

Iridium-Catalyzed Redistribution of Hydrodisilanes via a Silyl(silylene)iridium(III) Complex: Synthesis of a Donor-Stabilized Silyl(silylene)iridium(III) Complex

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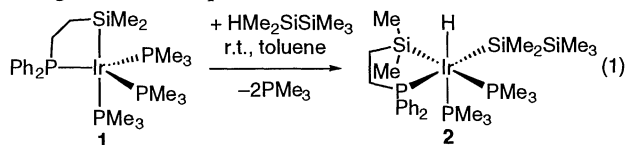
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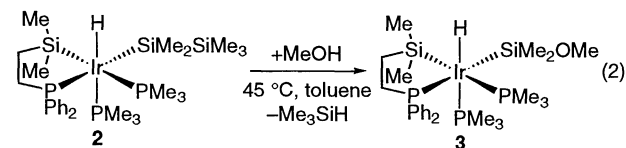
Thermolysis of HMePhSiSiMe_3 in the presence of a catalytic amount of $\text{Ir}\{\eta^2\text{-Me}_2\text{Si}(\text{CH}_2)_2\text{PPh}_2\}(\text{PMe}_3)_3$ resulted in the redistribution of substituents to give its isomer $\text{HMe}_2\text{SiSiMe}_2\text{Ph}$. We propose the mechanism involving the 1,3-Me-shift on a silyl(silylene) complex, which was trapped as a donor-stabilized silyl(silylene)iridium(III) complex.

A silyl(silylene) complex has been considered as a key intermediate in transition-metal-mediated redistribution reactions of organosilicon compounds.¹ It was found that photo-chemical conversion of $\text{CpFe}(\text{CO})_2\text{SiMe}_2\text{SiMeR}_2$ ($\text{R} = \text{Ph, Et, CD}_3$) to $\text{CpFe}(\text{CO})_2\text{SiMe}_{3-n}\text{R}_n$ ($n = 0, 1, 2$) is accompanied by the scrambling of substituents on silicon atoms, and Pannell et al. and we proposed a mechanism involving the 1,3-migration of substituents on silyl(silylene) intermediates.² Recently, Pannell et al. reported that thermolysis of hydrodisilanes with a catalytic amount of $(\eta^5\text{-C}_5\text{H}_5)\text{Fe}(\text{CO})(\text{PPh}_3)(\text{SiMe}_3)$ caused isomerization of disilanes, where they proposed a mechanism involving a silyl(silylene)iron intermediate.³ Tamao et al. described a $\text{Pd}(0)$ -catalyzed skeletal rearrangement of alkoxytrisilanes, and they proposed the mechanism involving a silyl(silylene)palladium(II) intermediate, which is stabilized by an intramolecular donor.⁴ In these catalytic reactions, however, nothing has been explored on the isolation of the intermediate, i.e., the silyl(silylene) complex or its stabilized form. We report here an $\text{Ir}(\text{I})$ complex $\text{Ir}\{\eta^2\text{-Me}_2\text{Si}(\text{CH}_2)_2\text{PPh}_2\}(\text{PMe}_3)_3$ (**1**)⁵ not only catalyzes the redistribution of hydrodisilanes but also forms a donor-stabilized bis(silylene)iridium complex, i.e., the stabilized form of the silyl(silylene) complex.

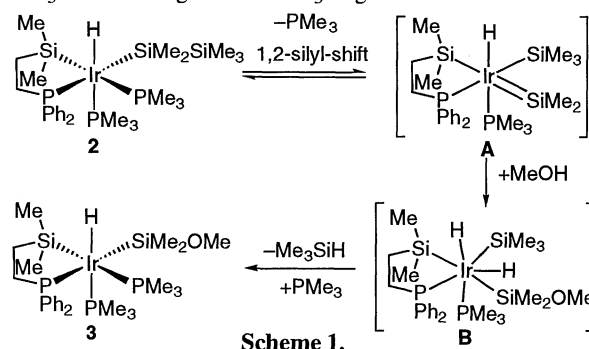
Treatment of **1** with 1 equiv. $\text{HSiMe}_2\text{SiMe}_3$ at room temperature gave a hydrido(disilanyl)iridium(III) complex $\text{Ir}(\text{H})(\text{SiMe}_2\text{SiMe}_3)\{\eta^2\text{-Me}_2\text{Si}(\text{CH}_2)_2\text{PPh}_2\}(\text{PMe}_3)_2$ (**2**) (eq 1).⁶ Recrystallization of **2** from toluene-hexane afforded colorless crystals of **2** in 89% yield. The ^1H , ^{29}Si , and ^{31}P NMR data established that **2** possesses three phosphorous atoms in a *fac*-configuration as in eq 1.



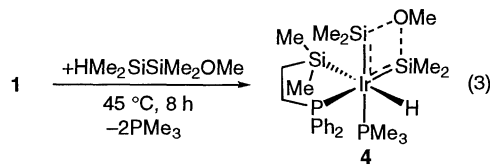
In the presence of MeOH , **2** was converted, on heating to 45°C for 2 h, to $\text{Ir}(\text{H})(\text{SiMe}_2\text{SiMe}_2\text{OMe})\{\eta^2\text{-Me}_2\text{Si}(\text{CH}_2)_2\text{PPh}_2\}(\text{PMe}_3)_2$ (**3**)⁷ in 73% isolated yield (eq 2),⁸ although **2** was thermally stable at this temperature in the absence of MeOH . In the ^1H NMR spectrum, a signal of SiOMe appears as a singlet at 3.45 ppm.



A plausible mechanism is illustrated in Scheme 1: The dissociation of a PMe_3 ligand followed by the reversible 1,2-silyl-shift gives a silyl(silylene) intermediate **A**. MeOH is known to work as an efficient trapping agent of silylene complex to give a hydrido(methoxysilyl) complex.⁹ Therefore, MeOH adds to the $\text{Ir}=\text{Si}$ double bond to give a seven-coordinate iridium(V) intermediate **B**, which then undergoes the reductive elimination of Me_3SiH ¹⁰ and ligation of PMe_3 to give **3**.

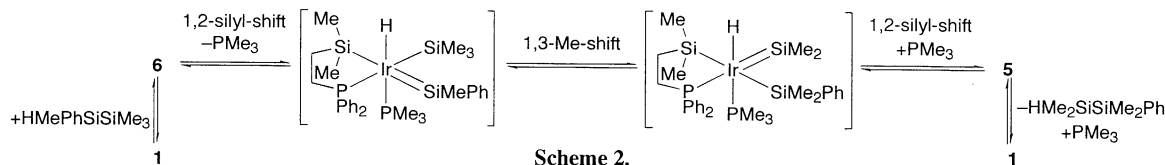


To confirm the existence of a silyl-silylene intermediate **A**, complex **1** was allowed to react with $\text{HMe}_2\text{SiSiMe}_2\text{OMe}$. The reaction proceeded cleanly at 45°C for 8 h to give hydridobis(silylene) complex **4** (eq 3).^{11,12} The geometry of **4** can be uniquely determined by the NMR data. The ^{31}P NMR spectrum shows an AX pattern at -60.6 ppm and 32.0 ppm ($J(\text{PP}_{\text{cis}}) = 28.9$ Hz), which is consistent with the geometry as shown in eq 3. In the ^{29}Si NMR spectrum, the signals of two silylene ligands appeared inequivalently at 62.8 ppm (dd, $J(\text{SiP}_{\text{trans}}) = 135.9$ Hz, $J(\text{SiP}_{\text{cis}}) = 11.8$ Hz) and 63.1 ppm (dd, $J(\text{SiP}_{\text{trans}}) = 136.9$ Hz, $J(\text{SiP}_{\text{cis}}) = 13.0$ Hz), which are shifted to significantly downfield from those of previously reported silyliridium compounds.^{5,9b,13} Moreover, the upfield shift of the ^1H NMR signal for the SiOMe group (2.77 ppm) is characteristic of methoxy-bridged bis(silylene) complexes.¹⁴

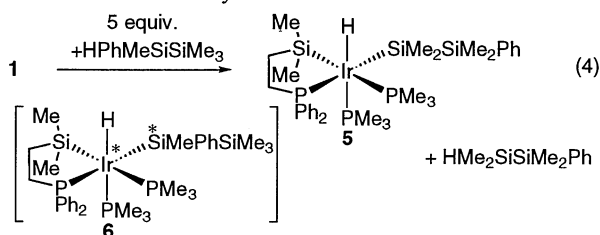


The formation of **4** can be also explained by the mechanism involving the 1,2-silyl-shift to the iridium center as shown in Scheme 1. In the case of $\text{HMe}_2\text{SiSiMe}_2\text{OMe}$, a silyl(silylene) intermediate corresponding to **A** can be stabilized by the bridging methoxy group to give a bis(silylene) complex **4**.

Reaction of **1** with 5 equiv. HPhMeSiSiMe_3 was carried out at room temperature for 2 h. Complex **1** reacted with 1 equiv. HPhMeSiSiMe_3 to give **5** via the rearrangement of substituents on a disilanyl ligand (eq 4).¹⁵ A small amount of $\text{HMe}_2\text{SiSiMe}_2\text{Ph}$ was also detected spectroscopically. The formation of **6**, a simple oxidative addition product, was



confirmed in the course of the reaction, but **6** finally disappeared.¹⁶ Furthermore, thermolysis of this solution at 45 °C underwent the rearrangement of hydrodisilane to give an isomeric mixture of HPhMeSiSiMe₃ and HMe₂SiSiMe₂Ph. After 5 days, the molar ratio of HPhMeSiSiMe₃ to HMe₂SiSiMe₂Ph became 2 : 3, and **5** finally decomposed.¹⁷ We treated HMe₂SiSiMe₂Ph under the same conditions in the presence of **1**, which resulted in the formation of a mixture of HPhMeSiSiMe₃ and HMe₂SiSiMe₂Ph in the ratio of 2 : 5. The decomposition of **5** formed in the reaction was also confirmed spectroscopically.¹⁷ Each reaction led to the isomeric mixture of hydrodisilane in the different ratio. This means that an active catalyst **5**¹⁸ decomposed before achieving the equilibrium.¹⁷ This isomerization reactions can be explained by the mechanism shown in Scheme 2. This mechanism also involves the generation of a silyl(silylene) intermediate, which causes a 1,3-Me-shift and 1,2-silyl-shift. The resulting disilanyl complex eliminates an isomeric hydrodisilane.



To detect the silyl-silylene intermediate in Scheme 2, we employed HMe(MeO)SiSiMe₃ instead of HPhMeSiSiMe₃. Thermolysis of **1** in the presence of HMe(MeO)SiSiMe₃ led to the clean formation of **4**. This apparently demonstrates that the skeletal rearrangement of hydrodisilanes take place via the silyl(silylene) intermediate, which causes the 1,3-Me-shift as shown in Scheme 2.

In this paper, we firstly succeeded in the isolation of a donor-stabilized silyl(silylene) species, which is a stabilized form of a key intermediate formed in the transition-metal-mediated redistribution of substituents of silicon atoms.

References and Notes

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- Selected data for **2**: ¹H NMR (300 MHz, C₆D₆) δ -12.78 (dt, *J*(HPtrans) = 105.1 Hz, *J*(HPcis) = 17.1 Hz, 1H, IrH). ²⁹Si NMR (59.6 MHz, C₆D₆) δ -51.3 (ddd, *J*(SiPtrans) = 115.6 Hz, *J*(SiPcis) = 11.4, 12.7 Hz, IrSiMe₂SiMe₃), -13.6 (t, *J*(SiPcis) = 11.4 Hz, SiMe₃), 10.5 (ddd, *J*(SiPtrans) = 116.2 Hz, *J*(SiPcis) = 11.5, 7.7 Hz, IrSiMe₂CH₂). ³¹P NMR (121.5 MHz, C₆D₆) δ -69.4 (t, *J*(PPcis) = 23.7 Hz, PMe₃), -61.4 (dd, *J*(PPcis) = 23.7, 17.3 Hz, PMe₃), 27.6 (dd, *J*(PPcis) = 17.3, 23.7 Hz, PPh₂). Anal. Found: C, 43.29; H, 7.00%. Calcd for C₂₇H₅₄IrP₃Si₃: C, 43.35; H, 7.28%.
- Selected data for **3**: ¹H NMR (300 MHz, C₆D₆) δ -12.82 (dt, *J*(HPtrans) = 99.0 Hz, *J*(HPcis) = 18.0 Hz, 1H, IrH), 3.45 (s, 3H, OMe). ³¹P NMR (121.5 MHz, C₆D₆) δ -65.5 (dd, *J*(PPcis) = 23.1, 25.5 Hz, PMe₃), -59.9 (dd, *J*(PPcis) = 20.2, 25.5 Hz, PMe₃), 25.6 (dd, *J*(PPcis) = 20.2, 23.1 Hz, PPh₂). Anal. Found: C, 42.94; H, 7.01%. Calcd for C₂₅H₄₈IrOP₃Si₂: C, 42.53; H, 6.85%.
- Formation of Me₃SiH was confirmed by ¹H NMR spectroscopy. ¹H NMR data of Me₃SiH (300 MHz, C₆D₆) δ 4.16 (sep., *J*(HH) = 3.6 Hz, 1H, SiH), 0.00 (d, 9H, SiMe₃).
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- There is another possibility that **B** eliminates HSiMe₂OMe instead of HSiMe₃, although the formation of HSiMe₂OMe was not observed in the ¹H NMR spectrum. This is attributable to the existence of the electron-withdrawing methoxy group on the silicon atom. An electron-withdrawing group on a silicon atom is known to strengthen a M-Si bond by increasing M-Si π-bonding involving Si d or σ* orbital.
- Selected data for **4**: ¹H NMR (300 MHz, C₆D₆) δ -9.88 (dd, *J*(HPcis) = 14.3, 19.7 Hz, 1H, IrH), 2.77 (s, 3H, SiOMe). ²⁹Si NMR (59.6 MHz, C₆D₆) δ 12.5 (dd, *J*(SiPcis) = 9.2, 11.7 Hz, silyl), 62.8 (dd, *J*(SiPtrans) = 135.9 Hz, *J*(SiPcis) = 11.8 Hz, silylene), 63.1 (dd, *J*(SiPtrans) = 136.9 Hz, *J*(SiPcis) = 13.0 Hz, silylene). ³¹P NMR (121.5 MHz, C₆D₆) δ -60.6 (d, *J*(PPcis) = 28.9 Hz, PMe₃), 32.0 (d, PPh₂). Exact mass (70 eV, DEI) *m/z* Calcd for C₂₄H₄₅IrOP₂Si₃: 688.1883. Found: 688.1889.
- At room temperature, the reaction of **1** and HMe₂SiSiMe₂OMe gave Ir(H)(SiMe₂SiMe₂OMe)(η²-Me₂Si(CH₂)₂PPh₂)(PMe₃)₃ corresponding to **2**. Spectroscopic data are similar to those for **2**.
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- a) H. Tobita, K. Ueno, M. Shimoi, and H. Ogino, *J. Am. Chem. Soc.*, **112**, 3415 (1990). b) H. Tobita, H. Wada, K. Ueno, and H. Ogino, *Organometallics*, **13**, 2545 (1994). c) T. Takeuchi, H. Tobita, and H. Ogino, *Organometallics*, **10**, 835 (1991).
- Selected data for **5**: ¹H NMR (300 MHz, C₆D₆) δ -12.79 (dt, *J*(HPtrans) = 104.4 Hz, *J*(HPcis) = 16.7 Hz, 1H, IrH). ²⁹Si NMR (59.6 MHz, C₆D₆) δ -51.2 (ddd, *J*(SiPtrans) = 119.0 Hz, *J*(SiPcis) = 11.3, 12.8 Hz, IrSiMe₂SiMe₂Ph), -16.0 (t, *J*(SiPcis) = 11.8 Hz, SiMe₂Ph), 10.3 (ddd, *J*(SiPtrans) = 115.6 Hz, *J*(SiPcis) = 7.5, 11.4 Hz, IrSiMe₂CH₂). ³¹P NMR (121.5 MHz, C₆D₆) δ -69.6 (t, *J*(PPcis) = 23.8 Hz, PMe₃), -61.9 (dd, *J*(PPcis) = 17.1, 23.8 Hz, PMe₃), 27.5 (dd, *J*(PPcis) = 17.1, 23.8 Hz, PPh₂). Anal. Found: C, 47.20; H, 6.93%. Calcd for C₃₂H₅₆IrP₃Si₃: C, 47.44; H, 6.97%.
- Complex **6** is a mixture of diastereomers **6a** and **6b**. ³¹P NMR (121.5 MHz, C₆D₆) data for **6a**: δ 28.1 (dd, *J*(PPcis) = 16.3 Hz, 24.2 Hz, PPh₂), -63.5 (dd, *J*(PPcis) = 16.3, 24.2 Hz, PMe₃), -72.6 (t, *J*(PPcis) = 24.2 Hz, PMe₃). **6b**: 29.4 (dd, *J*(PPcis) = 17.1, 23.2 Hz, PPh₂), -62.5 (dd, *J*(PPcis) = 17.1, 23.2 Hz, PMe₃), -71.8 (t, *J*(PPcis) = 23.2 Hz, PMe₃).
- Decomposition of **5** may be attributable to the reaction with a small amount of H₂O or thermal evolution of the SiR₂ moiety. The decomposition resulted in the formation of *fac*-[Ir(H)(SiMe₃)(η²-Me₂Si(CH₂)₂PPh₂)(PMe₃)₂], *fac*-[Ir(H)(SiMe₂Ph)(η²-Me₂Si(CH₂)₂-PPh₂)(PMe₃)₂], and some unidentified products.
- In the presence of an isolated **5**, the catalytic isomerization of the hydrodisilane, HPhMeSiSiMe₃ or HMe₂SiSiMe₂Ph, also took place in a similar manner. This means that **5** is one of active catalytic species.