, the contents were transferred and the volatiles removed. Ether was added, and the solution was filtered. Concentration to 2 mL and cooling to -78 °C afforded white crystals (0.097 g, 39%). IR (Nujol, cm⁻¹) 1605 (m), 1510 (m), 1475 (s), 1375 (s), 1320 (s), 1200 (w), 1170 (m), 1160 (s), 1120 (s), 1110 (m), 1075 (m), 1010 (w), 960 (s), 880 (s, br), 840 (m), 820 (s), 645 (w), 635 (s). Anal. Calcd for C₄₃H₈₅O₃F₃NSi₃Ta: C, 51.14; H, 8.69; N, 1.42. Found: C, 52.62; H, 8.86; N, 1.31.

8. (^tBu₃SiO)₃Ta=NLi (I=NLi). Glassware for this reaction was silylated in the following manner. After drying in an oven and cooling under vacuum, hexamethyldisilazane was introduced into a frit assembly against N₂ counterflow. The liquid was freeze/pump/thaw degassed three times and then warmed to reflux with a heat gun until it completely covered the inner surface of the glassware. The liquid was cooled and removed in vacuo, and the glassware transferred into a glovebox. One flask was charged with (silox)₃Ta=NH (I=NH, 0.473 g, 0.562 mmol) and neopentyl lithium (0.066 g, 0.845 mmol, 1.5 equiv). THF (25 mL) was transferred via vacuum transfer at -78 °C. The resulting solution went from yellow to colorless as it warmed to 23 °C, and after stirring for 1.5 h, the solution was filtered, concentrated to 5 mL, and cooled to -78 °C to afford colorless crystals (0.371 g, 78%). IR (nujol, cm⁻¹) 1460 (s), 1375 (s), 1010 (w), 900 (br, s), 820 (m), 625 (m). Anal. Calcd for C₃₆H₈₁O₃NLiSi₃Ta: C, 50.98; H, 9.63; N, 1.65. Found: C, 50.92; H, 9.84; N, 1.41.

9. (^tBu₃SiO)₃Ta=NSiMe₃ (I=NTMS). A flask attached to a calibrated gas bulb was silylated as described above. The flask was charged with (silox)₃Ta=NLi (I=NLi, 0.129 g, 0.152 mmol). THF (20 mL) was added at -78 °C followed by TMSCl (30 Torr in 91 mL, 0.15 mmol). Upon warming to 23 °C while stirring for 3 h, the volatiles were removed, and the solid was extracted into hexane. The solution was filtered, concentrated to 3 mL, and cooled to -78 °C, yielding white crystals (0.070 g, 50%). IR (nujol, cm⁻¹) 1475 (s), 1375 (m), 1245 (m), 1165 (s), 1010 (w), 950 (s), 870 (br, s), 820 (s), 745 (m), 630 (w), 625 (s). Anal. Calcd for C₄₀H₉₀O₃NSi₄Ta: C, 51.23; H, 9.92; N, 1.53. Found: C, 51.56; H, 9.94; N, 1.56.

10. (^tBu₃SiO)₃Ta=PH (I=PH). A solution of **1** (0.476 g, 0.575 mmol) in hexane (50 mL) was exposed to 5 equiv of PH₃, which was admitted via a calibrated gas bulb. After stirring for 1 h, the solution was filtered and transferred to a glass bomb reactor, where it was freeze/pump/thaw degassed three times, warmed, and placed in an oil bath at 70 °C. After 2 h the solution was again frozen at 77 K and evacuated. A large quantity of gas was observed. The process was repeated until no more gas was evolved (two more cycles). The volatiles were removed, and 10 mL of hexane added. The solution was filtered, concentrated to 5 mL, and cooled to -78 °C, affording pale orange microcrystals (0.301 g, 61%). IR (nujol, cm⁻¹) 2150 (w), 1450 (m), 1375 (m), 1000 (w), 960 (m), 930 (w), 865 (s), 815 (m), 625 (m). *M_r* found: 857(50); calcd: 859. Anal. Calcd for C₃₆H₈₂O₃PSi₃Ta: C, 50.32; H, 9.62. Found: C, 50.32; H, 10.05.

11. (^tBu₃SiO)₃Ta=PPh (I=PPh). The residue from the preparation of 2-HPPH was dissolved in hexane and transferred into a glass bomb reactor. The contents were freeze/pump/thaw degassed three times. The bomb was warmed to 23 °C, and the solution stirred for 18 h. The bomb was degassed and immersed in a 55 °C bath. It was stirred for 3 d with intermittent degassing until gas evolution ceased and the solution was deep red. The volatiles were removed and 10 mL of pentane was added. The solution was transferred, concentrated to 3 mL, and cooled to -78 °C, yielding red-violet crystals (0.293 g, 62% based on

remaining Ta). IR (nujol, cm⁻¹) 1580 (w), 1475 (m), 1375 (m), 1010 (w), 950 (m), 850 (s), 820 (m), 730 (w), 695 (w), 625 (m). Anal. Calcd for C₄₂H₈₆O₃PSi₃Ta: C, 53.93; H, 9.27. Found: C, 53.50; H, 9.44.

12. (^tBu₃SiO)₃Ta=AsPh (I=AsPh). To a flask charged with **1** (0.615 g, 0.743 mmol) was distilled 10 mL of toluene at -78 °C. A solution of phenyl arsine (84.5 μL, 0.744 mmol) in toluene (5 mL) was added via syringe. The blue color of the solution faded noticeably after 15 min and changed to yellow-orange after 1 h. Stirring was continued for an additional hour, and upon warming, the solution effervesced and turned red, then green. After the reaction subsided, the volatiles were removed, and the residue was triturated three times with 5 mL of hexane. Hexane (10 mL) was added, and the solution was filtered. The residual was washed, and the filtrate was concentrated to 6 mL, and cooled to afford green microcrystals (0.511 g). A second crop was also isolated (0.109 g, 85% total). IR (nujol, cm⁻¹) 1576 (w), 1470 (m), 1375 (m), 1010 (w), 945 (s), 850 (br, s), 820 (s), 725 (w), 690 (w), 620 (m). *M_r* found: 980(45); calcd: 979. Anal. Calcd for C₄₂H₈₆O₃Si₃AsTa: C, 51.51; H, 8.85. Found: C, 51.17; H, 8.56.

13. (^tBu₃SiO)₃Ta=AsH (I=AsH). Conducted on a Schlenk Line in a Hood. In a separate 2-neck flask attached to a needle valve and fitted with a bent tube containing solid Zn₃As₂ (34.4 mg, 0.0968 mmol, 1.2 equiv As) was frozen dilute sulfuric acid which was carefully degassed. The Zn₃As₂ was tapped into the acid at 0 °C to generate small quantities of AsH₃ (**Caution! Danger, Toxic**). The needle valve was opened and the AsH₃ was allowed to expand through a cold trap containing KOH and admitted to another flask attached to a frit assembly containing (silox)₃Ta (**1**, 0.200 g, 0.242 mmol) in hexane at -78 °C. The solution became orange immediately and the evolution of a gas, presumably H₂ was noted, along with the formation of a red precipitate. The red solid was removed by filtration and dark orange crystals were obtained from the solution. ¹H NMR analysis revealed the material to be composed of (silox)₃Ta=AsH (I=AsH) and (silox)₃Ta(AsH₂)₂ (3-(AsH₂)₂) in a 9:1 ratio. Any remaining AsH₃ was exposed to a cold NaOCl solution to oxidize the gas, and the Schlenk line was thoroughly evacuated through an LN₂ trap prior to further use.

14. (^tBu₃SiO)₃Ta=CH₂ (I=CH₂). To a small bomb reactor containing **1** (2.01 g, 2.40 mmol), 690 mg of PPh₃CH₂ was added 40 mL of THF (40 mL). The reaction mixture was stirred for 1 h at 23 °C, degassed, and transferred for workup. The solution was filtered, concentrated to 15 mL, and cooled to -78 °C to afford 1.43 g (~70%) of yellow crystals that contained ~6% PPh₃ by mass. This was often sufficient for further reactions. Recrystallization from THF, then Et₂O, yielded pure yellow crystals (29%) that were submitted for EA. Anal. Calcd for H₈₃C₃₇O₃Si₃Ta: C, 52.83; H, 9.94. Found: C, 52.91; H, 9.84.

15. (^tBu₃SiO)₃Ta=CPh₃ (I=CPh₃). To a 50 mL flask charged with **1** (1.96 g, 2.40 mmol) and 687 mg of PPh₃CH₂ at -78 °C was added 40 mL of THF by vacuum transfer. The solution was allowed to slowly warm to 23 °C over 12 h. The solution was degassed, filtered, concentrated to 15 mL, cooled to -78 °C, and filtered to yield 1.15 g **I=CH₂** (58%). The filtrate residue was dissolved in 5 mL of pentane and cooled to -78 °C to yield 371 mg of an off-white powder identified (NMR) as an equimolar mixture of PPh₃ and **I=CPh₃**. This material was dissolved in 5 mL of benzene and CH₃I (2 equiv, relative to PPh₃) was added and stirred overnight. A white precipitate was separated by filtration, and the solvent was removed in vacuo to yield 256 mg (10%, based on **1**) of **I=CPh₃** as a white powder. X-ray quality crystals could be grown from cooling a saturated pentane solution to -40 °C.

16. (silox)₃MeTaNHNH₂ (4-NHNH₂). To a 10 mL round-bottom flask charged with **I=CH₂** (560 mg, 0.665 mmol) and 8 mL of benzene was added 100 μL of hydrazine (3.1 mmol, 4.7 equiv) via syringe. The solution was stirred for 45 min, and

the solvent was removed in vacuo. The resultant off-white solid was triturated twice with 5 mL of THF, and 5 mL of pentane was added via vacuum transfer. The suspension was cold-filtered ($-78\text{ }^{\circ}\text{C}$), and the filter cake washed with 5 mL of pentane. Upon removal of the volatiles, the residual was dissolved in 20 mL of benzene and filtered twice through Celite, each time followed by a 20 mL benzene wash. The filtrates were combined, and the solvent was removed in vacuo to yield 360 mg of 4-NHNH₂ (49%) as a white powder. Anal. Calcd for C₃₇H₈₇N₂O₃Si₃Ta: C, 50.89; H, 10.04; N, 3.21. Found C, 50.94; H, 10.28; N, 3.09.

17. (silox)₃MeTaNH(-^cNCHMeCH₂) (4-NH(azir)). A 10 mL round-bottom flask was charged with 200 mg of **1**=CH₂ (0.24 mmol) and attached to a 180° valve adapter. The assembly was evacuated, and 2 mL of diethyl ether was added via vacuum transfer. To this solution was added 250 μL of *N*-amino-2-methylaziridine (3 mmol) at room temperature under argon counter-flow. The solution discolored over a period of 5 minutes, and the solvent and excess aziridine removed in vacuo, and the resulting waxy solid was left under dynamic vacuum for 45 min. The solid was recrystallized from pentane (0.5 mL) at $-40\text{ }^{\circ}\text{C}$ allowing for slow evaporation and washed with 0.5 mL of cold ($-40\text{ }^{\circ}\text{C}$) pentane to yield 70 mg (32%) of a thermally sensitive waxy solid.

18. [(silox)₂TaMe](μ-N_αHNβ)(μ-N_γHNδH)[Ta(silox)₂] (5). A small bomb was charged with 95 mg of 4-NHNH₂ (0.11 mmol), and 2 mL of benzene was added. The bomb was heated to 75 °C for 24 h and subsequently evacuated of all volatiles. The resulting oil was dissolved in pentane, removed from the bomb, and placed in a 4-dram vial where the solvent was removed in vacuo to yield a slightly yellow liquid (90 mg, 95%). ¹H NMR analysis indicated complete conversion. The complex could not be separated from 2 equiv of ¹Bu₃SiOH, and was characterized by multinuclear NMR spectroscopy.

19. [(silox)₃MeTa](μ-η²-N,N:η¹-C-NHNHCH₂CH₂CH₂)-[Ta(κ-O,C-OSi^tBu₂CMe₂CH₂)(silox)₂] (7). To a resealable NMR tube was added 61 mg of 4-NH(azir) (0.67 mmol) and 0.7 mL of C₆D₆. The tube was heated for 4 h at 80 °C until the starting material had disappeared. The resulting solution was removed from the tube and placed in a 1-dram vial, and a solution of (silox)₃Ta (58 mg in 1 mL, 0.070 mmol) was added. From the resulting green solution, yellow needles precipitated which were isolated by decanting the mother-liquor. The needles were washed with pentane and dried in vacuo to yield 15 mg of effectively insoluble material that could be recrystallized from hot THF (80 °C).

20. [(^tBu₃SiO)₂TaCl]₂(μ-N₂) (8-Cl). To a 50 mL round-bottom flask containing 1.000 g of [TaCl₃(THF)₂]₂N₂ (1.12 mmol) and 1.09 g of sodium silox (4.56 mmol, 4.06 equiv) at $-78\text{ }^{\circ}\text{C}$ was added 25 mL of THF via vacuum transfer. The flask was allowed to warm to 23 °C over 18 h during which time the color changed from red-orange to yellow. The solvent was removed in vacuo, and 10 mL of pentane was added. Filtering the suspension, concentrating the solution to 5 mL, and cooling to $-78\text{ }^{\circ}\text{C}$ afforded yellow microcrystals which were dried in vacuo. After collection of a second crop the total yield was 828 mg (56%).

21. [(^tBu₃SiO)₂TaMe]₂(μ-N₂) (8-Me). To a 25 mL flask charged with 100 mg of (0.075 mmol) [(silox)₂TaCl]₂N₂ at $-78\text{ }^{\circ}\text{C}$ was added 10 mL of Et₂O via vacuum transfer. After the solid dissolved, 0.08 mL of CH₃MgBr (2.0 M in Et₂O, 2.1 equiv) was added via syringe. The solution was allowed to warm to 23 °C over an 8 h period. After 30 min at 23 °C, a white precipitate formed, and the solution was filtered. The solution was filtered, concentrated to 2 mL, and cooled to $-78\text{ }^{\circ}\text{C}$, affording a light-yellow powder that was collected by filtration and dried in vacuo (50 mg, 52%).

NMR Tube Reactions. General Procedures. Solid (^tBu₃SiO)₃Ta (**1**) or (^tBu₃SiO)₃TaH₂ (**2-H**), typically 15–20 mg (0.017–0.025 mmol) was added to an NMR tube attached to a ground glass joint. Any solid reagents were also added along with

deuterated solvent ($\sim 0.6\text{--}0.8\text{ mL}$). Liquid reagents were added via microliter syringe. The tube was attached to a needle valve, and the tube was degassed and sealed on a vacuum line. Any gaseous reagents were added via calibrated gas bulbs, and solvents were added to solid reagents via vacuum distillation prior to sealing.

22. (silox)₃HTaAsH₂ (2-AsH). Conducted on a Schlenk Line in a Hood. In a separate 2-neck flask attached to a needle valve and fitted with a bent tube containing solid Zn₃As₂ was frozen dilute sulfuric acid which was carefully degassed. The Zn₃As₂ was tapped into the acid at 0 °C to generate small quantities of AsH₃ (**Caution! Danger, Toxic**). The needle valve was opened, and the AsH₃ was allowed to expand through a cold trap containing KOH and admitted to an NMR tube containing (silox)₃Ta (**1**), an equiv of 1,4-cyclohexadiene (added via vacuum transfer), and toluene-*d*₈. After the tube was sealed, any remaining AsH₃ was exposed to a cold NaOCl solution to oxidize the gas, and the Schlenk line was thoroughly evacuated through an LN₂ trap prior to further use. The NMR tube was kept at $-78\text{ }^{\circ}\text{C}$ for NMR monitoring.

General Kinetics. Solutions of (silox)₃HTaEHR (**2-EHR**) were generated in 2.0 mL volumetric flasks by one of three procedures: from isolated **2-EHR** and C₆D₆, dissolving (silox)₃Ta (**1**) in C₆D₆ and adding REH₂ via syringe, or dissolving (silox)₃TaH₂ (**2-H**) in C₆D₆ and adding REH₂ via syringe. An exception was (silox)₃HTaNHMe (**2-HNMe**), which required addition of MeNH₂ from gas bulbs to individual tubes of **1** in C₆D₆. TMS₂O ($\sim 0.5\text{ }\mu\text{L}$) was added as an internal integration standard. The samples ($\sim 0.6\text{ mL}$) were transferred to NMR tubes sealed to 14/20 ground glass joints and attached to 180° needle valves. The tubes were freeze/pump/thaw degassed three times and sealed with a torch. Samples were kept in a constant temperature oven at 24.8(4)°C or were thermolyzed in a Tamson TX9 constant temperature bath (**2-NH₂**). The rate of disappearance of **2-EHR** was typically monitored via disappearance of the hydride ¹H NMR resonance (see Table 1 for exceptions) for 5–6 half-lives. Single-transient spectra were used for reproducibility of integration. In all cases where appearance of **1**=ER was monitored, the rate was found to be the same as disappearance of **2-EHR**. Rates and uncertainties were obtained from unweighted, non-linear, least-squares fitting of the exponential form of the rate expression (averaged where simultaneous runs were obtained).

Solutions of (silox)₃HTaAsPh (**2-HAsPh**) were generated in the following manner. PhAsH₂ was syringed into volumetric flasks which were then attached to needle valves. The flasks were cooled to 77 K, evacuated, and toluene-*d*₈ was added via vacuum transfer. The resulting solutions were added to NMR tubes containing **1** via vacuum transfer at 77 K, and 1,4-cyclohexadiene was added from gas bulbs. The tubes were sealed with a torch and maintained at 77 K until immediately prior to each run. The tubes were warmed to $-78\text{ }^{\circ}\text{C}$ in dry ice/acetone baths and agitated until the blue color of **1** had bleached. The samples were transferred to the NMR probe which had been previously cooled to $-78\text{ }^{\circ}\text{C}$. Initial spectra were obtained to confirm the complete formation of **14-AsHPh**. The temperature of the probe was then raised to $-9.5\text{ }^{\circ}\text{C}$ and single-transient spectra were recorded at measured time intervals.

Single-Crystal X-ray Diffraction Studies. Upon isolation, the crystals were covered in polyisobutenes and placed under a 173 K N₂ stream on the goniometer head of a Siemens P4 SMART CCD area detector (graphite-monochromated Mo K α radiation, $\lambda = 0.71073\text{ \AA}$). The structures were solved by direct methods (SHELXS). All non-hydrogen atoms were refined anisotropically unless stated, and hydrogen atoms were treated as idealized contributions (Riding model).

23. (^tBu₃SiO)₃Ta=NPh (1=NPh). A colorless block (0.3 × 0.3 × 0.4 mm) was obtained from a hot heptane solution, and was sealed in a 0.5 mm glass capillary. A total of 5077 reflections

were collected with 4665 being symmetry independent ($R_{\text{int}} = 0.0254$), and 3371 were greater than $2\sigma(I)$. A semiempirical absorption correction from equivalents was applied, and the refinement utilized $w^{-1} = \sigma^2(F_o^2) + (0.0549p)^2 + 7.4307p$, where $p = (F_o^2 + 2F_c^2)/3$. The ^tBu groups of one silox group were disordered and modeled accordingly.

24. (^tBu₃SiO)₃Ta=CPPh₃ (**1=CPPh₃**). A colorless block (0.15 × 0.20 × 0.25) was obtained from cooling a saturated pentane solution to −40 °C. A total of 73,173 reflections were collected with 17,745 being symmetry independent ($R_{\text{int}} = 0.0766$) and 12,757 were greater than $2\sigma(I)$. A semiempirical absorption correction from equivalents was applied, and the refinement utilized $w^{-1} = \sigma^2(F_o^2) + (0.0260p)^2 + 0.0173p$, where $p = (F_o^2 + 2F_c^2)/3$.

25. [(silox)₃MeTa](μ - η^2 -N,N: η^1 -CNHNHCH₂CH₂CH₂)[Ta-(k-O,C-OSi^tBu₂CMe₂CH₂)(silox)₂] (**7**). A colorless plate (0.03 × 0.10 × 0.20) was obtained from cooling a hot THF solution. A total of 57,689 reflections were collected with 17,150 being symmetry independent ($R_{\text{int}} = 0.0762$) and 11,852 were greater than $2\sigma(I)$. A semiempirical absorption correction from equivalents was applied, and the refinement utilized $w^{-1} = \sigma^2(F_o^2) + (0.0624p)^2 + 15.5864p$, where $p = (F_o^2 + 2F_c^2)/3$. Disordered solvent was SQUEEZED from the cell, and a disorder in the ^tBu groups of one silox ligand was modeled appropriately.

26. [(^tBu₃SiO)₂TaCl]₂(μ -N₂) (**8-Cl**). A colorless block (0.15 × 0.20 × 0.25) was obtained from slow evaporation of a 1:1

pentane/hexamethyldisiloxane solution at −78 °C over a period of 4 h. A total of 58,495 reflections were collected with 12,659 being symmetry independent ($R_{\text{int}} = 0.0686$), and 8,578 were greater than $2\sigma(I)$. A semiempirical absorption correction from equivalents was applied, and the refinement utilized $w^{-1} = \sigma^2(F_o^2) + (0.0605p)^2 + 73.4015p$, where $p = (F_o^2 + 2F_c^2)/3$. The disorder in the ^tBu groups of one silox was modeled appropriately. The electron density in the region between the two tantalums did not refine well, and the nitrogen atoms were refined isotropically. The distances and angles between the two tantalums are inadequately determined (see text).

Calculations. To obtain minima for these complexes, full geometry optimizations, without any metric or symmetry restrictions, were performed using the Gaussian 03⁹⁰ package, and these employed density functional theory (DFT), specifically the BLYP functional.⁹¹ Atoms were described with the Stevens effective core potentials (ECPs) and attendant valence basis sets (VBSs).⁹² This scheme, dubbed CEP-31G(d), entails a valence triplet zeta description for the transition metals, a double- ζ -plus-polarization basis set for main group elements, and the −31G basis set for hydrogen. This level of theory was selected on the basis of previous work.^{18,83}

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Supporting Information Available: CIF files for **1=NPh**, **1=CPPh₃**, **7**, and **8-Cl**. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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