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A Hydride-Substituted Homoleptic Silylborate: How Similar is it to its Diborane(6)-Dianion Isostere?

Jannik Gilmer,^[a] Michael Bolte,^[a] Alexander Virovets,^[a] Hans-Wolfram Lerner,^[a] Felipe Fantuzzi,^{*[b]} and Matthias Wagner^{*[a]}

Abstract: The B-nucleophilic 9*H*-9-borafluorene dianion reacts with 9-chloro-9-silafluorene to afford air- and moisture-stable silylborate salts $M[Ar_2(H)B-Si(H)Ar_2]$ (M[HBSiH], M=Li, Na). Li[HBSiH] and Me₃SiCl give the B-pyridine adduct $Ar_2(py)B-Si-(H)Ar_2$ ((**py)BSiH**) or the chlorosilane Li[Ar_2(H)B-Si(Cl)Ar_2] (Li[HBSiCl]) in C₆H₆-pyridine or THF. In both cases, the first step is H⁻ abstraction at the B center. The resulting free borane subsequently binds a py or thf ligand. While the py adduct is stable at room temperature, the thf adduct undergoes a 1,2-H

shift via the cyclic B(μ -H)Si intermediate **BHSi**, which is afterwards attacked at the Si atom by a Cl⁻ ion to give Li[**HBSiCI**]. DFT calculations were employed to support the proposed reaction mechanism and to characterize the electronic structure of **BHSi**. Treatment of Li[**HBSiCI**] with the N-heterocyclic carbene IMe introduces the neutral donor at the Si atom and leads to Ar₂(H)B–Si(IMe)Ar₂ (**HBSi(IMe)**), a donor-acceptor-stabilized silylene.

Introduction

The use of diboranes, in particular bis(pinacolato)diboron (B₂pin₂), for CH-borylation^[1] and functionalization of alkenes and alkynes^[2] has greatly advanced organic synthesis. In a similar vein, disilanes, especially the highly reactive Si₂Cl₆, are currently establishing themselves as useful reagents for the preparation of organosilanes with a wide range of applications, for example, in silicone chemistry.^[3] The next systematic step in the extension of this chemistry is the progression from homoelement compounds to the corresponding heteroelement compounds, namely silylboranes R₂B–SiR'₃.^[2b,4] Silylboranes are suitable for the introduction of boryl and/or silyl groups into a given substrate molecule (Scheme 1). As an example, Ito and coworkers used pinB–SiMe₂Ph (A) as storage form of the [Bpin]⁻ anion, which they released in situ by heterolysis of the B–Si bond with KOMe to prepare **B**.^[5] Conversely, Wang, Uchiyama

[a] J. Gilmer, Dr. M. Bolte, Dr. A. Virovets, Dr. H.-W. Lerner, Prof. Dr. M. Wagner Institut für Anorganische und Analytische Chemie J. W. Goethe-Universität Frankfurt Max-von-Laue-Str. 7, 60438 Frankfurt (Main) (Germany) E-mail: Matthias.Wagner@chemie.uni-frankfurt.de Homepage: https://www.uni-frankfurt.de/58708118/wagner
[b] Dr. F. Fantuzzi School of Chemistry and Forensic Science University of Kent Park Wood Rd, Canterbury CT2 7NH (UK) E-mail: f.fantuzzi@kent.ac.uk

Homepage: https://www.felipefantuzzi.org/

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Chem. Eur. J. 2022, e202203119 (1 of 8)

Scheme 1. The widely employed silylborane pinB–SiMe₂Ph (A) can be used to introduce either Bpin (B), or SiMe₂Ph (C) or both groups (D) into a substrate molecule without the need for a transition metal catalyst. (i) 1.5 equiv. Ar–Br, 1.2 equiv. KOMe, DME, 30 °C, 1 h, 50–85 % yield; (ii) 0.5 equiv. Ar–F, 1.5 equiv. NaOtBu, THF, room temperature, 12 h, 34–89% yield; (iii) 1.0 equiv. styrene, 10 mol% KOtBu, THF, room temperature, 1 h, 91% yield.

and coworkers introduced the SiMe₂Ph group into benzene derivatives through the activation of the B–Si bond with NaOtBu (cf. C).^[6] Again, Ito et al. performed the silaboration of olefins by means of the pinB–SiMe₂Ph/KOtBu system (cf. D).^[7]

Although more and more evidence for the general utility of silylboranes is emerging, fundamental properties of pristine B–Si systems, largely unperturbed by electronic substituent



effects (±I, ±M), still remain underexplored.^[8] Due to a lack of availability, the parent H₂B–SiH₃ has been studied only theoretically:^[9] Silylene (SiH₂) forms an adduct with monoborane (BH₃), which contains a B(μ -H)Si three-membered ring (2-electron-3-center, 2e3c, bond);^[10] the associated stabilization energy was computed to fall in the range 37–50 kcal mol⁻¹, depending on the level of theory applied. The primary product represents only a local minimum and tends to undergo a 1,2-H shift with an activation barrier of only 1–4 kcal mol⁻¹ to furnish silylborane H₂B–SiH₃ with an exothermicity of about 3–7 kcal mol⁻¹.^[9c] Remarkably, the free borane-silylene complex H₃B-SiH₂ without bridging H atom does not exist, but its formal adduct H₃B-SiH₂·NHC was synthesized by Rivard and coworkers (NHC = 1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene).^[11]



Scheme 2. Analogy between the B–B and B–Si species M[BHB] and BHSi, formally obtained by MH elimination from the isolable isosteres M₂[HBBH] and M[HBSiH]. Carbon atoms marked with asterisks bear *t*Bu substituents.

With the aim of validating the theoretical predictions by experiments, we searched for a synthetically accessible derivative of H₂B-SiH₃ and decided to rely on the same design principles as previously employed for the synthesis of isolable B-B bonded analogues.^[12] We finally settled on the silylborate $[Flu(H)B-Si(H)Flu]^{-}$ ($[HBSiH]^{-}$, $HBFlu/H_2SiFlu = bora-/silafluorene;$ Scheme 2),^[13] the formal H^- adduct of FluB–Si(H)Flu, as the entry point of our studies. Its diborane(6) analogue [HBBH]²⁻ (Scheme 2) exists and has been proven useful for studying the diborane(6) B–B-bond reactivity in dianions [Flu(H)B–B(H)Flu]^{2–.[14,15]} Specifically, H⁻ abstraction from $[HBBH]^{2-}$ leads to the isolable $B(\mu-H)B$ diborane(5) anion $[BHB]^{-[15]}$, which is valuable for modeling the parent $[B_2H_5]^{-}$ anion,^[16] a valence-isoelectronic congener of H₂B–SiH₃. Herein, we disclose the synthesis protocol of [HBSiH]-, describe how the B-bonded, the Si-bonded, or both H⁻ ions can be replaced by other ligands, and show that BHSi is the key intermediate in observed SiH/ligand exchange reactions.

Results and Discussion

Synthesis and characterization of M[HBSiH]

The arguably best established synthetic route to B-Si bonds uses electrophilic boron species and nucleophilic silanide reagents.^[13d,17] We also initially relied on this approach and selected HBFlu·thf^[14] and Cl(H)SiFlu^[18] as starting materials.^[19] Since all our attempts to convert Cl(H)SiFlu to its Li⁺ salt Li[HSiFlu] resulted in overreduction with formation of the 9silafluorenyl-dianion salt Li₂[SiFlu],^[20] we tested the suitability of Li₂[SiFlu] for B-Si-bond formation, but to no avail: the reaction scenario was dominated by redox processes and led to complex product mixtures. It was therefore necessary to resort to an "Umpolung" approach, for which we could use the exact same starting materials: Two-electron reduction of HBFlu-thf with alkali metal in THF gives our recently reported boron-centered nucleophile M_2 [HBFlu] (M = Li, Na).^[15,21,22] Freshly prepared M₂[HBFlu] reacted with Cl(H)SiFlu at room temperature in THF to afford the silylborate salts M[HBSiH] in yields of 83% (M = Li) and 56% (M = Na; Scheme 3). We wanted to have both salts



Scheme 3. Left: Synthesis of M[HBSiH] (M = Li, Na) through reaction of M_2 [HBFlu] with Cl(H)SiFlu. Carbon atoms marked with asterisks bear tBu substituents. Right: Solid-state structure of [Li(thf)₂][HBSiH]; C-bonded H atoms, all tBu substituents, and two non-coordinating C₆H₆ molecules are omitted for clarity. H: gray, Li: magenta, B: green, C: black, O: red, Si: blue.

Chem. Eur. J. 2022, e202203119 (2 of 8)

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available, because previous experience indicates a pronounced influence of the counter cation on the reactivity of subvalent boron compounds.^[23,24]

Since the NMR data of M[HBSiH] (M = Li, Na; [D₈]THF) show only the observed negligible usually cation dependencies,^[15,16a,23] it suffices here to discuss the spectra of Li[HBSiH]: In both the ¹H and ¹³C{¹H} NMR spectra, one sees two signal sets of magnetically inequivalent 2,2'-biphenyldiyl fragments (proton-integral ratio = 1:1). The ¹¹B NMR spectrum contains a doublet at -22.7 ppm (¹J(B,H) = 89 Hz), testifying to the presence of tetracoordinated boron nuclei.^[25] Upon proton decoupling, the doublet collapses to a singlet; the corresponding B-bonded proton gives rise to a 1:1:1:1 quartet at $\delta({}^{1}H) =$ 2.72 ppm. A ²⁹Si NMR signal was not directly detectable, likely because it is severely broadened due to ¹J(Si,B) coupling with the quadrupolar ^{10/11}B nuclei. Yet, a ¹H,²⁹Si-HMBC NMR spectrum showed an intense cross-peak between δ ⁽²⁹Si) = -22.7 ppm and the resonances of the ortho-H atoms of one of the 2,2'biphenyldiyl signal sets (which can thereby be assigned to the silafluorene backbone). A ¹H NMR signal at 4.50 ppm with clearly visible ²⁹Si satellites arises from the Si-bonded proton $(^{1}J(H,Si) = 157.0 \text{ Hz})$. The IR spectrum (recorded by ATR) shows characteristic B-H and Si-H stretches at 2272.7 (calcd. 2273.4) and 2032.6 (calcd. 2053.5) cm⁻¹, respectively.^[18]

The molecular structure of [Li(thf)₂][**HBSiH**] was determined by single-crystal X-ray crystallography (SCXD; Scheme 3). The length of the central B–Si bond is 2.028(3) Å (calcd. 2.030 Å).^[18] The two heteroatom-bonded H atoms were located in the difference Fourier map and freely refined. They adopt an s-*trans* conformation with an H–B–Si–H torsion angle of 180(2)°. [Li(thf)₂][**HBSiH**] forms contact-ion pairs in the solid state, in which the [Li(thf)₂]⁺ ca tion is chelated by the BH-, one *ipso*-C-, and one *ortho*-C atom of the HBFlu unit.

Both M[**HBSiH**] salts are stable toward air and moisture in [D₈]THF. More detailed investigations were conducted using Na[**HBSiH**], which turned out to tolerate even added H₂O (NMR-spectroscopic control).^[18] Until to-date, only very few comparably inert B–Si compounds have been reported,^[26] and also the [**HBSiH**]⁻ isostere [**HBBH**]²⁻ is exceedingly sensitive toward hydrolysis.^[18] Treatment of Na[**HBSiH**] with ethereal HCI as the stronger acid led to protonolysis of the B–Si bond with formation of HBFlu-thf and H₂SiFlu (approx. 90% selectivity by NMR). Both the B–H and the Si–H bond remained untouched. Thus, since we wanted to address primarily these potentially reactive sites, it seemed advisable to replace the Brønsted acid with a simple Lewis acid, such as Me₃SiCI.

By switching from the diborane(6) dianion $[HBBH]^{2-}$ to the silylborate monoanion $[HBSiH]^-$, a break in symmetry is induced. On the one hand, the resulting perturbation of the system should be small (cf. the diagonal relationship of the periodic table). On the other hand, H⁻ abstraction from $[HBSiH]^-$ results either in a neutral borane or in a formal silylium-cation fragment, which should result in significant reactivity differences between the B–H and Si–H bonds. A particularly important question in this context would be to what extent the H atom that remains after H⁻ abstraction can adopt a bridging position between B and Si (BHSi in Scheme 2;

compare the B(μ -H)B 2e3c bond in the crystallographically characterized isostere [**BHB**]⁻).

Outcomes of H⁻ abstractions from M[HBSiH] with Me₃SiX (X = Cl, Br) in the presence of no ligand, the strong ligand py, or the weak ligand thf

We first treated Li[HBSiH] with 1 equiv. of Me₃SiCl in the noncoordinating solvent $\mathsf{C}_6\mathsf{D}_{6^r}$ but obtained no indication of BHSiand only observed the unselective decomposition of the starting material (along with Me₃SiH as the only Me₃Sicontaining species detectable by NMR). To test whether at least the initial step of the reaction cascade(s) can be conducted selectively, we repeated the experiment in the presence of 3 equiv. of pyridine (py) and quantitatively obtained the B-py adduct (py)BSiH (Scheme 4). Next, we confirmed that (py)BSiH generation indeed begins with B-H-bond activation and not with Si-H-bond activation accompanied by a subsequent 1,2-H shift: An isotopic-labeling experiment using Li[DBSiH] and Me_3SiCl fittingly furnished (py)BSiH and Me_3SiD as sole products. Can the strong donor py be replaced with the weaker ligand thf to approach the putative free borane BHSi more closely? When H⁻ abstraction from Li[HBSiH] was carried out in [D₈]THF, Li[HBSiCI] was generated, while a significant amount of



Scheme 4. Divergent reaction behavior after H⁻ abstraction from Li[HBSiH] with Me₃SiX (X = Cl, Br). Carbon atoms marked with asterisks bear tBu substituents. (i) 1 equiv. Me₃SiCl, 3 equiv. py, C₆D₆, room temperature, quantitative conversion to (**py**)BSiH; (ii) 5 equiv. Me₃SiBr, [D₈]THF, -30° C, characterized by NMR; (iii) *without 12-c-4*: 10 equiv. Me₃SiCl, [D₉]THF, room temperature, 90% conversion to Li[HBSiCl]; *with 12-c-4*: 2 equiv. 12-c-4, 30 equiv. Me₃SiCl, THF, room temperature, quantitative conversion to [Li(12-c-4)₂][HBSiCl], 85% yield.

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residual starting material Li[HBSiH] remained (approx. ratio = 2:1 after 2 d). Full conversion of Li[HBSiH] was reached by using 10 equiv. rather than 1 equiv. of Me₃SiCl. By this protocol, Li[HBSiCI] formed with about 90% selectivity (Scheme 4);^[27] among the two side products, we identified HBFlu-thf. In the presence of 12-crown-4, these side products do no longer occur ([Li(12-c-4)₂][HBSiCI] was isolated in 85% yield; the amount of Me₃SiCl required to drive the reaction to completion increased from 10 to 30 equiv.). All in all, we conclude that the two pairs Li[HBSiH]/Me₃SiCl and Li[HBSiCI]/Me₃SiH are engaged in a dynamic equilibrium, as can be rationalized by the fact that the underlying transformation comes close to an identity reaction (see also the corresponding DFT calculations below).

The most obvious route to Li[HBSiCI] would proceed via mere H/Cl exchange at the Si center of Li[HBSiH] and thus, in stark contrast to the first case, require an initial Si-H-bond activation. As an alternative, the reaction could again be initiated by H⁻ abstraction from the B atom, followed by a rearrangement sequence eventually furnishing Li[HBSiCI]. To distinguish between both mechanistic scenarios, we monitored the conversion by low-temperature NMR spectroscopy.^[28] Given that even at the moderately low temperature of -30 °C, Li[HBSiH] and Me₃SiCl (5 equiv.) behaved inert toward each other, we replaced Me₃SiCl by the more reactive Me₃SiBr. Now, the reaction mixture gave rise to prominent signals assignable to the (py)BSiH-analogue (thf)BSiH already at -30°C (Scheme 4). Upon warming to room temperature, the primary intermediate (thf)BSiH was converted to Li[HBSiBr] (NMRspectroscopic control). Since we have no reasons to assume that the reactions with Me₃SiCl and Me₃SiBr follow qualitatively different pathways, we postulate (thf)BSiH as the common intermediate of Li[HBSiX] formation (X=Cl, Br). From this it follows that the reaction between Li[HBSiH] and Me₃SiX in C₆D₆/py and [D₈]THF always yields an initial donor adduct (do)BSiH (do = py, thf). In the case of the stronger ligand py, (py)BSiH can be isolated; in the case of the weaker donor thf, (thf)BSiH reacts with the anionic donor X⁻ present in the solution to give Li[HBSiX] (presumably via the H-bridged intermediate BHSi).

So far, we have developed ways to remove one H⁻ substituent from the [HBSiH]⁻ anion. Is it possible to exchange both H⁻ ligands as well? Since the reaction of Li[HBSiH] in [D₈]THF stops at the Li[HBSiCl] monohydride stage even in the presence of a vast excess of Me₃SiCl, the amount of added H⁻ scavenger alone does not appear to be a sufficient handle. The key to success ultimately lies in the counter-cation effects mentioned above, as Na[HBSiH] could indeed be converted almost quantitatively to the isolable H⁻-free neutral product (thf)BSiCI (10 equiv. Me₃SiCl, THF; yield losses occurred during workup due to the high solubility of (thf)BSiCl in n-hexane; Scheme 5). Na[HBSiCI] was observed as an intermediate by in situ NMR monitoring at room temperature. These results indicate that the first steps of the reaction cascade are the same as for the Li⁺ salt, while the necessary driving force for the follow-up reaction Na[HBSiCI]→(thf)BSiCI must come from the precipitation of NaCl, which is essentially insoluble in THF.^[29] As



Scheme 5. Synthesis of (thf)BSiX by twofold H⁻ abstraction from Na[HBSiH] using Me₃SiX (X=CI, Br). Carbon atoms marked with asterisks bear tBu substituents. (i) 10 equiv. Me₃SiX (X=CI, Br), THF, room temperature, >95% conversion to (thf)BSiX, X=CI: 55% yield, X=Br: 42% yield.

an aside, the use of Me_3SiBr instead of Me_3SiCl furnishes the bromosilane (thf)BSiBr (Scheme 5).^[18]

Characterization of the H⁻-abstraction products (py)BSiH, $[Li(12-c-4)_2][HBSiCI]$, and (thf)BSiCI

NMR spectra of **(py)BSiH** and $[Li(12-c-4)_2]$ [**HBSiCI**]/(**thf)BSiCI** were recorded in C₆D₆ and [D₈]THF, respectively.

The ¹¹B NMR resonance of (**py)BSiH** appears at -2.1 ppm, downfield-shifted by about 20 ppm compared to the starting material Li[**HBSiH**], but still in a range characteristic of tetracoordinated boron nuclei.^[25] The ²⁹Si chemical shift value of -26.6 ppm remains largely the same as in Li[**HBSiH**]. A signal corresponding to a B-bonded H⁻ substituent is no longer detectable in the ¹H NMR spectrum of (**py)BSiH**, whereas a singlet at 5.33 ppm, possessing ²⁹Si satellites and integrating 1 H, can be assigned to the remaining Si*H* proton. The ¹*J*(H,Si) coupling constant of 174 Hz is larger by 17 Hz than ¹*J*(H,Si) in Li[**HBSiH**], which, according to Bent's rule,^[30] is consistent with a higher group electronegativity of the boryl moiety in (**py)BSiH**. The relative integral values of the three py-proton resonances are fitting to the introduction of one py ligand.^[31]

As expected, the changes in the NMR parameters of the HBSiCl core in [**HBSiCl**]⁻ are complementary to those of the (py)BSiH core in (**py**)**BSiH**: While the signals of the HB fragment at $\delta({}^{1}\text{H}) = 2.97$ ppm and $\delta({}^{11}\text{B}) = -21.1$ ppm (d, ${}^{1}J(\text{B},\text{H}) = 94$ Hz) remain close to those of [**HBSiH**]⁻, the signal of the silyl fragment at $\delta({}^{29}\text{Si}) = 23.7$ ppm is shifted to lower field strengths by about 50 ppm and an SiH resonance is no longer visible. An intense singlet at $\delta({}^{1}\text{H}) = 3.41$ ppm (32 H) corresponds to 2 equiv. of crown-ether ligands and thus supports the proposed ion-pair structure [Li(12-c-4)₂][**HBSiCl**]. The presence of one Cl substituent was confirmed by the characteristic isotope pattern of the [**HBSiCl**]⁻-anion peak at m/z = 603.36 in the LDI mass spectrum.^[18]

Upon going from [HBSiCI]⁻ to (thf)BSiCI, the resonance of the BH substituent vanishes; the ¹¹B NMR signal loses its doublet multiplicity and shifts from -21.1 to 7.1 ppm. All these observations point toward successful abstraction also of the second H⁻ substituent, accompanied by the formation of a moderately strong thf adduct. ¹H NMR signals of the original protio-thf ligand, which stems from the synthesis of (thf)BSiCI in [H₈]THF, were found and integration was proportional to

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1 equiv. of protio-thf.^[32] The ²⁹Si nuclei (δ = 8.6 ppm) are shielded by 15.1 ppm relative to those of [**HBSICI**]⁻.

Compounds (py)BSiH, (thf)BSiCl, and (thf)BSiBr were structurally authenticated by SCXD (Figure 1). (py)BSiH crystallizes with two crystallographically independent molecules in the asymmetric unit ((py)BSiH^A, (py)BSiH^B), which vary slightly due to different environments in the crystal packing. The B-Si bond lengths are 2.039(3) and 2.061(3) Å, respectively. The sums of CBC and CBSi bond angles amount to 316.6(4)° ((py)BSiH^A) and 324.4(4)° ((py)BSiH^B), showing a pronounced pyramidalization of the B atom due to coordination of the py ligand. The B-N distances of 1.608(4) and 1.604(3) Å are similar to the B-N distance in BF₃·py (1.604(5) Å).^[33] The Si-bonded H atom was located in the difference Fourier map and freely refined. In (thf)BSiCl, the key bond lengths are B-Si = 2.026(3) Å, B–O=1.584(4) Å, and Si–Cl=2.1106(12) Å. The pyramidalization of the B atom ($\Sigma_{CBE} = 324.8(4)^{\circ}$; E = C or Si) is comparable to that in (py)BSiH^B. The torsion angles COG^B-B-Si-COG^{Si} in (py)BSiH^{A/B} and (thf)BSiCl fall in the range $35 \pm 4^{\circ}$ (COG^{B/Si}: centroid of the respective five-membered ring). Bromosilane (thf)BSiBr is isostructural with (thf)BSiCl in the crystalline state and therefore does not merit further discussion.

Assessment of the reaction Li[HBSiH]/Me₃SiCl and characterization of the key intermediate BHSi by theory

All energies were calculated at the SMD(THF)/PBE0-D3(BJ)/def2-TZVP level of theory, with optimized structures at SMD(THF)/ PBE0-D3(BJ)/def2-SVP.

From our previous experiments, we concluded that the reaction between Li[**HBSiH**] and Me₃SiCl in THF leads to a dynamic equilibrium involving the intermediates (**thf)BSiH** and **BHSi** as well as an H⁻ shift between B and Si. For the overall reaction [**HBSiH**]⁻ + Me₃SiCl \rightarrow [**HBSiCl**]⁻ + Me₃SiH, we computed a free energy of $\Delta G_{298} = -0.2 \text{ kcal mol}^{-1}$ (equilibrium constant K = 1.4); inclusion of the complex cation [Li(thf)₂]⁺ gave a similar picture with $\Delta G_{298} = -0.6 \text{ kcal mol}^{-1}$ (K = 2.8).^[34] Both K values are close to the experimentally determined $K \approx 2$.

To exclude any necessity of considering counter cations, the energetics of the 1,2-H shift were calculated starting from the NMR-characterized, neutral adduct (thf)BSiH (Scheme 6): Barrierless dissociation of the thf ligand affords BSiH with a tricoordinated B center and a terminal SiH atom. The free



Figure 1. Solid-state structures of (**py)BSiH**^A (left; one of two crystallographically independent molecules), (**thf)BSiCI** (middle), and (**thf)BSiBr** (right). C-bonded H atoms, all tBu substituents, and non-coordinating solvent molecules are omitted for clarity. H: gray, B: green, C: black, N: light blue, O: red, Si: blue, Cl: yellow, Br: brown.

Chem. Eur. J. 2022, e202203119 (5 of 8)



Scheme 6. *Top:* Calculated reaction mechanism for the 1,2-H shift from Si to B via BHSi. The reaction starts from (thf)BSiH and finally yields HBSi(thf). Carbon atoms marked with asterisks bear tBu substituents. Energies were calculated at the SMD(THF)/PBE0-D3(BJ)/def2-TZVP level of theory, with optimized structures at SMD(THF)/PBE0-D3(BJ)/def2-SVP. *Bottom:* Intrinsic bond orbital of the B(*u*-H)Si core in the key intermediate BHSi. Total contributions are related to the fraction of electrons of the doubly occupied IBO assigned to each individual atom.

energy of **BSiH** is +13.3 kcal mol⁻¹ relative to the starting point. The key intermediate **BHSi** lies 25.9 kcal mol⁻¹ above (**thf)BSiH**, in a shallow minimum between **TS1** and **TS2** (28.2 kcal mol⁻¹ and 28.6 kcal mol⁻¹, respectively). The H-atom shift from the bridging position in **BHSi** to the terminal position in **HBSi** is exothermic by 8.4 kcal mol⁻¹. We note that the tricoordinated Si atom in **HBSi** has a close intramolecular contact with one of the *ipso*-C atoms of the borafluorene moiety (Si···C = 2.173 Å), which can be traced back to some stabilizing hyperconjugation between the *ipso*-C–B σ bond and the Si(p_z) acceptor orbital.^[18] Finally, an incoming thf ligand binds to the Si center of **HBSi** without a barrier and generates **HBSi(thf)**, which has essentially the same energy as its isomer (**thf)BSiH**.

The central H atom of **BHSi** interacts with both the B and the Si atom as shown by the intrinsic bond orbital (IBO) of the heteroatom core (total contributions: H=0.91, B=0.64, Si=0.41; Scheme 6).^[35] The Wiberg/Mayer bond indices for the B–H and Si–H bond contributions were computed as 0.604/0.453 and 0.333/0.481, respectively (cf. the Mayer bond orders for B₂H₆: B-(μ -H)=0.47, B–H=0.98).^[36,37] These results consistently indicate the existence of a B(μ -H)Si 2e3c bond supporting the B–Si 2e2c bond in **BHSi**.

Synthesis and characterization of HBSi(IMe)

We have so far observed that, after H^- abstraction, the neutral donor (do⁰) py tends to prefer the neutral B center, while the anionic donor Cl⁻ aims for the formal silvlium cation. Is it

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nevertheless possible to force the system into an isolable HBSi(do⁰) structural motif? To reach this goal, we selected the sterically undemanding 1,3-dimethylimidazol-2-ylidene ligand (do⁰ = IMe), because its carbene-C atom also possesses properties of a carbanionic ligand (cf. the pronouncedly zwitterionic resonance structures of IMe).^[38] In a nucleophilic substitution approach, a mixture of the appropriately pre-organized [Li(12-c-4)₂][HBSiCI] and LiH^[39] in [D₈]THF was treated at room temperature with the imidazolium iodide [HIMe]I for the in situ release of IMe (Scheme 7). After 5 d, 90% conversion to HBSi(IMe) was achieved. In the NMR spectra, Cl/IMe exchange leads to an upfield shift of the ²⁹Si NMR signal from 23.7 to -16.0 ppm,



Scheme 7. Treatment of [HBSiCI]⁻ with exc. LiH/[HIMe]I in [D_a]THF at room temperature gives HBSi(IMe) (>90% conversion after 5 d). HBSi(IMe) can plausibly be described as donor-acceptor-stabilized silylene, which would render it the NHC-coordinated BSi analog of [HBB]⁻ (dashed box), a crucial isomer to explain observed reactivity patterns of [BHB]⁻ (similar arguments hold for HBSi and HBSi(thf)). Carbon atoms marked with asterisks bear *t*Bu substituents.



Figure 2. a) Molecular structure of **HBSi(IMe)** in the solid state. C-bonded H atoms, all *t*Bu substituents, and non-coordinating solvent molecules are omitted for clarity. H: gray, B: green, C: black, N: light blue, Si: blue. b) Plots of the deformation densities $\Delta \rho_1$ (Si—IMe bond) and $\Delta \rho_{1'}$ (B—Si bond) of the main pairwise contributions associated with the orbital interaction term ΔE_{orb} (kcal mol⁻¹) in **HBSi(IMe)**. Interacting fragments: *left*: [Flu(H)B-SiFlu] with [:IMe]; *right*: [HBFlu] with [:Si(IMe)Flu]. Values in parentheses are the percentage of the pairwise orbital interaction with respect to the total ΔE_{orb} contribution. The eigenvalues $\nu_{\pm k}$ (k = 1/1' for the Si—IMe/B—Si case) correspond to a charge transfer from orbital ψ_{-k} to orbital ψ_{+k} in the NOCV representation. Isovalues: 0.003. Charge flows from red to blue. All C-bonded H atoms and *t*Bu substituents are omitted for clarity.

while the ¹H and ¹¹B resonances of the HB moiety shift by less than 0.2 and 1 ppm, respectively. Integration of the ¹H NMR spectrum confirms the presence of one IMe ligand in **HBSi(IMe)**; the ¹³C resonance of the carbene-C atom appears at 158.4 ppm. The molecular structure of **HBSi(IMe)** was characterized by SCXD (Figure 2); since the crystals obtained were only very weakly diffracting, we refrain from a detailed discussion of structural parameters.^[18]

With (py)BSiH and HBSi(IMe), we have the two possible forms of neutral donor adducts at our disposal, but still with different ligands. To evaluate which of the isomers (do⁰)BSiH or HBSi(do⁰) with do⁰ = py or IMe has the lower energy, we also computed these two pairs (cf. Scheme 6 for the (thf)BSiH/ HBSi(thf) pair). The minimum structure (py)BSiH is energetically favored over its isomer HBSi(py) by 4.2 kcalmol⁻¹, consistent with our experimental observations. The isomer pair (IMe)BSiH/ HBSi(IMe) exhibits a similar trend with an even more pronounced energy difference of 8.1 kcalmol⁻¹. Thus, B-coordination is apparently thermodynamically preferred for strong neutral donors. The isolation of HBSi(IMe) is most likely enabled by kinetics: We assume that the Si-bonded Cl⁻ ion is replaced by IMe via an S_N2-type mechanism. Once coordinated to the Si center, the dissociation of IMe required for the rearrangement HBSi(IMe)→(IMe)BSiH would be associated with a prohibitively high energy penalty (compare the energy difference between **HBSi**(IMe) and **HBSi**/IMe of 41.4 kcal mol⁻¹).

The conceptual role model [BHB]⁻ (cf. Scheme 2) of the BHSi chemistry outlined herein shows a reactivity that is best explained by postulating the rearranged isomer [HBB]⁻ as crucial intermediate (Scheme 7) that acts as synthetically useful source of [:BFlu]⁻ anions.^[16,40] This suggests that the sp³-sp² diborane [HBB]⁻ should be described as an acceptor-stabilized boryl anion. By the same token, HBSi(IMe) could be viewed as a donor-acceptor-stabilized silylene (Scheme 7). We have answered the question of how valid this view is with the help of energy decomposition analysis in combination with the natural orbitals for chemical valence method (EDA-NOCV) based on the optimized structure for HBSi(IMe): The interaction between neutral closed-shell fragments, i.e., [Flu(H)B-SiFlu] with [:IMe] and [HBFlu] with [:Si(IMe)Flu], rather than between ionic openshell fragments resulted in the least negative orbital interaction terms $\Delta E_{\rm orb}$.^[18] This reveals that the choice of neutral fragments leads to a better description of the considered bonds in HBSi(IMe). For these, only one NOCV pairwise interaction contributes majorly to the corresponding bonding situations. The resulting deformation densities indicate electron-charge flows from the electron-rich to the electron-poor fragments, which are representative of donor-acceptor coordination (Figure 2). Taken together, these results demonstrate that the bonding situation in HBSi(IMe) can best be delineated as that of a donor-acceptor-stabilized silylene.

Conclusion

Molecules containing B–Si bonds are not only interesting in their own right, but are also becoming increasingly important as

building blocks for organoelement compound synthesis. Compared to silvlboranes, silvlborates are still more scarce; herein, we disclosed an access route to both compound classes. As decisive asset, the parent silvlborate of our studies, M[Flu(H)B-Si(H)Flu] (M[HBSiH], M=Li, Na), bears exclusively H and aryl substituents that allow key properties of the B-Si core to be evaluated in their purest form and largely unbiased by ligand artifacts. Apart from B vs. Si, both halves of the molecule are identical, so that their reactivity is only influenced by the different nature of the heteroatoms. Furthermore, the anion [HBSiH]- is an isostere of the known diborane(6) dianion [Flu(H)B-B(H)Flu]²⁻ ([HBBH]²⁻),^[14,15] so that a direct comparison of both species is possible. Important differences are: (i) [HBSiH]⁻ is stable under an ambient atmosphere, whereas [HBBH]²⁻ is extremely sensitive toward hydrolysis. (ii) H⁻ abstraction from [HBBH]²⁻ yields the isolable product $[BHB]^{-}$, which contains a central three-membered ring with B(μ -H)B 2e3c bond; the analogous BHSi with $B(\mu$ -H)Si 2e3c bond is a transient intermediate, but nevertheless very useful for the preparation of differently substituted silylborates and -boranes. H⁻ abstraction from [HBSiH]⁻ always occurred at the B center, but a subsequent 1,2-H shift of the remaining Si-bonded H atom is facile. Considering its three sterically unencumbered reactive sites, i.e., the B-H, Si-H, and B-Si bonds, we are optimistic that [HBSiH]⁻ will prove to be a similarly fruitful entry into subvalent boron-silicon chemistry as [HBBH]²⁻ has been for diborane chemistry.[14-16]

X-ray Crystallography

Deposition numbers 2203993 (for $[Li(thf)_2][HBSiH]$), 2203991 (for $[Na(thf)_4][HBSiH]$), 2203990 (for (py)BSiH), 2203995 (for (thf)BSiCI), 2203992 (for (thf)BSiBr), 2203994 (for HBSi(IMe)), and 2203989 (for Cl₂SiFlu) contain the supplementary crystallographic data for this paper. These data are provided free of charge by the joint Cambridge Crystallographic Data Centre and Fachinformationszentrum Karlsruhe Access Structures service.

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Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

The data that support the findings of this study are available in the supplementary material of this article.

Keywords: diagonal relationship • hydride abstraction silylborane • subvalent compounds • umpolung

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RESEARCH ARTICLE



A 1,2-dihydride-substituted silylborate was synthesized via an reactivities

"Umpolung" approach using a chlorosilane and a boron-centered nucleophile. Except for the B–Si vs. B–B cores, the silylborate is identical to a prominent diborane(6) dianion,



J. Gilmer, Dr. M. Bolte, Dr. A. Virovets, Dr. H.-W. Lerner, Dr. F. Fantuzzi*, Prof. Dr. M. Wagner*

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A Hydride-Substituted Homoleptic Silylborate: How Similar is it to its Diborane(6)-Dianion Isostere?