

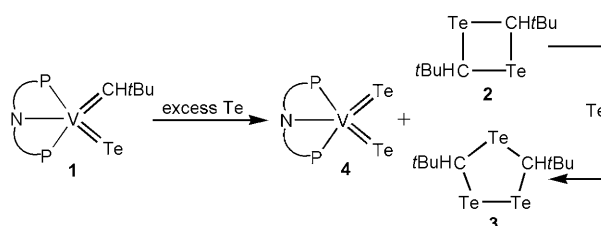
Tellus in, Tellus out: The Chemistry of the Vanadium Bis(telluride) Functionality**

Uriah J. Kilgore, Jonathan A. Karty, Maren Pink, Xinfeng Gao, and Daniel J. Mindiola*

Terminal telluride ligands bound to transition metals still represent a rare functionality,^[1–8] the first example (*trans*-[(Me₃P)₄W(Te)₂]) being reported by Rabinovich and Parkin in 1991.^[1] A terminal telluride on a 3d transition metal, namely the complex [(Me₃SiCH₂CH₂)₃N]V=Te) was later reported by Schrock and co-workers in 1994.^[2] Even though a few other examples of terminal early-transition-metal tellurides have been documented,^[3–8] the only structurally characterized terminal 3d transition metal/telluride complexes consist of [(η⁴-Me₈taa)M'=Te] (M' represents Ti and V, Me₈taa^{2–} = octamethyldibenzotetraaza[14]annulene).^[7] Therefore it is of no surprise that little is known about the reactivity of the M=Te functionality in question, especially when 3d transition metals are involved.^[2,8] For the few 3d examples known,^[2,8] it has been proposed that the M=Te linkage decomposes by reductive elimination of Te⁰, whereby the fate of the reduced metal complex M^{2–} can sometimes result in formation of dinuclear species.^[9,10] Only in the case of [(η⁴-Me₈taa)M'=Te] or [Cp^{*}2M=Te(NC₅H₅)], could the terminal Te^{2–} ligand be replaced with O^{2–} (derived from O₂ or N₂O), Se₂^{2–} (derived from elemental Se), and 2 Cl[–] from the addition of two equivalents of Me₃SiCl.^[7,8] Although not discussed, it can be argued that these transformations proceed by oxidation of the M=Te bond with concurrent Te⁰ ejection.

Based on the chemistry known for terminal tellurides, we hypothesized that the M'=Te functionality could be utilized as a labile group by the reductive elimination of the low-valent M^{n–2} fragment with concurrent extrusion of Te⁰. This feature is counterintuitive when considering the reactivity of the ubiquitous oxo ligand, given its propensity to stabilize high-valent oxidation states, and would imply that the metal center that has a terminal Te atom should be amenable to substitution and/or redox processes. Herein, we demonstrate that a vanadium alkylidene/telluride complex, [(PNP)V=

CH*t*Bu(Te)] (**1**) (PNP[–] = N(2-PiPr₂-4-methylphenyl)₂),^[11] prepared from [(PNP)V(CH₂*t*Bu)₂] and Te⁰ via the intermediate [(PNP)V=CH*t*Bu] (**A**), can reductively eliminate a telluroaldehyde fragment to afford the formation of 2,4-di-*tert*-butyl-1,3-ditellurethane (**2**) and/or 3,5-di-*tert*-butyl-1,2,4-tritellurolane (**3**; Scheme 1). The hypothetical “(PNP)V” fragment can be trapped with excess Te⁰ to afford the first 3d



Scheme 1. Synthesis of complex **2**, and the di- and tritellurolanes.

transition metal/bis(telluride) complex (PNP)V(Te)₂ (**4**).^[1,5,6] Through a series of reactivity studies we demonstrated the [Te=V^V=Te] moiety in **4** to be a synthon of the [Te=V^V=X] and [X=V^V=X] scaffolds, which are formed by oxidation of the V=Te bonds with concomitant deposition of Te⁰. In addition, we also show the ability of **4** to reductively eliminate one Te ligand to furnish a vanadium(III) complex that contains a terminal telluride moiety.

We recently reported that the complex (PNP)V(CH₂*t*Bu)₂ could be oxidized by chalcogen sources to promote α-hydrogen abstraction and ultimately form the alkylidene chalcogenides (PNP)V=CH*t*Bu(X) (X = O, S, Se, and Te) and CH₃*t*Bu.^[11] Complex **1** is remarkably stable in the solid state and in solution, but it was noted that the presence of traces of Te⁰ promoted gradual transformation of **1** into another vanadium species together with cyclic ditellurethane [(TeCH*t*Bu)₂] (**2**) and tritellurolane [Te(TeCH*t*Bu)₂] (**3**).^[12] Optimization of the reaction to yield these new products involved treatment of [(PNP)V(CH₂*t*Bu)₂] with an excess of Te powder (4–8 equivalents) under a static vacuum over 4 days at 90 °C; this procedure consistently yielded the dark-green insoluble vanadium complex together with the by-products **2** and **3** (Scheme 1). Evacuation of the solution was necessary in order to avoid activation of atmospheric nitrogen, which results in the formation of the known dinitrogen-bridged vanadium dimer [(PNP)V=CH*t*Bu]₂(μ₂;η¹,η¹-N₂).^[11a] The reaction mixture was vacuum dried and the organotellurium complexes **2** and **3** were extracted by washing the solids with hexanes or pentane. The identities of compounds **2** and **3** have been confirmed by a combination of ¹H and ¹³C NMR spectra and chemical ionization (CI) mass

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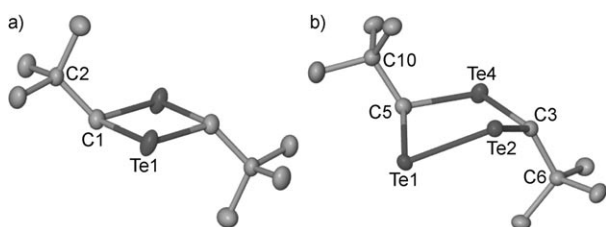


Figure 1. The molecular structures of compounds **2** (a) and **3** (b) showing thermal ellipsoids at the 50% probability level. Hydrogen atoms are omitted for clarity.

spectrometry. Given the rarity of ditelluretane^[13] and tritellurolane^[14a] derivatives in general, we collected single-crystal X-ray diffraction structures of each compound.^[12] Figure 1 shows the head-to-tail isomers of the four- and five-membered rings **2**^[15] and **3**,^[16] respectively. While the structure of **2** reveals a planar four-membered ring residing on an inversion center, compound **3** incorporates a puckered five-membered ring with a Te1–Te2 distance of 2.7190(3) Å.^[17] It was also determined that lower concentrations of Te (ca. 4–5 equivalents) with [(PNP)V(CH₂tBu)₂] greatly enhanced the yields of **2**, while 6–8 equivalents of the chalcogen facilitated formation of **3**. Previous work has speculated that the analogous 1,2,4-triselenolane formation derives from the insertion of Se⁰ into 1,3-diselenetane.^[14b] In our case, treatment of the reaction mixture containing **2** and **3** (after removal of the green vanadium-containing precipitate) with Te⁰ increased the concentration of **3**, thus supporting the original hypothesis that precursors such as **2** can ring expand by the uptake of elemental chalcogen. A handful of ditelluretanes that originate from transient telluroketones or perfluorinated telluracarbonyl compounds exist; however, to the best of our knowledge, the ditelluretane **2** is the first example of an isolated ditelluretane that originates from a transient telluroaldehyde [Te=CHtBu] monomer.^[13,14]

The insoluble green material that resulted from the reaction of **1** with an excess of Te⁰ (see above) was extracted with THF, filtered, and dried under reduced pressure to afford a green powder, which was collected in 74 % yield. The ¹H, ³¹P (δ = 142 ppm), and ⁵¹V (δ = 2293 ppm, triplet, Δν_{1/2} = 490, 299, 460 Hz) NMR spectra of the insoluble green material were consistent with a C₂-symmetric “[(PNP)V]” scaffold. As expected from the inverse electronegativity dependence of high-valent vanadium complexes, the ⁵¹V resonance of this new vanadium product is significantly downfield from that of complex **1** (δ = 1249 ppm, Δν_{1/2} = 864 Hz).^[18] Unfortunately, the low solubility of this material in haloarenes and THF (unstable in CHCl₃ and CH₂Cl₂) prevented us from observing a ¹²⁵Te NMR spectroscopic signal. Crystals suitable for X-ray diffraction could be grown from solution of THF that was heated at reflux, the data confirmed formation of an unprecedented vanadium-bis(telluride) monomeric complex [(PNP)V(Te)₂] (**4**, Figure 2).^[19] The terminal nature of the V=Te bond is clearly manifested by the short bond lengths of 2.4225(5) and 2.4223(4) Å, which are comparable to the only other crystallographically characterized terminal vanadium–telluride complex [(η⁴-Me₈taa)V=Te] (2.433(1) and 2.435(1) Å based on two crystallographically independent

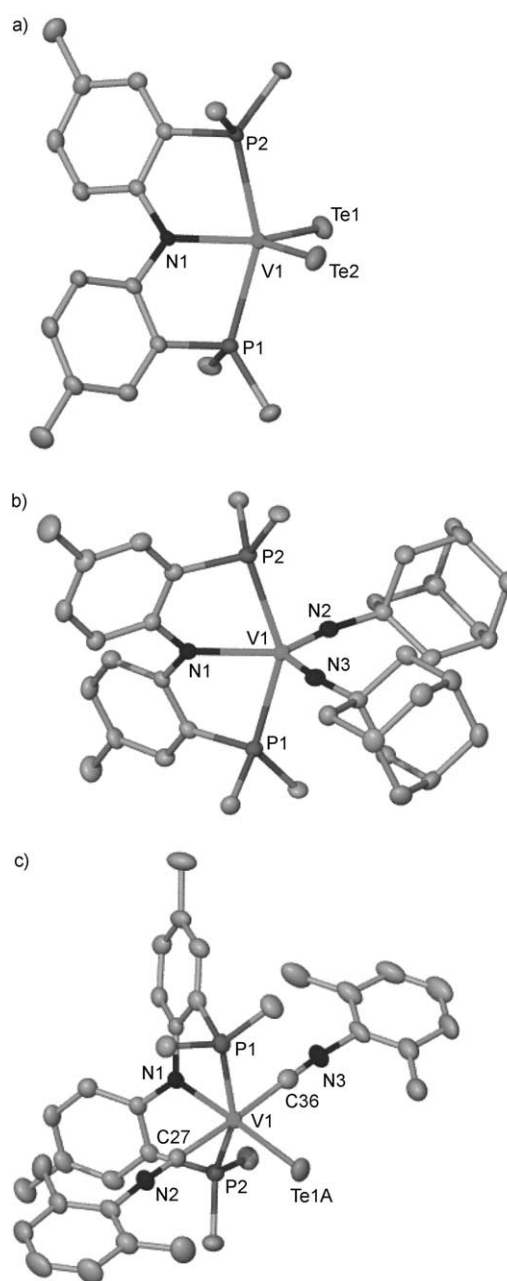
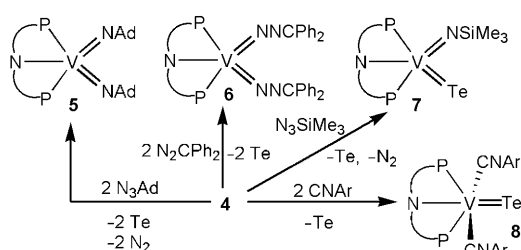


Figure 2. The molecular structures of complexes **4** (a), **5** (b), and **8** (c) with thermal ellipsoids at the 50% probability level. *i*Pr methyl groups on P, solvent, and hydrogen atoms have been excluded for clarity.

molecules).^[7] In addition, the intramolecular Te⋯Te distance in **4** is too long (4.0190(3) Å) for a η²-Te₂²⁻-type ligand bound to a vanadium(III) center. Some intermolecular distances of 4.0190(3) Å between Te atoms are observed in the X-ray crystal structure of **4**. The gross geometry about the metal center in **4** closely resembles other Group 5 complexes that have the same ancillary ligand PNP but contain bisalkylidene moieties composed of CH₂,^[20] CHtBu,^[21] and CHSiMe₃.^[21] It was independently found that complex **4** could be prepared from a one-pot reaction of [(PNP)VCl₂]^[11] with excess Mg and excess Te powder, albeit not in analytically pure form.^[12]

Although complex **4** appears to be resistant to thermal decomposition, the choice of reactant can result in the clean substitution of one or two Te ligands. Accordingly, treatment of **4** with two equivalents of N_3Ad (Ad = 1-adamantyl) immediately results in excision of N_2 with concomitant formation of the complex $[(\text{PNP})\text{V}(\text{NAd})_2]$ (**5**) as yellow crystals in 41 % yield (Scheme 2). The X-ray crystal structure of **5** confirms the existence of a five-coordinate vanadium(V) bisimide complex, which is clearly indicated by the linear V–N–C 159.3(2) and 164.5(2)° angles and short V=N bond lengths of 1.703(3) and 1.696(3) Å (Figure 2).^[22] The V=N metrical parameters are comparable to the few documented monomeric vanadium complexes that have two terminal imide ligands.^[23] Likewise, treatment of **4** with N_2CPh_2 also promotes release of two equivalents of Te by forming the bis(diphenylmethylene hydrazido) complex $[(\text{PNP})\text{V}(\text{N}_2\text{CPh}_2)_2]$ (**6**; Scheme 2). Complex **6** was characterized by a combination of multinuclear NMR spectroscopy (^1H , ^{13}C , ^{31}P , and ^{51}V) and CI mass spectrometry.^[12] Much like complex **5**, room-temperature NMR spectroscopic studies of **6** are indicative of a C_2 -symmetric system. The CI mass spectrum provides further evidence for the proposed structure, whereby N_2 has not been released, with a peak corresponding to the molecular ion at m/z 867.3718.^[12]

Treatment of **4** with one equivalent of N_3SiMe_3 resulted in the substitution of one telluride ligand by the trimethylsilylimide group to form $[(\text{PNP})\text{V}=\text{NSiMe}_3(\text{Te})]$ (**7**) in 32 % yield (Scheme 2).^[12] The addition of excess azide does not promote



Scheme 2. Synthesis of complexes **5–8** from reductive elimination of Te in **4**.

elimination of a second equivalent of Te, even under forcing conditions (80 °C, 12 h). The formation of compound **7** was confirmed by multinuclear NMR spectroscopy (^1H , ^{13}C , ^{31}P , and ^{51}V) and CI MS and, unlike **5** or **6**, the ^1H and ^{13}C NMR spectra are consistent with a C_1 -symmetric molecule because of the substitution of only one Te ligand, coupled with the skewing of the aryl moieties in the PNP ligand. The CI mass spectrum of **7** is also consistent with its structure, and shows the molecular ion peak at m/z 696.1624 as well as a distinctive isotopic mass distribution that arises from the telluride ligand.^[12]

Complex **4** fails to react with a variety of reagents, apart from oxidants, such as $\text{P}(\text{SiMe}_3)_3$, OCNAr (Ar = 2,6- $\text{Me}_2\text{C}_6\text{H}_3$), B_2Pin_2 (Pin = pinacol), benzophenone, 2,3-dimethylbutadiene, and diphenylacetylene. However, treatment of **4** with two equivalents of isonitrile CNAr results in the immediate ejection of metallic Te and the simultaneous

formation of the first V^{III} terminal telluride complex $[(\text{PNP})\text{V}=\text{Te}(\text{CNAr})_2]$ (**8**) as a dichroic (purple–green) material isolated in 50 % yield. The NMR spectra of **8** are in agreement with a diamagnetic vanadium telluride species (^{51}V NMR: δ = 2019 ppm, $\Delta\nu_{1/2}$ = 1581 Hz; ^{125}Te NMR: very broad signal at δ = 3966 ppm), while the IR spectrum of **8** clearly displays two strong ν_{CN} signals (2022 and 2002 cm^{-1}) for the bound isocyanides, which are red-shifted from the ν_{CN} signal of free isocyanide (2116 cm^{-1}). The X-ray crystal structure of **8**^[24] predicates a pseudo octahedral V^{III} species (P1–V1–P2 152.58(5)°; N1–V1–Te1A 177.72(10)°; C36–V1–C27 177.23(19)°), which results from one Te^{2-} ligand in precursor **4** being substituted by two neutral isocyanides, presumably by virtue of a reductive elimination step (Figure 2). The most notable feature in the structure of **8** is the V=Te distance of 2.4603(9) Å, which is consistent with that of a terminal chalcogenide ligand.^[2] In marked contrast, Parkin and Rabinovich have observed isocyanide-induced reductive coupling of two telluride ligands to form an η^2 -ditellurido ligand.^[5,6]

The present work illustrates how the chemistry of the alkylidene and telluride functionalities can ultimately afford rare examples of ditellurethane and tritellurolane, as well as the first vanadium–bis(telluride) complex **4**. Ditellurethanes are exceedingly rare, and our work clearly establishes a method of C=Te bond formation and subsequent dimerization of a transient telluroaldehyde [$\text{Te}=\text{CHtBu}$], as well as dimerization/Te insertion chemistry to form a ring-expanded tritellurolane. We also demonstrate that one or both telluride ligands in precursor **4** can be readily substituted with oxidants such as azides and diphenyldiazomethane. Likewise, reductive elimination of Te^0 can be instigated with isonitriles to afford a rare example of a vanadium(III)–telluride complex. Therefore, our work has established that terminal telluride ligands have the ability to mask high- or low-valent vanadium fragments, presumably because of the weak bond dissociation enthalpy predicted for Te and 3d early-transition metals.^[9]

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