



From Silylone to an Isolable Monomeric Silicon Disulfide Complex**

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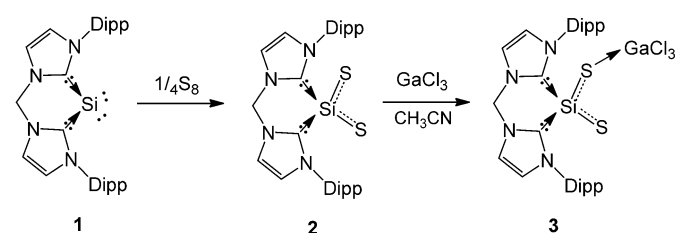
Dedicated to Professor Manfred Scheer on the occasion of his 60th birthday

Abstract: The synthesis and characterization of the first bis-N-heterocyclic carbene stabilized monomeric silicon disulfide (bis-NHC)SiS₂ **2** (bis-NHC = H₂C[NC(H)C(H)N(Dipp)]C:]₂, Dipp = 2,6-*i*Pr₂C₆H₃) is reported. Compound **2** is prepared in 89 % yield from the reaction of the zero-valent silicon complex ('silylone') **1** [(bis-NHC)Si] with elemental sulfur. Compound **2** can react with GaCl₃ in acetonitrile to give the corresponding (bis-NHC)Si(S)S → GaCl₃ Lewis acid–base adduct **3** in 91 % yield. Compound **3** is also accessible through the reaction of the unprecedented silylone–GaCl₃ adduct [(bis-NHC)Si → GaCl₃] **4** with elemental sulfur. Compounds **2**, **3**, and **4** could be isolated and characterized by elemental analyses, HR-MS, IR, ¹³C- and ²⁹Si-NMR spectroscopy. The structures of **3** and **4** could be determined by single-crystal X-ray diffraction analyses. DFT-derived bonding analyses of **2** and **3** exhibited highly polar Si–S bonds with moderate p_π–p_π bonding character.

Silicon disulfide SiS₂ is a textbook compound which was first prepared from elemental silicon and sulfur by Berzelius in 1824.^[1] Under normal conditions SiS₂ adopts a polymeric structure owing to the extremely high polarity of the Si–S bond.^[2] Monomeric SiS₂ (S=Si=S) is a heavier analogue of CO₂ and CS₂ but far more reactive, and it undergoes spontaneous polymerization as a result of the relatively weak p_π–p_π bond between the silicon and sulfur atoms. Since monomeric SiS₂ is metastable its isolation requires the utilization of particular experimental techniques and methods. Schnöckel and co-workers showed that molecular SiS₂ can be synthesized and isolated under matrix isolation conditions at very low temperatures and studied spectroscopically.^[3] In an alternative approach, the isolation of monomeric SiS₂ even at room temperature could be envisaged by taking advantage of the concept of donor–acceptor stabilization and

formation of a L → SiS₂ complex with L acting as a donor ligand towards the strongly electrophilic silicon atom in S=Si=S. To our knowledge, such a complex is currently unknown.

The concept of kinetic and/or thermodynamic stabilization has enabled great achievements in the chemistry of low-coordinate silicon compounds as shown through the isolation and structural characterization of disilenes, Ar₂Si=SiAr₂ (Ar = 2,4,6-Me₃C₆H₂), by West et al.^[4] Remarkable examples also include the isolation of stable silylenes,^[5] disilynes,^[6] and other interesting low-coordinate silicon species.^[7a–c] Moreover, a striking N-heterocyclic carbene (NHC) supported disilicon(0) complex could be synthesized by Robinson et al. by taking advantage of the concept of donor–acceptor stabilization.^[7d] Unusual silicon species featuring elusive terminal Si=O and Si=S moieties^[4a,8] and E=Si=E bonds (M = C, Si, Ge)^[9] could as well be stabilized using Lewis donors. Very recently, Roesky et al. and our group successfully isolated the first zero-valent silicon complexes ('silylones') by utilizing two cyclic alkyl(amino) carbenes (cAACs)^[10a] and the bis-NHC chelating ligand (bis-NHC = H₂C[NC(H)C(H)N(Dipp)]C:]₂, Dipp = 2,6-*i*Pr₂C₆H₃),^[10b] respectively. Owing to the stronger σ-donor but weaker π-acceptor character of the bis-NHC ligand, the silylone **1** bearing two lone pairs of electrons on the silicon atom (Scheme 1) is more electron-rich than Roesky's (cAAC)₂Si as



Scheme 1. Formation of the SiS₂ complex **2** and its GaCl₃ adduct **3**, starting from the bis-NHC stabilized silylone **1** (Dipp = 2,6-*i*Pr₂C₆H₃).

confirmed by DFT calculations. The presence of an electron-rich silicon(0) atom in the pocket of the sterically protecting bis-NHC ligand of **1** prompted us to investigate whether the monomeric silicon disulfide complex **2** could be synthesized by direct oxidation of **1** with elemental sulfur. Herein we report the synthesis and characterization of the first SiS₂ complex (bis-NHC)SiS₂ **2** and its GaCl₃ adduct **3**, (bis-NHC)Si(S)S → GaCl₃, starting from silylone **1** (Scheme 1).

Treatment of a dark red solution of **1** in THF with 0.25 molar equivalents of elemental sulfur (S₈) at room temperature leads immediately to a color change of the

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reaction mixture to pale orange, and subsequent formation of a colorless precipitate (Scheme 1). Filtration and washing of the precipitate with small amounts of THF affords the desired (bis-NHC)SiS₂ compound **2** as a colorless powder in 89% yield. The compound is insoluble in hydrocarbons and ethereal solvents. Its composition has been confirmed by correct elemental analysis and high-resolution (HR) electrospray ionization (ESI) mass spectrometry (MS; see Supporting Information). The HR-ESI-MS spectrum of **2** shows the molecular ion peak at *m/z* 561.25220 (calcd 561.25364). The solid state CP/MAS ²⁹Si{¹H} NMR spectrum of **2** displays a signal at $\delta = -32.5$ ppm which is in good agreement with the value of $\delta = -29.8$ ppm predicted by DFT calculations (Table 1). The experimental ²⁹Si{¹H} NMR chemical shift is

Table 1: Calculated ²⁹Si NMR chemical shifts (in ppm vs. TMS; experimental values are given in parentheses),^[a] NPA charges (*q*),^[b] Wiberg bond indices (WBI),^[b] and Mayer bond orders (MBO)^[b] for compounds **2'**, **3'**, and **4'**.

Parameter	2'	3'	4'
$\delta(^{29}\text{Si})$	-29.8 (-32.5)	-42.0 (-40.0)	-131.3 (-119.0)
<i>q</i> (Si)	+1.14	+1.17	+0.17
<i>q</i> (S _{ax/1})	-0.85	-0.69	—
<i>q</i> (S _{eq/2})	-0.84	-0.77	—
<i>q</i> (GaCl ₃)	—	-0.46	-0.73
WBI(Si-S _{ax/1})	1.19	0.97	—
WBI(Si-S _{eq/2})	1.34	1.41	—
MBO(S-S _{ax/1})	1.65	1.24	—
MBO(Si-S _{eq/2})	1.75	1.86	—

[a] B3LYP/IGLO-III(H,C,Si,N,S,Cl)/def2-TZVPP(Ga)//B3LYP-D3(BJ)/def2-TZVPP results. [b] B3LYP/def2-TZVPP results. As expected, owing to the different definitions of bond orders, MBOs exhibit a larger overall weight of covalent σ -bonding than WBIs, but the same trends when going from **2'** to **3'**.

upfield from the value of $\delta = -19.5$ ppm observed for polymeric and polycrystalline SiS₂.^[11] However, it is very close to the chemical shifts of $\delta = -33.5$ and -34.9 ppm observed for the NHC-stabilized Si=S complexes L(NHC)SiS (L = N(Dipp)C(=CH₂)CH=C(Me)N(Dipp)) with four-coordinate silicon centers.^[12a]

Compound **2** is almost insoluble in common organic solvents, presumably due to the extremely high polarity of the two Si=S moieties. Thus attempts to grow single crystals of **2** for X-ray diffraction analysis failed. However, complexation of **2** with GaCl₃ in acetonitrile at ambient temperature leads to the formation of the soluble (bis-NHC)Si(S)S→GaCl₃ Lewis acid–base adduct **3** in 91% yield after work-up. Single crystals of **3** suitable for X-ray diffraction analysis were obtained in a THF solution at -30°C . Complex **3** crystallized in the monoclinic space group *Cm*. The analysis revealed the presence of a bis-NHC-ligand-supported monomeric silicon disulfide molecule which is coordinated to the GaCl₃ Lewis acid by a S→Ga bond (Figure 1). Accordingly, the Si atom adopts a distorted tetrahedral geometry and is in a slightly puckered six-membered C₃N₂Si ring with the chelating bis-NHC ligand. The C1–Si1–C1' angle of $92.5(2)^\circ$ is slightly larger than the corresponding angle in silylone **1** ($89.1(2)^\circ$).^[10b] The Si–C distances of $1.930(4)$ Å are elon-

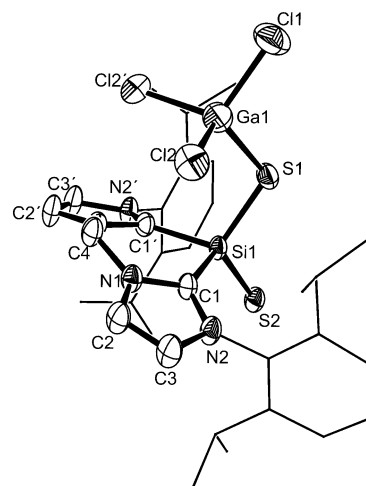
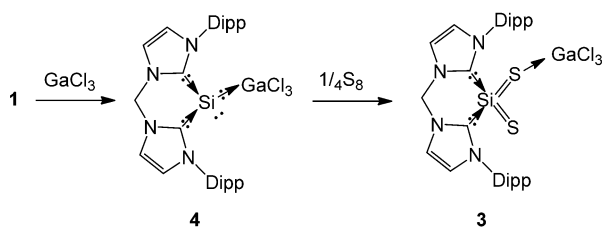


Figure 1. Molecular structure of **3**. Thermal ellipsoids are set at 50% probability. H atoms and one THF lattice solvent molecule are omitted for clarity. Symmetry transformations used to generate equivalent atoms with ('): *x*, $-y$, *z*. Selected interatomic distances [Å] and angles [$^\circ$]: Si1–S1 2.106(2), Ga1–S1 2.262(2), Si1–S2 2.006(2), Ga1–Cl1 2.169(2), Ga1–Cl2 2.190(1), Si1–C1 1.930(4), N1–C1 1.352(4), N1–C2 1.378(5), N1–C4 1.454(4), C1–N2 1.338(5), N2–C3 1.396(5), C2–C3 1.351(6); Si1–S1–Ga1 109.32(7), C1–Si1–C1' $92.5(2)$, C1–Si1–S2 $113.3(1)$, C1–Si1–S1 $110.3(1)$, S2–Si1–S1 $115.03(8)$, Cl1–Ga1–Cl2 $109.28(5)$, Cl2–Ga1–Cl2' $106.97(7)$, Cl1–Ga1–S1 $107.86(8)$, Cl2–Ga1–S1 $111.71(4)$.

gated compared with the value observed for **1** (1.869 Å). As expected, the two Si–S distances are significantly different owing to the coordination of the GaCl₃ moiety to the S1 atom. While the Si1–S2 bond length of $2.006(2)$ Å is identical to the value of $2.006(1)$ Å found for the Si=S bond in L(NHC)Si=S mentioned above,^[12a] the Si1–S1 bond of $2.106(2)$ Å is only slightly shorter than the Si–S single bond in [PhC(NtBu)₂]Si(S)SrBu ($2.131(7)$ Å).^[12b] In addition, the S–Si–S angle of $115.03(8)^\circ$ in **3** deviates pronouncedly from the value of 180° for a linear S=Si=S structure of “free” SiS₂, apparently because of the coordination of the bis-NHC ligand to the highly electrophilic silicon center.

Complex **3** is insoluble in hexane, toluene, and diethyl ether, but sparingly soluble in THF and acetonitrile which enabled us to record its ¹H NMR spectrum in CD₃CN solutions (see Supporting Information). In the HR-ESI-MS spectrum of **3**, only the molecular ion peak of **2** at *m/z* 561.25470 (calcd 561.25364) could be observed. The solid-state CP/MAS ²⁹Si{¹H} NMR spectrum of **3** shows a signal at $\delta = -40.0$ ppm which is upfield shifted compared with **2** ($\delta = -32.5$ ppm). The change of the ²⁹Si shift of **3** is somewhat unexpected because GaCl₃ coordination withdraws charge from the silicon center (cf. more positive Si NPA charge in Table 1). Analysis of molecular-orbital (MO) contributions in this context (see below) show that changes in the ²⁹Si shielding are dominated by Si–S σ -bonding MOs (these have far larger MO coefficients at Si than the very polar Si–S π -bonding MOs; cf. Supporting Information for details), resulting in an overall stronger shielding of silicon in **3**.

Akin to **2**, the electron-rich zero-valent Si atom in silylone **1** is able to form a Lewis acid–base adduct with GaCl₃. Accordingly, treating a THF solution of **1** with one molar



Scheme 2. Formation of the silylone- GaCl_3 adduct **4**, starting from silylone **1** and GaCl_3 , and its direct oxidation with elemental sulfur to **3**.

equivalent of GaCl_3 results in the formation of the (bis-NHC) $\text{Si} \rightarrow \text{GaCl}_3$ complex **4**, which precipitates from the reaction solution as a yellow solid and can be isolated in 58 % yield (Scheme 2). Even with molar excess of GaCl_3 , only compound **4** is formed (^1H NMR). Compound **4** is the first Lewis acid-base adduct of a silylone. It crystallized as yellow blocks in the monoclinic space group $P2_1/c$ in acetonitrile solutions (Figure 2). X-ray diffraction analysis revealed

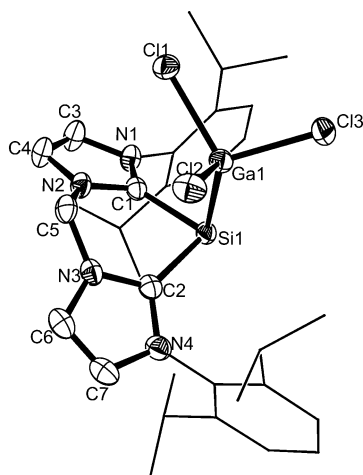


Figure 2. Molecular structures of **4**. Thermal ellipsoids are set at 50% probability. H atoms are omitted for clarity. Selected interatomic distances [Å] and angles [°]: Ga1-Cl3 2.2319(6), Ga1-Cl2 2.2516(6), Ga1-Cl1 2.2884(5), Ga1-Si1 2.4588(6), Si1-C2 1.942(2), Si1-C1 1.942(2), N1-C1 1.351(3), N1-C3 1.390(3), C1-N2 1.351(3), N2-C4 1.377(3), N2-C5 1.445(3), C2-N4 1.346(3), C2-N3 1.351(3), N3-C5 1.450(3), C3-C4 1.340(3), Cl3-Ga1-Cl2 102.42(2), Cl3-Ga1-Cl1 104.55(2), Cl2-Ga1-Cl1 97.45(2), Cl3-Ga1-Si1 113.32(2), Cl2-Ga1-Si1 120.82(2), Cl1-Ga1-Si1 115.83(2), C2-Si1-C1 88.59(9), C1-Si1-Ga1 92.16(9), C2-Si1-Ga1 92.42(9).

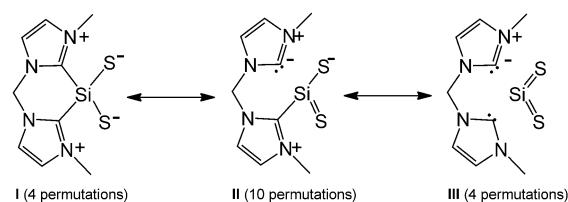
a monomeric structure with the GaCl_3 moiety directly coordinated to the zero-valent Si atom. Thus the three-coordinate silicon atom features a pseudotetrahedral coordination geometry with a lone pair of electrons occupying the vertex. This implies that one of the lone pairs of the silicon center donates charge to the GaCl_3 molecule. The C-Si-C angle of $88.59(9)^\circ$ is quite close to the corresponding angle found in silylone **1** ($89.1(2)^\circ$). The sum of the bond angles around the silicon atom is 273.17° . The Si-C distance of $1.942(2) \text{ Å}$ is slightly longer than the value in **1** (1.869 Å),^[10b]

but similar to the distance observed in **3**. The Si1-Ga1 distance of $2.4588(6) \text{ Å}$ is similar to the Si-Ga values observed in a hypersilylated polygallium cluster $[\text{Ga}_{10}\{\text{Si}(\text{SiMe}_3)_3\}_9]$ ($2.443(1)$ to $2.468(1) \text{ Å}$).^[13]

The ^1H NMR spectrum of **4** shows one doublet and one broad signal for the methyl groups of the isopropyl substituents, presumably, caused by the coordination of GaCl_3 and the associated steric congestion which hinders the free rotation of the isopropyl groups. In line with this, the $^{29}\text{Si}\{^1\text{H}\}$ NMR spectrum exhibits a broad signal at $\delta = -119.0 \text{ ppm}$, strongly upfield from that of **1** ($\delta = -83.8 \text{ ppm}$). Interestingly, the reaction of **4** with elemental sulfur enables an alternative approach to **3** (Scheme 2). In fact, mixing **4** with 0.25 molar equivalent of S_8 in THF at room temperature affords **3** in 50 % yield after work-up.

To elucidate the electronic structure of **2** and **3**, DFT calculations at the B3LYP level of theory have been performed for truncated model systems (**2'**, **3'**, cf. Supporting Information for details). Table 1 lists the calculated ^{29}Si chemical shifts (which are close to the computed values for the full, untruncated systems), as well as NPA charges and Wiberg/Mayer bond orders.

To achieve closer insight into bonding in **2'** and **3'** in terms of resonance, analyses based on natural resonance theory (NRT)^[14] have been performed. Detailed NRT analyses are provided in Schemes S1 and S2 in the Supporting Information. Concisely summarizing the main results, Scheme 3 shows



Scheme 3. Predominant electronic-structure situation for compound **2'** (I) and no-bond/double-bond resonance structures (II, III). Permutation numbers include only resonance structures above 0.5 % weight.

that the electronic structure is best described by semi-polar Si-S bonds and a delocalization of positive charge into the bis-carbene ligand framework. The underlying NRT structures account for about 80–85 % of the one-particle density matrix. Si-S multiple-bonding character is built in through no-bond/double-bond resonance structures (Scheme S1, S2; cf. Ref. [15] for related analyses in main-group chemistry). These contributions correspond to the notion of negative hyperconjugation in MO language. The overall weight of such resonance structures is about 11–13 % in **2'** and **3'**, consistent with relatively polar Si-S π -bonding (S_1 = the axial S atom, S_2 = the equatorial S atom). In **2'**, such resonance structures pertaining to the equatorial Si- S_2 bond contribute 9 %, those to the axial Si- S_1 bond approximately 4 %. GaCl_3 coordination to the axial S_1 (made favorable by its more negative sulfur charge, Table 1) eliminates the Si- S_1 contributions but enhances the weight of equatorial Si- S_2 multiple-bonding NRT structures to 11 %. These observations are fully consistent with Wiberg bond indices (WBI, Table 1), which

exhibit moderate π bonding for both Si–S bonds in **2'** (more for Si–S₂) but a clear reduction of bond order for Si–S₁ and a slight increase for Si–S₂ in **3'** (note also the large negative charges on sulfur). NPA charges clearly confirm the expected charge donation to the GaCl₃ Lewis acid (Table 1).

In summary, the first isolable monomeric silicon disulfide complex **2** could be synthesized by taking advantage of the donor–acceptor concept, starting from the electron-rich silylone **1** and elemental sulfur. One of the highly polar Si–S bonds in **2** is able to form a Lewis acid–base adduct, as shown by the reaction with GaCl₃ to give the (bis-NHC)Si(S)S–Ga complex **3**. Compound **3** is also accessible from direct oxidation of the unprecedented silylone–GaCl₃ adduct **4** with elemental sulfur. Bonding analyses indicated highly polar Si–S bonds with moderate multiple-bonding character made possible by negative hyperconjugation (no-bond/double-bond resonance). Notably, the multiple bonding character of the axial Si–S₁ bond in **2'** is lower than that of the equatorial Si–S₂ bond. In **3'**, GaCl₃ coordinates to the more negatively charged axial S₁. This quenches much of the π -bonding character in the Si–S₁ bond but enhances that in the Si–S₂ bond. Featuring a zero-valent silicon atom, the silylone **1** and its adduct **4** are also very sensitive towards other oxidation reagents such as dioxygen and N₂O which may enable the synthesis of isolable complexes of monomeric silicon oxides.^[16] Respective investigations are currently in progress.

Keywords: hyperconjugation · multiple bonds · N-heterocyclic carbenes · silicon · sulfur

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