A Fischer-Type Silylene Complex of Platinum: [trans-(Cy₃P)₂(H)Pt=Si(SEt)₂]BPh₄

Steven D. Grumbine and T. Don Tilley*

Department of Chemistry 0506 University of California at San Diego 9500 Gilman Drive La Jolla, California 92093-0506

Frederick P. Arnold and Arnold L. Rheingold*

Department of Chemistry University of Delaware Newark, Delaware 19716

Received May 19, 1993

Since the discovery of transition-metal carbene complexes, $L_n M = CR_2$, in the early 1960s,¹ chemists have pursued possible synthetic routes to the congeneric silicon compounds, L_nM=SiR₂ (silylene complexes). Interest in silylene complexes intensified as it became clear that the carbene derivatives could play important roles in synthesis.² Indeed, silylene complexes have often been invoked as unobserved intermediates in proposed mechanisms.³ However, attempted syntheses of silvlene complexes have met with little success, despite development of routes to well-defined germylene, stannylene, and plumbylene analogs.⁴ Significant advances occurred in 1987, when separate reports described the base-stabilized complexes (CO)₄Fe=Si(O^tBu)₂[OP(NMe₂)₃] and $[Cp^{*}(PMe_{3})_{2}Ru=SiPh_{2}(NCMe)]^{+}BPh_{4}^{-}(Cp^{*} = \eta^{5}-C_{5}^{-}$ Me₅).^{5,6} A number of related adducts containing tetrahedral silicon are now known.7

The marked difficulty encountered in obtaining base-free silvlene complexes raises questions concerning the fundamental nature of metal-silylene bonding interactions and highlights the necessity for defining the structural and electronic properties of these elusive species. The recent isolation of stable, base-free silvlene complexes of the type $Cp^{*}(PMe_{3})_{2}Ru=Si(SR)_{2}^{+8}$ and dynamic studies of their acetonitrile adducts9 suggest that thiolate groups have a stabilizing influence on the silvlene ligand, perhaps

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Scheme I



via 3p-3p π -donation from sulfur to silicon.¹⁰ However, structural information has remained unavailable for these compounds. More recently, we have established the presence of a planar, sp²hybridized silicon center in a complex with two transition metals, $Cp^{*}(PMe_{3})_{2}Ru-Si(STol-p)Os(CO)_{4}$ (Tol = tolyl).¹¹ In search of new transition-metal silvlene complexes which could be structurally characterized, we have turned our attention to platinum derivatives. Such systems should also yield more information on known catalytic processes, since platinum provides many of the most widely used catalysts for transformations of organosilicon compounds.³

The starting - juare-planar platinum silyl complex is obtained by oxidative addition of $HSi(SEt)_3$ to $Pt(PCy_3)_2$ (Cy = cyclohexyl), as shown in Scheme I. The resulting product has cis stereochemistry (by ¹H and ³¹P NMR spectroscopy), but it is quantitatively isomerized to the trans geometry upon photolysis (3 h, 450-W Hg lamp) or heating (over ca. 8 h at 65 °C in benzene- d_6).¹² cis-(Cy₃P)₂Pt(H)Si(SEt)₃ reacts with trimethylsilyl triflate, $Me_3SiOTf(OTf = OSO_2CF_3)$, in a pentane/diethyl ether solution over 3 days to produce the silyl complex trans-(Cy₃P)₂-Pt(H)Si(SEt)₂OTf in 57% yield. The latter triflate derivative has inequivalent methylene protons that appear in the ¹H NMR spectrum at room temperature as overlapping multiplets centered at δ 3.30 (benzene- d_6). These resonances coalesce at 40 °C, probably by a mechanism involving dissociation of the triflate anion to transiently produce the silylene complex $[(Cy_3P)_2(H)Pt=Si(SEt)_2]^+OTf^-$, in which the methylene hydrogens are equivalent. Related behavior has been observed for the ruthenium complex Cp*(PMe₃)₂RuSi(SEt)₂OTf, for which the diastereotopic methylene hydrogens coalesce at 21 °C (toluened₈ solution).⁸ The labile triflate group in trans-(Cy₃P)₂Pt(H)- $Si(SEt)_2OTf$ is displaced in benzene- d_6 by the neutral two-electron donor (dimethylamino)pyridine (DMAP), to afford the donor-

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(12) Selected data are as follows. cis-(PCy₃)₂(H)PtSi(SEt)₃: ³¹Pf¹H}NMR (benzene-d₆, 121.5 MHz, 23 °C) δ 39.31 (mult with ¹⁹⁵Pt satellites as dd for each P, J_{PPt} = 6454 Hz, J_{PP} = 33 Hz, and J_{PPt} = 4409 Hz, J_{PP} = 33 Hz), IR (Nujol, CsI, cm⁻¹) 2088 m (PtH). trans-(PCy₃)₂(H)PtSi(SEt)₃: ³¹P{¹H}NMR (benzene- d_6 , 121.5 MHz, 23 °C) δ 39.61 (s with ¹⁹⁵Pt satellites, $J_{PPt} = 2646$ Hz); IR (Nujol, CsI, cm⁻¹), 2010 w (PtH). trans-(PCy₃)₂(H)PtSi(SEt)₂-The first of the set z_2 , z_1 , z_2 , z_2 , z_3 , z_5 , z_5 , z_5 , z_5 , z_5 , z_5 , z_7 , Si(SEt)₂][BPh₄]: 31 P{¹H} NMR (dichloromethane- d_2 , 121.5 MHz, 23 °C) δ 51.51 (s with 195 Pt satellites, $J_{PP1} = 2279$ Hz), IR (dichloromethane, KBr, cm-1), 2010 w (PtH).

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Figure 1. Structure of the cation in $[(Cy_3P)_2(H)Pt=Si(SEt)_2]BPh_4-CH_2-$ Cl₂.

stabilized silvlene complex [trans-(Cy₃P)₂(H)Pt=Si(SEt)₂-(DMAP)]OTf as the kinetic product. During crystallization of this compound from dichloromethane/diethyl ether, rearrangement to the cis isomer occurred (Scheme I).

The title compound was formed in the reaction of trans-(Cy₃P)₂-Pt(H)Si(SEt)₂OTf with NaBPh₄ in dichloromethane (Scheme I). Precipitation of NaOTf produces a solution of the cationic silvlene complex, which was crystallized from dichloromethane/ diethyl ether in 56% yield. NMR spectra for the compound are consistent with the structure shown in Scheme I, with trans hydride and silylene ligands. Of most interest is the ²⁹Si chemical shift of 308.65 ppm, which is dramatically downfield-shifted from that for the triflate trans-(Cy₃P)₂Pt(H)Si(SEt)₂OTf, 89.69 ppm. Such low-field chemical shifts are found for the previously reported base-free silvlene complexes $Cp^{*}(PMe_{3})_{2}Ru=Si(SR)_{2}^{+}$, and lowfield ¹³C shifts are also associated with related carbene complexes.² The ${}^{1}J_{PtSi}$ coupling contant for the silylene complex (1558 Hz) is lower than the ${}^{1}J_{PtSi}$ value for the triflate derivative (1825 Hz), implying that the Si(SEt)₂OTf ligand, being a better σ -dono., interacts more with the Pt 6s orbital than does Si(SEt)2.13

The structure of [trans-(Cy₃P)₂(H)Pt=Si(SEt)₂]⁺, shown in Figure 1, reveals a square-planar coordination geometry for platinum.14 There are no short interionic or ion-solvent interactions in the solid state. Planarity of the silylene fragment (Si is only 0.04 Å out of the PtS_2 plane) and the summation of angles at silicon (359.9°) reflect sp² hybridization.¹⁵ The plane of the silvlene ligand is rotated 76° out of the least-squares plane of platinum donor atoms, implying that π -donation from the d_{xy} orbital, rather than from the somewhat lower-energy d_{xz} level, contributes more to the molecular bonding. This "perpendicular" orientation is observed for most square-planar platinum carbene complexes (exceptions occur when the carbene group is constrained as part of a metallacycle).¹⁶ As in the latter cases, this geometry appears to result largely from steric interactions between the silylene substituents and bulky phosphine ligands.^{16,17} Scheme II gives the principal resonance structures for [trans-(Cy₃P)₂-(H)PtSi(SEt)₂]+.

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Scheme II



The Pt-Si bond length, 2.270(2) Å, is short in comparison with those in related complexes such as trans-(Cy₃P)₂Pt(H)Si- $(SEt)_3 (2.379(1) \text{ Å})^9$ and trans- $(Cy_3P)_2Pt(H)SiH_3 (2.382(3) \text{ Å}).^{18}$ The Si-S distances (2.092(4) and 2.074(4) Å) are also short, since Si-S bonds in Cp*(PMe₃)₂Ru-Si(SR)XY complexes generally fall in the range 2.17-2.20 Å, 6,9,11 and the Si-S distances in trans-(Cy₃P)₂Pt(H)Si(SEt)₃ average to 2.17 Å. The Si-S bond shortening observed for the silvlene complex may be ascribed to the lower coordination number for silicon and some degree of π -bonding. The presence of some π -bonding is suggested by acute dihedral angles between the Si-S-C and SiS₂ planes of 9.9 and 10.7°. This results in roughly cis and trans orientations for the ethyl groups relative to the Pt atom in the solid-state structure, as depicted in Scheme II. However, since the integrity of these orientations is not upheld in dichloromethane solution down to -80 °C, there must be rapid rotation about the Si-S bonds. This contrasts with observations for analogous platinum carbenes, such as $(Ph_3P)_2(I)Pt=C(SMe)_2$, for which the rotational barrier about the C-S bonds is high enough for distinct rotamers to be observed at room temperature.¹⁹

Preliminary computational results based on the Fenske-Hall method²⁰ suggest that the low-coordinate silicon atom is stabilized more by π -donation from the sulfur atoms than by donation from platinum. While significant overlap exists between the silylene p orbital and orbitals of appropriate symmetry on the platinum and sulfur centers, only Si-S (and not Si-Pt) π -bonds are populated. An overlap population of 0.43 electrons per sulfur atom, or 0.86 electrons total, exists between the unhybridized silylene p orbital and the sulfur lone pairs. The computational results therefore suggest that the structure is best described as containing a low-valent silicon center stabilized primarily by π -interactions with the sulfur lone pairs and much less by π -bonding to platinum. In order to more quantitatively describe the bonding of this system, a correlated method such as MCSCF will be necessary.²¹ These calculations are in progress.

Initial reactivity studies show that [trans-(Cy₃P)₂(H)Pt= Si(SEt)₂]⁺BPh₄⁻ does not bind acetonitrile in dichloromethane solution but does bind the stronger base DMAP. We are currently focusing on defining the chemical properties of this complex as they pertain to known and potential catalytic cycles.

Acknowledgment is made to the National Science Foundation for financial support, and we thank Johnson Matthey Inc. for a loan of K_2 PtCl₄. We would also like to thank Dr. Thomas Albright for his helpful discussions.

Supplementary Material Available: Experimental procedures and characterization data for complexes, tables of crystal, data collection, and refinement parameters, bond distances and angles, anisotropic displacement parameters, and hydrogen atom coordinates (17 pages); listings of observed and calculated structure factors (24 pages). Ordering information is given on any current masthead page.

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