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Tetracyclic silaheterocycle formed through a pericyclic reaction cascade including a two-fold intramolecular C–C bond activation[†]

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with 1.5 equiv. of Mg*.

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Reductive debromination of the tribromoamidosilane 2 gave the tetracyclic silaheterocycle 3 through a unique reaction cascade involving unprecedented two-fold intramolecular cycloaddition by transient silylenes. Experimental and computational analyses of the reaction mechanism allowed the identification of the key intermediates that lead to the silaheterocycle 3 or, alternatively, to the cyclotrisilene 19.

During the past decade, the synthesis and subsequent use of very bulky amido ligands, such as {N(SiR₃)Ar} (R = Me, ⁱPr; Ar = aryl group), have paved the way for kinetic stabilisation of many low-valent complexes of heavier main group elements and those in the group 14 particularly.¹ This has led to the characterisation of, *inter alia*, the first examples of amido-substituted digermynes and distannynes with long Ge–Ge or Sn–Sn single bonds, formally bis(tetrylenes),² stable two-coordinate acyclic silylenes and silylsilylenes,³ and multiply bonded amidodigermynes.⁴ While these species are of fundamental interest, they also show fascinating reactivity with small molecules. For example, singly bonded amido-substituted digermynes and distannynes have been reported to reduce both CO and CO₂.⁵

Recently, we examined the reduction of {N(SiMe₃)Dipp}SiBr₃ (Dipp = 2,6-ⁱPr₂-C₆H₃) with 1.5 equiv. of activated magnesium (Mg*).⁶ This was found to give the cluster Si₄{N(SiMe₃)Dipp}₄ I, formally a dimer of two amidodisilynes, with a butterfly-type structure.⁷ Further studies showed that I reacts readily with heavier chalcogens to give amido-substituted cage compounds,

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while its thermolysis led to the six-vertex silicon cluster Si₆{N(SiMe₃)Dipp}₄ II with lone pair character at the ligand-free

vertices.8 These results led us to pose the question of the

influence of the steric bulk on the structure of I. Specifically,

could the butterfly-type Si4 skeleton of I be forced into

planarity,⁹ or would steric strain lead to stabilisation of other amido-substituted silicon rings or cages. As an attempt to

answer the above question, we synthesised the tribromoamido-

silane {N(SiMe₂Ph)Dipp}SiBr₃ 2 and carried out its reduction

Me₃S

Dipp

Reaction of the lithium amide Li{N(SiMe₂Ph)Dipp} 1 with SiBr₄

in Et₂O gave the tribromoamidosilane {N(SiMe₂Ph)Dipp}SiBr₃ 2 in

good yield (ESI⁺), similarly to the corresponding trichlorosilane.¹⁰

The NMR data of 2 are consistent with its formulation and its

(Scheme 1 and ESI⁺). Extraction with *n*-hexane gave a vellow

residue that contained two primary products based on the NMR data (ESI†). The tetracyclic silaheterocycle **3** was subsequently

crystallized as yellow rods from toluene in low but reproducible

Compound 2 was reacted with 1.5 equiv. of Mg* in THF

structure was confirmed by single crystal X-ray diffraction (ESI⁺).

Dipp

 $\label{eq:scheme1} \begin{array}{l} \mbox{Reductive debromination of the tribromoamidosilane 2 to the silaheterocycle 3 (Dipp = 2,6-^i Pr_2C_6H_3). \end{array}$

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Fig. 1 Solid-state structures of 3 (left), 8 (middle), and 19' (right) with thermal ellipsoids drawn at 50% probability level and hydrogen atoms and cocrystallized solvent molecules omitted for clarity. Full structural details are provided in the ESI.†

yields (Fig. 1). The combination of six silicon atoms, three amido groups, and one bromine atom in 3 suggests that the reductive debromination of 2 gives, in sequence, the tetrabromodisilane 4, the dibromodisilene 5, and the disilyne 6 (Scheme 2). This is supported by literature syntheses of [{N(SiMe₃)Dipp}SiBr₂]₂,^{3a} analogous to 4, and [{N(SiMe₃)Ar*}Ge]₂, similar to 6 but with bulkier substituents (Ar* = C₆H₂ⁱPr{C(H)Ph₂}₂-4,2,6, C₆H₂Me{C(H)Ph₂}₂-4,2,6).^{2a,4} Presumably, the conversion of 6 to 3 takes place by a reaction of the former with the bromosilylene 7, generated by an equilibrium with 5, followed by an unprecedented intramolecular two-fold cycloaddition to a single phenyl substituent on one of the SiMe₂Ph groups. Dynamic disilene \leftrightarrow silylene equilibria are well-known in the literature.¹¹

Direct experimental support for the steps outlined above was sought by trapping the proposed intermediates. The dibromodisilene intermediate 5 could be trapped through [2+2] cycloaddition with ethylene (Scheme 2 and ESI†), yielding the disilacyclobutane 8 (Fig. 1). A related cycloaddition product has been obtained from reductive dehalogenation of $\{N(SiMe_3)Dipp\}SiBr_3$ with lithium naphthalenide.^{3a} In the presence of 2,3-dimethyl-1,3-butadiene (dmbd), the reductive debromination of 2 gave the [4+2] cycloaddition product $\{N(SiMe_2Ph)Dipp\}SiBr(dmbd)$ 9 as evidence for the equilibrium between 5 and 7 (Scheme 2 and ESI†). Conceivably, 9 could also arise from successive reduction of 2 in the presence of dmbd. Evaluation of the equilibrium between 5 and 7 with computational methods showed that the dibromodisilene is favoured only by 19 kJ mol⁻¹ in the gas phase, supporting the coexistence of 5 and 7 in solution.

Attempts to trap other intermediates, such as **6**, were met with failure. For this reason, the reaction mechanism connecting **6** to **3** was investigated computationally. Two related pathways were identified (Scheme 3). In pathway 1, the combination of **6** and 7 gives the acyclic intermediate **10** that then undergoes two consecutive intramolecular cycloadditions coupled by a 1,2-bromine shift to give **3**. In pathway 2, an internal cycloaddition of **6** first gives the silacycloheptatriene (silepin) **18** that then reacts with **7** to give **12**, an intermediate common to both pathways, that ultimately transforms to **3** *via* 1,2-bromine shift and second cycloaddition.

The results of DFT calculations (Scheme 3 and ESI[†]) show that pathways 1 and 2 have low and equal activation barriers (highest $\Delta G^{\ddagger} = 59$ and 54 kJ mol⁻¹, respectively). However, considering that the initial reduction of **2** is carried out at low temperature, the barrierless and exergonic formation of **10** can give an advantage to pathway **1**. The facile nature of this



Scheme 2 Trapping of intermediates 5 and 7 en route from 2 to 3 (Dipp = 2,6- $^{i}Pr_{2}C_{6}H_{3}$, dmbd = 2,3-dimethyl-1,3-butadiene).





reaction step agrees with the electronic structure of **6** that, like its heavier germanium analogues,^{2*a*,4} was found to have two minima in agreement with disilyne and bis(silylene) formulations, that is, with short and long Si–Si bonds, respectively (**6**_{short} and **6**_{long}, ESI†). As the two isomers are separated only by 27 kJ mol⁻¹, with the bis(silylene) structure lower in energy, the reaction between **6** and **7** can be viewed as a combination of two silylenes to generate a disilene functionality. Overall, the computational work indicates that the formation of **3** from **6** should be rapid and exergonic ($\Delta G = -219$ kJ mol⁻¹). The first steps from **6** to **14** are governed almost exclusively by silylene reactivity of the intermediates, whereas the last steps connecting **14** to **3** involve rearrangement of the sila(cyclopropyl) product from the second cycloaddition.

Further support to the pathways outlined in Scheme 3 can be found from the literature. Silylenes are well-known to undergo thermally or photochemically activated intermolecular cycloadditions with several aromatic substrates. These typically give either sila(cyclopropyls) or silepins, such as **III** and **IV**, analogous to **11/17** and **12/18**.¹² Similar reactivity has recently been described for an aluminyl anion stabilised by a xanthenebased diamido ligand that reversibly activates benzene even at room temperature.¹³ Intermolecular two-fold cycloadditions of silylenes to aromatic frameworks also generate sila(cyclopropyls) and silepins (*e.g.* **III**),^{12a} a notable exception being the reaction between a silylene and pyrazine that leads to ring expansion to a cyclooctatriene analogue.¹⁴



The fact that the conversion of 6 to 3 takes place so readily can be explained by the intramolecular nature of the two-fold cycloaddition and the associated negligible entropic penalty. To our knowledge, there are only two prior examples of intramolecular cycloadditions between silylenes and aromatic Dipp substituents and one example of reversible cycloaddition between a disilene and Cp* (Cp* = C_5Me_5),¹⁵ but 3 is the only example in which the ligand framework is attacked twice. In this respect, the unprecedented structure of 3 is reminiscent of two-fold borylated products, such as **V**, that have been obtained from defluorination of difluoro(diorganylamino)boranes with Na/K alloy in the presence of benzene derivatives.¹⁶ While there are no detailed mechanistic data available, the formation of **V** is thought to proceed *via* double attack of *in situ* generated borylenes on the aromatic substrates, in similar fashion as outlined for the formation of 3 in Scheme 3.

The reaction cascade in Scheme 3 is consistent with our inability to trap 6 or other intermediates en route to 3 using bases, hydrocarbons, or transition metal complexes. In this respect, we also considered the possibility that 7 would undergo an internal cycloaddition analogous to that connecting 6 and 18. Calculations showed, however, that the product is not a stable species on the potential energy surface due to its strained geometry. The same is also true if the cycloaddition would involve a Dipp substituent in place of SiMe₂Ph. In similar fashion, detailed potential energy surface scans indicated that the intermediates 17 and 18 will not undergo cycloadditions involving the free silylene moiety and the dangling aromatic substituents adjacent to it. Such reactivity is, however, possible for 13, but in this case, selective attack of the silylene to the silepin is ensured by the negligible activation barrier between **13** and **14** (ΔG^{\ddagger} = 12 kJ mol⁻¹) and the stability of the product $(\Delta G = -165 \text{ kJ mol}^{-1}).$

Having examined the most likely pathways available for the intermediates in Scheme 3, we considered the possibility that **10** cyclises to the cyclotrisilene **19**. This parallels the reactivity reported for a related base-stabilized disilenyl silylene **VI** that is in an equilibrium with the corresponding cyclotrisilene **VII** (and the free base) in solution.¹⁷ The results of computational work showed that the ease of cyclisation of **10** depends on the relative orientation of the amido substituents. When the SiMe₂Ph and Dipp groups are perfectly poised to allow Si–Si bond formation, the activation barrier for ring closure is as low as 11 kJ mol⁻¹ (ESI⁺). Furthermore, the formation of **19** was

found to be highly exergonic ($\Delta G = -181 \text{ kJ mol}^{-1}$) but less than that calculated for 3. Attempts to find reaction pathways connecting **19** to 3 were met with failure due to the energy demand associated with opening the Si₃ ring without an external base (*cf.* equilibrium **VI** \leftrightarrow **VII**). This suggests that **19** is a potential thermodynamic sink, and, therefore, a likely candidate for the unidentified reaction product.



Despite several attempts, compound 19 could not be crystallized from the product mixture. Consequently, the synthesis of its SiMe3 analogue 19' was attempted by carrying out the reduction of {N(SiMe₃)Dipp}SiBr₃ under highly diluted conditions. The replacement of Ph with Me eliminates the possibility for intramolecular cycloaddition and lowers the kinetic barrier associated with cyclisation, whereas the low concentration increases the likelihood of the formation of 19' over the Si₄ ring I. In good agreement with the above, the cyclotrisilene 19' was isolated, albeit in minute quantities, allowing its spectroscopic and structural characterisation (Fig. 1 and ESI[†]). Halogen-substituted cyclopropene analogues are known for germanium,¹⁸ but **19**' is the first example of a corresponding silicon species. When the reduction of 2 is carried out under similar conditions used in the synthesis of 19', an analysis of the reaction mixture by ²⁹Si{¹H}-IG-NMR spectroscopy showed two signals in 2:1 ratio at $\delta = -5.1$ and -10.9 ppm that are comparable to data for 19', -7.5 and -16.8 ppm, respectively (ESI[†]). This lends strong support to the proposed mechanism and the key role played by intermediate 10 with its ability to either cyclise, giving 19, or undergo intramolecular cycloaddition, leading to 3.

In summary, we describe the synthesis and characterisation of the tetracyclic silaheterocycle 3 that was obtained from the reductive debromination of the tribromoamidosilane 2. Experimental and computational analyses of the reaction mechanism implicate that the disilyne 6, generated by the reduction of 2, reacts with the bromosilylene 7 to give the transient intermediate 10. Successive pericyclic reactions coupled by a 1,2-bromine shift result in a two-fold intramolecular C-C bond activation of a single phenyl substituent on one of the SiMe₂Ph moieties. The calculated reaction energies agree with the facile formation of 3, while the reactivity of the silylene functionality in 10 corroborates the formation of a second product, the cyclotrisilene 19. Indirect support for the proposed pathway was obtained via complete characterisation of 19'. Efforts to further fine-tune the steric bulk of the amido ligand to allow a highyielding synthesis of novel halogen-substituted cyclotrisilenes analogous to 19 and 19' are currently underway.

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Conflicts of interest

The authors declare no conflicts of interest.

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