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Thermodynamic versus kinetic control in substituent redistribution reactions of silylium ions steered by the counteranion†

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An in-depth experimental and theoretical study of the substituent exchange reaction of silylium ions is presented. Apart from the substitution pattern at the silicon atom, the selectivity of this process is predominantly influenced by the counteranion, which is introduced with the trityl salt in the silylium ion generation. In contrast to Müller's protocol for the synthesis of triarylsilylium ions under kinetic control, the use of Reed's carborane anions leads to contact ion pairs, allowing selective formation of trialkylsilylium ions under thermodynamic control. DFT calculations finally revealed an unexpected mechanism for the rate-determining alkyl exchange step, which is initiated by an unusual 1,2-silyl migration in the intermediate *ipso*-disilylated arenium ion. The resulting *ortho*-disilylated arenium ion can then undergo an alkyl transfer *via* a low-barrier five-centered transition state.

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Introduction

Silvlium ions (R₃Si⁺) have recently emerged as useful and versatile catalysts for synthetically attractive transformations.^{1,2} The most commonly used approach to generate silylium ions is the Bartlett-Condon-Schneider reaction,3 that is the silicon-tocarbon hydride transfer from a hydrosilane to the trityl cation (Ph₃C⁺) paired with a weakly coordinating counteranion.^{4,5} However, substituent redistribution of the hydrosilane starting material can occur under these highly Lewis acidic reaction conditions, leading to undesired mixtures of various silicon compounds.6-8 Hence, hydrosilanes containing three identical substituents, e.g. Et₃SiH or iPr₃SiH, are usually employed in this reaction.9 Conversely, Müller and co-workers have turned this unselective process into a useful synthetic route to triarylsilylium ions (Scheme 1, top).10 When sterically demanding methyl(diaryl)silanes MeAr₂SiH are used in the hydride abstraction with Ph₃C⁺[B(C₆F₅)₄]⁻, the formation of otherwise difficult to prepare triarylsilylium ions $Ar_3Si^{\dagger}[B(C_6F_5)_4]^{-}$ is

Herein, we report that treatment of hydrosilanes of type Me_2RSiH (R=aryl, benzyl) with Reed's carborane-based trityl salt $Ph_3C^{\dagger}[CHB_{11}H_5Br_6]^{-}$ (ref. 13) results in substituent exchange reactions selectively forming the elusive trimethylsilylium ion $Me_3Si^{\dagger}[CHB_{11}H_5Br_6]^{-}$ (Scheme 1, bottom). This method thus complements Müller's approach and offers a practical route to Me_3Si^{\dagger} , avoiding the use of gaseous and highly flammable $Me_3SiH.^{14}$ A systematic experimental and computational investigation was performed to gain a full

Scheme 1 Divergence in the generation of silylium ions by substituent redistribution (x + y + z = 4).

R = aryl, benzyl

observed.¹¹ Notably, the useof less bulky hydrosilanes such as $MePh_2SiH$ or $Me(o\text{-}Tol)_2SiH$ does not give triarylsilylium ions but mixtures of different silicon cations.¹²

Müller (2011): $Ph_3C^{+}[B(C_6F_5)_4]^{-}$ MeAr₂SiH $Ar_3Si^{+}[B(C_6F_5)_4]^{-}$ C_6D_6 rt, 1 h (1.6 equiv) ²⁹Si NMR (C₆D₆) - Ph₃CH δ 216–230 ppm - Me₃SiH Ar = 2,6-disubstituted phenyl this work: Ph₃C⁺[CHB₁₁H₅Br₆]⁻ Me₂RSiH Me₃Si⁺[CHB₁₁H₅Br₆]⁻ toluene ²⁹Si NMR (o-Cl₂C₆D₄) rt. overniaht (2 equiv) - Ph₃CH δ 93 ppm – Me_xR_vSiH_z

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[‡] Responsible for X-ray diffraction analysis.

mechanistic picture of this phenomenon. DFT calculations revealed an unexpected mechanism and suggested an active

role of the carborane counteranion in the outcome of these reactions.

Results and discussion

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Generation of the trimethylsilylium ion by substituent redistribution

When a mixture of Me₂PhSiH and Ph₃C⁺[CHB₁₁H₅Br₆] in toluene was stirred overnight at room temperature, a white suspension was obtained. The solid was collected by filtration, washed with n-pentane, and dissolved in o-Cl2C6D4 for NMR spectroscopic analysis. Unexpectedly, only a singlet at 0.83 ppm was detected in the ¹H NMR spectrum, while no aromatic resonances except for those of the deuterated solvent were observed. The low-field ²⁹Si NMR chemical shift of 93 ppm in the corresponding ¹H/²⁹Si HMQC spectrum, which is characteristic of trialkylsilylium ions, indicated clean formation of Me₃Si⁺[CHB₁₁H₅Br₆] (Fig. 1). The structural integrity of the carborane counteranion was confirmed by 11B NMR spectroscopy.

Unambiguous evidence for the structure of Me₃Si⁺[CHB₁₁-H₅Br₆] was eventually provided by its crystallographic characterization (Fig. 2).15 Single crystals suitable for X-ray diffraction analysis were obtained by vapor diffusion with n-hexane from a solution of the silylium salt in o-F₂C₆H₄ at room temperature. In accordance with reported molecular structures of silylium carboranes,16 one bromine atom at the pentagonal belt of the icosahedral anion is bound to the silicon cation. Both the Si-Br bond distance of 2.435(6) Å and the sum of all C-Si-C bond angles of 346.3(6)° are comparable to the larger $\mathrm{Et_3Si}^+[\mathrm{CHB_{11}H_5Br_6}]^-.$

In contrast to the clean formation of Me₃Si⁺, the non-polar *n*pentane filtrate contained several tri- and tetraorganosilanes, such as Ph₄Si, MePh₃Si, Ph₃SiH, Me₂Ph₂Si, MePh₂SiH, Me₃PhSi, and Me₂PhSiH, as verified by GC-MS analysis. Since silylium

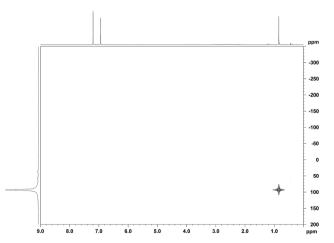


Fig. 1 1 H/ 29 Si HMQC NMR spectrum (500/99 MHz, o-Cl₂C₆D₄, 298 K, optimized for J = 7 Hz) of Me₃Si⁺[CHB₁₁H₅Br₆]⁻ from the reaction of Me₂PhSiH with Ph₃C⁺[CHB₁₁H₅Br₆]⁻.

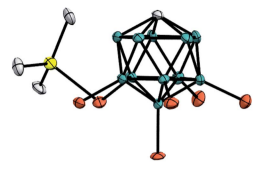


Fig. 2 Molecular structure of Me₃Si⁺[CHB₁₁H₅Br₆]⁻ (thermal ellipsoids at the 50% probability level; H atoms omitted for clarity).

ions are known to promote substituent redistribution,8 this result did not come as a surprise but raised the question why Me₃Si⁺ was selectively formed in this reaction mixture, whereas Müller's conditions cleanly afford sterically congested triarylsilylium ions.10

Influence of the substituent pattern at the silicon atom on the selectivity of the substituent redistribution reaction

To understand the differences between Müller's protocol¹⁰ and our findings, we systematically studied the hydride transfer reaction of various hydrosilanes of type MeAr₂SiH and Me₂ArSiH using trityl salts Ph₃C⁺[B(C₆F₅)₄] and Ph₃- $C^{+}[CHB_{11}H_{5}Br_{6}]^{-}$ (Table 1). Depending on the counteranion, slightly modified procedures were applied for the generation of the silicon cations (see the ESI† for details). For all reactions, an excess of hydrosilane (4 equiv.) was used, thereby

Table 1 Silylium ion generation by substituent redistribution: effect of the hydrosilane and counteranion (Si = triorganosilyl)

$$R_3Si-H \xrightarrow{Ph_3C^+[X]^-} R_3Si^+[X]^-$$

$$-Ph_3CH - Me_xAr_ySiH_z$$

$$R_3Si^+[X]^-$$

Entry ^a	Si-H (4 equiv.)	[X] ⁻	Si ⁺	δ(²⁹ Si) ^b [ppm]
1	Me(C ₆ Me ₅) ₂ SiH	$[B(C_6F_5)_4]^-$	$(C_6Me_5)_3Si^+$	217
2	$Me(C_6Me_5)_2SiH$	$[CHB_{11}H_5Br_6]^-$	$(C_6Me_5)_3Si^+$	217
3	MePh ₂ SiH	$[B(C_6F_5)_4]^-$	c	_
4	MePh ₂ SiH	$[CHB_{11}H_5Br_6]^-$	MePh ₂ Si ⁺ / Me ₂ PhSi ^{+d}	57/76
5	Me ₂ PhSiH	$[B(C_6F_5)_4]^-$	c	_
6	Me ₂ PhSiH	[CHB ₁₁ H ₅ Br ₆]	Me ₃ Si ⁺	93
7	Me ₂ (C ₆ Me ₅)SiH	$[B(C_6F_5)_4]^-$	$(C_6Me_5)_3Si^+$	217
8^e	$Me_2(C_6Me_5)SiH$	$[CHB_{11}H_5Br_6]^-$	Me ₃ Si ⁺	93

^a All reactions were performed according to General Procedure (GP) 1 for $X^- = [B(C_6F_5)_4]^- (C_6D_6$, room temperature, 60 min) or GP 2 for X^- = $[CHB_{11}H_5Br_6]^{-1}$ (toluene, room temperature, 18–24 h). See the ESI for details. b Measured in o-Cl₂C₆D₄. c A complex mixture was obtained as a result of counteranion decomposition.17 d Ratio of 79:21 determined by ¹H NMR spectroscopy. ^e Reaction performed at Chemical Science Edge Article

excluding any influence of stoichiometry on the product formation. In accordance with Müller's report, bulky methyl(diaryl)silane Me(C₆Me₅)₂SiH was converted to the corresponding triarylsilylium ion, regardless of which counteranion was used (entries 1 and 2). In contrast, hydride abstraction from sterically less hindered MePh2SiH with Ph₃C⁺[B(C₆F₅)₄]⁻ led to a complex reaction mixture as a result of anion decomposition (entry 3). 12,17 The use of the carborane counteranion [CHB₁₁H₅Br₆] furnished the unscrambled silylium ion MePh₂Si⁺[CHB₁₁H₅Br₆]⁻, as confirmed by X-ray diffraction analysis (entry 4; see the ESI† for the molecular structure of MePh₂Si⁺[CHB₁₁H₅Br₆]⁻).¹⁵ However, the formation of the MePh₂Si⁺ cation was accompanied by a substantial amount of a second silylium ion, which was found to be the Me₂PhSi⁺ cation. Notably, longer reaction times (7 days) or elevated temperatures (50 °C for 72 h) did not significantly change the product ratio of \sim 79 : 21 (not shown). In all cases, the generation of Me₃Si⁺ was not observed. We then turned our attention to dimethyl(aryl)silanes (entries 5-8). The reaction of Me₂PhSiH with Ph₃C⁺[B(C₆F₅)₄] again resulted in decomposition of the borate counteranion (entry 5).17 Conversely, treatment of Me₂PhSiH with trityl carborane Ph₃C⁺[CHB₁₁H₅Br₆] exclusively afforded silylium salt Me₃-Si⁺[CHB₁₁H₅Br₆] without detectable formation of MePh₂Si⁺ or Me₂PhSi⁺ (entry 6). Strikingly, hydride abstraction from sterically more demanding Me₂(C₆Me₅)SiH led to the corresponding triarylsilylium ion in the presence of the borate counteranion (entry 7), while substituent redistribution into the 'opposite direction' was induced by the carborane anion,

now affording Me₃Si[†][CHB₁₁H₅Br₆]⁻ (entry 8).¹⁹ However, heating of the reaction at 50 °C for 72 h was necessary.

Overall, these results indicate that hydride abstraction from hydrosilanes of type Me_2ArSiH with a carborane-based trityl salt tends to form the trimethylsilylium ion, whereas hydrosilanes of type $MeAr_2SiH$ with a bulky aryl substituent favor triarylsilylium ion generation.

Mechanism of the substituent redistribution reaction with Me₂PhSiH

To gain insight into the reaction mechanism and to understand why the treatment of Me₂PhSiH with Ph₃C⁺[CHB₁₁H₅Br₆]⁻ exclusively gives Me₃Si⁺[CHB₁₁H₅Br₆]⁻, we constructed a complete reaction energy profile using DFT calculations at the M06/cc-pVTZ(-f)//6-31G** level of theory (Fig. 3; see the ESI† for details of the computational method).20 The hydride abstraction from Me₂PhSiH with Ph₃C⁺[CHB₁₁H₅Br₆] was found to have a barrier of 15.5 kcal mol⁻¹ and is therefore expected to occur rapidly at room temperature (not shown). In the condensed phase, the resulting silvlium ion Me₂PhSi⁺ (6A), which is located at a relative free energy of 6.5 kcal mol⁻¹, is stabilized through coordination by the solvent, another hydrosilane molecule, or by the counteranion (see the ESI† for a comparison of the association energies).8e,21 Coordination of one of the bromine atoms of the carborane counteranion to the silicon cation results in the highest binding energy, and the resulting ion pair 6A' is predicted to be at a relative free energy of $-24.1~
m{kcal}~
m{mol}^{-1}$. Silylium ion $6A~
m{can}$ also interact with another equivalent of Me2PhSiH to form hydride-bridged adduct 7A,21

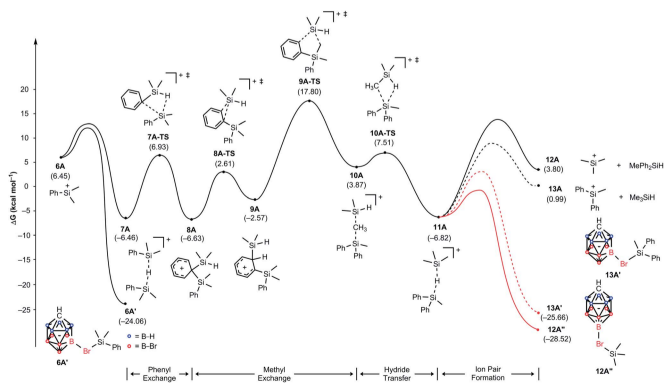


Fig. 3 Energy (kcal mol^{-1}) profile of the substituent redistribution in the reaction of Me₂PhSiH (1A) with Ph₃C⁺[CHB₁₁H₅Br₆]⁻ (2A). The energies are relative to the starting materials 1A and 2A.

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located at -6.5 kcal mol⁻¹. Note that these energies are not adjusted for the different concentrations of the components and assume normal conditions. Given that Me₂PhSiH (1A) is present in excess, these normal energies suggest that adduct 7A will be encountered easily in significant quantities.

Hydride-bridged adduct ion 7A can undergo a phenyl group transfer to arrive at phenyl-bridged adduct 8A7c,8b,22 via the fourcentered transition state 7A-TS, associated with a barrier of 13.4 kcal mol⁻¹. Surprisingly, the subsequent methyl group transfer does not proceed via another typical four-membered transition state.²³ Instead, our calculations suggest that 1,2migration of the silicon group in 8A occurs via the low barrier transition state 8A-TS, leading to ortho-disilylated arenium ion 9A. This seemingly unfavorable intermediate is only 4.1 kcal mol⁻¹ higher in energy than arenium ion 8A. Finally, 9A facilitates the exchange of one methyl group, passing through five-centered transition state 9A-TS with an overall barrier of 24.3 kcal mol⁻¹ relative to 7A. This energetically most demanding reaction step forms methonium ion 10A, which is metastable and rapidly rearranges to hydridebridged adduct 11A via low barrier transition state 10A-TS. The hydrosilane-stabilized silvlium ions 7A and 11A are almost isoenergetic ($\Delta G = 0.4 \text{ kcal mol}^{-1}$), suggesting that both structures coexist in equilibrium. The formal dissociation of 11A gives either Me₃Si⁺ or MePh₂Si⁺, the former being calculated to be 2.8 kcal mol⁻¹ higher in energy. However, coordination by the carborane anion changes the energy landscape decisively, as ion pair formation reverses the energy ordering. $Me_3Si^+[CHB_{11}H_5Br_6]^-$ (12A"), which is located at -28.5 kcal mol⁻¹, is 2.9 kcal mol⁻¹ lower in energy than MePh₂Si⁺[CHB₁₁H₅Br₆] $^-$ (13A') and also 4.5 kcal mol⁻¹ more stable than $Me_2PhSi^+[CHB_{11}H_5Br_6]^-$ (6A'), thus predicting the silylium salt 12A" as the major product of the substituent redistribution reaction.

It should be noted that silylium ions are significantly more stabilized by coordination of the carborane counteranion than by formation of solvent adducts such as $R_3\text{-}Si(\text{toluene})^{\dagger}[\text{CHB}_{11}\text{H}_5\text{Br}_6]^-$. Moreover, the energy differences between these arenium ions are small, predicting a mixture of different silylium ions in the absence of the carborane counteranion (see the ESI† for details). This result was supported by independent control experiments (Scheme 2). The hydride abstraction from Me_2PhSiH with borate-based trityl salt $\text{Ph}_3\text{C}^{\dagger}[\text{B}(\text{C}_6\text{F}_5)_4]^-$ was repeated but stopped after stirring for 10 min in toluene (cf. Table 1, entry 5). NMR spectroscopic analysis of the polar phase in o-Cl_2C_6D_4 revealed the formation of a mixture of Me_3Si^{\dagger}[\text{B}(\text{C}_6\text{F}_5)_4]^- and Me_2PhSi^{\dagger}[\text{B}(\text{C}_6\text{F}_5)_4]^- in a ratio of \sim 51 : 49 along with small amounts of byproducts arising from counteranion

Scheme 2 Influence of the counteranion on the selectivity of the trimethylsilylium ion formation.

decomposition. In contrast, stopping the reaction of Me_2 -PhSiH with $Ph_3C^{\dagger}[CHB_{11}H_5Br_6]^{-}$ after stirring for 10 min in toluene furnished $Me_3Si^{\dagger}[CHB_{11}H_5Br_6]^{-}$ as the major product along with only small amounts of unscrambled Me_2 -PhSi $^{\dagger}[CHB_{11}H_5Br_6]^{-}$ (ratio \sim 84 : 16). In both reactions, full conversion of the trityl salt was observed.

As shown in Fig. 4, the silylium ions can be bound either to the apical or one of the equatorial bromine atoms of the carborane counteranion, with a slight preference of 1.1 kcal mol⁻¹ for the apical position in $Me_3Si^+[CHB_{11}H_5Br_6]^-$ (12A"). This result is in contrast to the molecular structure in the solid state, which shows the equatorial isomer (cf. Fig. 2). We speculate that either packing effects or a statistical preference for the equatorial isomer is the reason for this discrepancy. Notably, the equatorial isomer 12A' is still 1.8 kcal mol⁻¹ lower in energy than the equatorial isomer of MePh₂Si⁺[CHB₁₁H₅Br₆]⁻ (13A'). The higher ion pairing energy in 12A' can be ascribed to the low steric demand of Me₃Si⁺, leading to a closer carborane coordination and to attractive van der Waals interactions between the methyl moieties and the carborane anion. Especially in the apical position, the methyl functionality can interact with the highly polarizable bromine atoms. In contrast, the molecular fit of the sterically more demanding silylium ions Me₂PhSi⁺ (6A) and MePh₂Si⁺ (13A) with the carborane counteranion is less

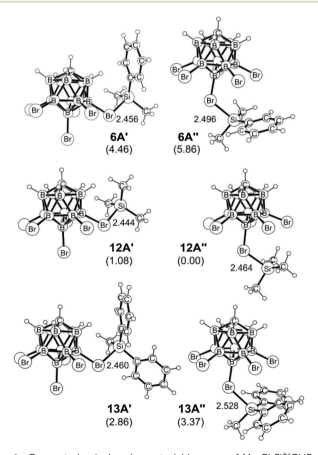


Fig. 4 Computed apical and equatorial isomers of Me₂PhSi⁺[CHB₁₁-H₅Br₆]⁻ (top), Me₃Si⁺[CHB₁₁H₅Br₆]⁻ (middle) and MePh₂Si⁺[CHB₁₁H₅-Br₆]⁻ (bottom). Si–Br bond lengths are given in Å and relative free energy differences (kcal mol⁻¹) are shown in parentheses.

tight, and the ion pairing is therefore slightly less favorable. This trend is reflected in the corresponding Si–Br bond lengths of these silylium carborane salts, which were computed to be shortest in both isomers of $Me_3Si^+[CHB_{11}H_5Br_6]^-$ (12A′ and 12A″). Hence, this ion pair is the most stable silylium salt despite the lack of stabilizing phenyl groups. Both isomers of $Me_2PhSi^+[CHB_{11}H_5Br_6]^-$ (6A′ and 6A″) are higher in energy than the corresponding $MePh_2Si^+[CHB_{11}H_5Br_6]^-$ (13A′ and 13A″), indicating that the stabilization of these silylium carborane salts is determined by a delicate balance of electronic and steric effects. It should also be noted here that the DFT optimized structures for $Me_3Si^+[CHB_{11}H_5Br_6]^-$ (12A′) and $MePh_2Si^+[CHB_{11}H_5Br_6]^-$ (13A′) are in good agreement with the corresponding molecular structures obtained by X-ray diffraction analysis (see the ESI† for details).

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Mechanism of the substituent redistribution reaction with MePh₂SiH

To understand why the reaction of MePh₂SiH with Ph₃-C[†][CHB₁₁H₅Br₆]⁻ does not furnish Me₃Si[†][CHB₁₁H₅Br₆]⁻, we constructed again a complete energy profile employing DFT simulations (Fig. 5). The initial hydride transfer of the hydrosilane to the trityl cation has a calculated barrier of 14.3 kcal mol⁻¹ (not shown), which is 1.2 kcal mol⁻¹ lower in energy compared to the case of Me₂PhSiH due to the slightly higher hydride donor strength of MePh₂SiH (see Table S1 in the

ESI† for details). The resulting silylium ion MePh₂Si⁺ (6B) with a relative free energy of 0.8 kcal mol⁻¹ is almost isoenergetic to the reactant state. Adduct formation with another equivalent of MePh₂SiH affords hydrosilane-stabilized silylium ion 7B, which undergoes a phenyl/methyl exchange reaction following a very similar reactivity pattern as described above, leading to scrambled hydride-bridged adduct 11B. The transformation of 7B to 11B via intermediates 8B, 9B, and 10B is again reversible, since 7B and 11B have similar free energies ($\Delta G =$ $0.7 \text{ kcal mol}^{-1}$). As before, the methyl group transfer *via* fivemembered transition state 9B-TS shows the highest barrier, which is 24.2 kcal mol⁻¹ relative to 7B. In this equilibrium, unscrambled MePh₂Si⁺[CHB₁₁H₅Br₆]⁻ (**6B**') with a relative free energy of -25.9 kcal mol⁻¹ is predicted to be the major species, followed by scrambled Me₂PhSi⁺[CHB₁₁H₅Br₆]⁻ (12B') and $Ph_3Si^+[CHB_{11}H_5Br_6]^-$ (13B"), which are basically isoenergetic at -24.6 kcal mol⁻¹ and -24.7 kcal mol⁻¹, respectively. This finding is in good agreement with the experimental observation of unscrambled MePh₂Si⁺[CHB₁₁H₅Br₆] being the main product of the reaction (cf. Table 1, entry 4).25

Our calculations suggest that a subsequent methyl exchange reaction leading to Me_3Si^+ is unlikely ($\mathbf{11B} \to \mathbf{18B}$, gray energy profile in Fig. 5). The transition state for this methyl group transfer, $\mathbf{16B}$ -TS, is located 26.7 kcal mol^{-1} relative to $\mathbf{11B}$, which is 1.8 kcal mol^{-1} higher in energy than the barrier of the backward reaction via transition state $\mathbf{9B}$ -TS. Consequently, the

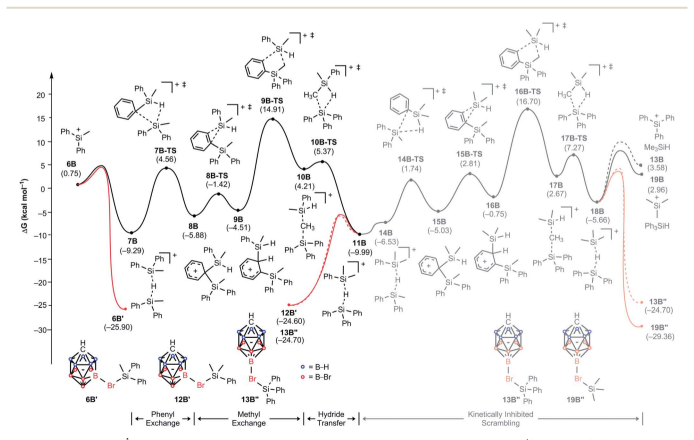


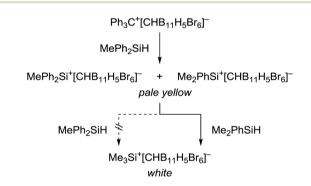
Fig. 5 Energy (kcal mol^{-1}) profile of the substituent redistribution in the reaction of MePh₂SiH (1A) with Ph₃C⁺[CHB₁₁H₅Br₆]⁻ (2B). The energies are relative to the starting materials 1B and 2B.

reaction of MePh₂SiH with Ph₃C[†][CHB₁₁H₅Br₆]⁻ stops at the above-mentioned mixture of silicon cations rather than undergoing exhaustive substituent redistribution to furnish low energy Me₃Si[†][CHB₁₁H₅Br₆]⁻.

This kinetic inhibition was further proven by another mechanistic control experiment (Scheme 3). When a mixture of $\mathrm{Ph_3C^+[CHB_{11}H_5Br_6]^-}$ and $\mathrm{MePh_2SiH}$ in toluene was stirred overnight at room temperature, a pale yellow suspension was obtained, which is characteristic of silylium ions with aromatic substituents (*cf.* Table 1, entry 4). Addition of less bulky $\mathrm{Me_2-PhSiH}$ to this mixture resulted in a quick decolorization and formation of a white suspension. NMR spectroscopic analysis of the solid now confirmed exclusive formation of $\mathrm{Me_3Si^+[CHB_{11}H_5Br_6]^-}$.

Scope of the substituent redistribution reaction

The hydride abstraction from various dialkyl(phenyl)silanes with $Ph_3C^{\dagger}[CHB_{11}H_5Br_6]^{-}$ finally revealed that the redistribution reaction is not restricted to methyl groups (Table 2). Although Et_2PhSiH reacted much slower compared to $Me_2-PhSiH$, exclusive formation of trialkylsilylium ion $Et_3-Si^{\dagger}[CHB_{11}H_5Br_6]^{-}$ was observed (entries 1 and 2). Employing more bulky iPr_2PhSiH led to clean generation of unscrambled dialkyl(aryl)silylium ion $iPr_2PhSi^{\dagger}[CHB_{11}H_5Br_6]^{-}$, as verified by X-ray crystallography (entry 3; see the ESI^{\dagger} for the molecular



Scheme 3 Probing the kinetic inhibition in the substituent redistribution reaction with $MePh_2SiH$.

Table 2 Silylium ion generation from hydrosilanes of type R₂PhSiH

	R ₂ PhSiH (2 equiv)	tol		·[CHB ₁₁ H ₅ Br ₆] ⁻ or si ⁺ [CHB ₁₁ H ₅ Br ₆] ⁻
Entry		R	Si ⁺	$\delta(^{29}\mathrm{Si})^b$ [ppm]
1		Me	Me ₃ Si ⁺	93
2^c		Et	$\mathrm{Et_{3}Si}^{^{+}}$	100
3		iPr	iPr_2PhSi^+	76
4		<i>t</i> Bu	d	_

 $[^]a$ All reactions were performed according to GP 2. See the ESI for details. b Measured in o-Cl₂C₆D₄. c With 4 equiv. of Et₂PhSiH and 7 days reaction time. d No reaction; only Ph₃C[†][CHB₁₁H₅Br₆] was recovered.

Table 3 Silylium ion generation from hydrosilanes of type Me₂RSiH

	Me ₂ RSiH (2 equiv)	t	CHB ₁₁ H ₅ Br ₆] oluene overnight	$ m Me_3Si^{\dagger}[CHB_{11}H_5Br_6]^{-}$ or $ m Me_2RSi^{\dagger}[CHB_{11}H_5Br_6]^{-}$
Entry		R	Si ⁺	$\delta(^{29}\mathrm{Si})^b$ [ppm]
1 2		Ph Bn	Me_3Si^+ Me_3Si^+	93
3		<i>t</i> Bu	Me ₃ si Me ₂ tBu	93 Si ⁺ 98

^a All reactions were performed according to GP 2. See the ESI for details. ^b Measured in o-Cl₂C₆D₄.

structure of $iPr_2PhSi^+[CHB_{11}H_5Br_6]^-$).¹⁵ These results are in accordance with our calculations, predicting high energy barriers for the transfer of bulky alkyl groups. Sterically even more shielded tBu_2PhSiH then completely thwarted the hydride abstraction, and only the trityl salt was recovered from the reaction mixture (entry 4).

To investigate whether the phenyl group in Me₂PhSiH can be replaced by other 'leaving groups', we also tested a benzyl and an alkyl substituent in Me₂RSiH (Table 3). As in the case of Me₂PhSiH (entry 1), clean formation of Me₃Si⁺[CHB₁₁H₅Br₆]⁻ was observed with Me₂BnSiH (entry 2), showing that the phenyl group is not essential for the exchange process. In contrast, the bulky *tert*-butyl group in Me₂tBuSiH completely prevented substituent redistribution, and silylium ion Me₂tBuSi⁺[CHB₁₁-H₅Br₆]⁻ was formed as the only product (entry 3). This result again demonstrates that the intermolecular substituent exchange reaction is sensitive towards sterically demanding alkyl groups (*cf.* entry 3 in Table 2).

Conclusion

It has been known for decades that silylium ions can undergo redistribution reactions of their substituents. The present combined experimental and detailed computational study finally provides a full mechanistic picture of this phenomenon. The mechanism involves a series of phenyl and alkyl exchange reactions, the latter being calculated to be the energetically most demanding steps. While the transfer of phenyl groups proceeds *via* common four-centered transition states, the corresponding alkyl exchange was found to pass through unusual five-membered transition states. These are accessible after 1,2-silyl migration at the stage of the intermediate disilylated arenium ions.

Additionally, our DFT calculations revealed that the silicon cations are significantly more stabilized by ion pair formation with the carborane counteranion ($R_3Si^+[CHB_{11}H_5Br_6]^-$) than by formation of toluenium ($R_3Si(toluene)^+[CHB_{11}H_5Br_6]^-$) or hydrosilane-stabilized silylium ions ([$R_3Si-H-SiR_3$] $^+[CHB_{11}H_5-Br_6]^-$). More importantly, purely aliphatic silylium carboranes with small substituents, *i.e.*, methyl or ethyl groups, were found to be distinctly lower in energy than the corresponding mixed aliphatic/aromatic or purely aromatic silylium ion pairs as

a result of stronger attractive interactions ($\Delta G \ge 2.9 \text{ kcal mol}^{-1}$ for R = Me). These energy differences account for the highly selective formation of $Me_3Si^+[CHB_{11}H_5Br_6]^-$ and $Et_3Si^+[CHB_{11}H_5Br_6]^-$ from the reaction of the corresponding hydrosilanes R_2PhSiH (R = Me, Et) with $Ph_3C^+[CHB_{11}H_5Br_6]^-$ under thermodynamic control.

The phenyl group in Me₂PhSiH turned out to be replaceable by other 'leaving groups', such as a benzyl or even a sterically demanding C₆Me₅ group. However, two alkyl groups must be preinstalled in the hydrosilane starting material to steer the reaction towards formation of Me₃Si⁺[CHB₁₁H₅Br₆]⁻. In contrast, hydride abstraction from MePh₂SiH with only one alkyl substituent leads to a mixture of different silylium ions, as exhaustive scrambling to Me₃Si⁺ is kinetically inhibited. Exchanging the phenyl groups in MePh₂SiH by 2,6-disubstituted aryl groups (*e.g.* C₆Me₅) eventually provides access to sterically congested triarylsilylium ions, as previously demonstrated by Müller and co-workers.¹⁰

These general trends provide a solid foundation for the mechanistic understanding of the substituent redistribution of silylium ions, thereby enabling the prediction of the outcome of these exchange reactions. Thus, this process can be used as a reliable synthetic route not only to triaryl- but also to trialkylsilylium ions by deliberate choice of the hydrosilane and counteranion of the trityl salt.

Conflicts of interest

Chemical Science

There are no conflicts to declare.

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Notes and references

- 1 For general reviews of silylium ion chemistry, see: (*a*) V. Y. Lee and A. Sekiguchi, in *Organosilicon Compounds*, ed. V. Y. Lee, Academic Press, Oxford, 2017, vol. 1, pp. 197–230; (*b*) T. Müller, in *Structure and Bonding*, ed. D. Scheschkewitz, Springer, Berlin, 2014, vol. 155, pp. 107–162; (*c*) T. Müller, in *Science of Synthesis Knowledge Updates* 2013/3, ed. M. Oestreich, Thieme, Stuttgart, 2013, pp. 1–42.
- 2 For silylium ions in catalysis, see: (a) H. F. T. Klare, ACS Catal., 2017, 7, 6999–7002; (b) A. Schulz and A. Villinger, Angew. Chem., Int. Ed., 2012, 51, 4526–4528; (c) H. F. T. Klare and M. Oestreich, Dalton Trans., 2010, 39, 9176–9184
- 3 (a) P. D. Bartlett, F. E. Condon and A. Schneider, *J. Am. Chem. Soc.*, 1944, **66**, 1531–1539; (b) J. Y. Corey and R. West, *J. Am.*

- Chem. Soc., 1963, **85**, 2430–2433; (c) J. Y. Corey, J. Am. Chem. Soc., 1975, **97**, 3237–3238.
- 4 For recent reviews of weakly coordinating anions, see: (a) I. M. Riddlestone, A. Kraft, J. Schaefer and I. Krossing, *Angew. Chem., Int. Ed.*, 2018, 57, DOI: 10.1002/anie.201710782; (b) T. A. Engesser, M. R. Lichtenthaler, M. Schleep and I. Krossing, *Chem. Soc. Rev.*, 2016, 45, 789–899.
- 5 For further strategies to generate silylium ions, see: (a) J. B. Lambert, Y. Zhao, H. Wu, W. C. Tse and B. Kuhlmann, J. Am. Chem. Soc., 1999, 121, 5001–5008 (allyl-leaving-group approach); (b) M. J. MacLachlan, S. C. Bourke, A. J. Lough and I. Manners, J. Am. Chem. Soc., 2000, 122, 2126–2127 (ring-opening protonolysis); (c) A. Schäfer, M. Reißmann, A. Schäfer, M. Schmidtmann and T. Müller, Chem.-Eur. J., 2014, 20, 9381–9386 (silylene protonation); (d) A. Simonneau, T. Biberger and M. Oestreich, Organometallics, 2015, 34, 3927–3929 (cyclohexadienyl-leaving-group approach); (e) Q.-A. Chen, H. F. T. Klare and M. Oestreich, J. Am. Chem. Soc., 2016, 138, 7868–7871 (hydrosilane protonation).
- 6 For a review of substituent redistribution reactions at silicon, see: D. R. Weyenberg, L. G. Mahone and W. H. Atwell, *Ann. N. Y. Acad. Sci.*, 1969, **159**, 38–55.
- 7 For Lewis acid-catalyzed substituent redistribution reactions of hydrosilanes, see: (a) J. L. Speier and R. E. Zimmerman, J. Am. Chem. Soc., 1955, 77, 6395-6396; (b) M. Khandelwal and R. J. Wehmschulte, Angew. Chem., Int. Ed., 2012, 51, 7323-(c) A. Feigl, I. Chiorescu, K. Deller, S. U. H. Heidsieck, M. R. Buchner, V. Karttunen, A. Bockholt, A. Genest, N. Rösch and B. Rieger, Chem.-Eur. J., 2013, 19, 12526-12536; (d) R. J. Wehmschulte, M. Saleh and D. R. Powell, Organometallics, 2013, 32, 6812-6819; (e) R. Labbow, F. Reiß, A. Schulz and A. Villinger, Organometallics, 2014, 33, 3223-3226; (f) J. Chen and E. Y.-X. Chen, Angew. Chem., Int. Ed., 2015, 54, 6842-6846; (g) Y. Ma, L. Zhang, Y. Luo, M. Nishiura and Z. Hou, J. Am. Chem. Soc., 2017, 139, 12434-12437.
- 8 For substituent redistribution reactions of silicon cations, see: (a) C. Eaborn, P. D. Lickiss, S. T. Najim and W. A. Stańczyk, J. Chem. Soc., Chem. Commun., 1987, 1461–1462; (b) N. Choi, P. D. Lickiss, M. McPartlin, P. C. Masangane and G. L. Veneziani, Chem. Commun., 2005, 6023–6025; (c) N. Lühmann, H. Hirao, S. Shaik and T. Müller, Organometallics, 2011, 30, 4087–4096; (d) K. Müther, P. Hrobárik, V. Hrobáriková, M. Kaupp and M. Oestreich, Chem.-Eur. J., 2013, 19, 16579–16594; (e) S. J. Connelly, W. Kaminsky and D. M. Heinekey, Organometallics, 2013, 32, 7478–7481; (f) Ref. 7e; (g) L. Albers, S. Rathjen, J. Baumgartner, C. Marschner and T. Müller, J. Am. Chem. Soc., 2016, 138, 6886–6892.
- For seminal reports, see: (a) J. B. Lambert, S. Zhang,
 C. L. Stern and J. C. Huffman, *Science*, 1993, 260, 1917–1918; (b) C. A. Reed, Z. Xie, R. Bau and A. Benesi, *Science*, 1993, 262, 402–404.
- 10 (a) A. Schäfer, M. Reißmann, A. Schäfer, W. Saak, D. Haase and T. Müller, Angew. Chem., Int. Ed., 2011, 50, 12636-

Edge Article

12638; (b) A. Schäfer, M. Reißmann, S. Jung, A. Schäfer, W. Saak, E. Brendler and T. Müller, Organometallics, 2013,

- 32, 4713-4722.
- 11 (a) J. B. Lambert and Y. Zhao, Angew. Chem., Int. Ed. Engl., 1997, **36**, 400–401; (b) K.-C. Kim, C. A. Reed, D. W. Elliot, L. J. Mueller, F. Tham, L. Lin and J. B. Lambert, Science, 2002, 297, 825-827; (c) J. B. Lambert and L. Lin, J. Org. Chem., 2001, 66, 8537-8539.
- 12 Hydride abstraction from MePh₂SiH with Ph₃C⁺[B(C₆F₅)₄] was reported as a clean reaction: J. B. Lambert, S. Zhang and S. M. Ciro, Organometallics, 1994, 13, 2430-2443. However, this result could not be reproduced by Müller (cf. ref. 10b) and us.
- 13 (a) C. A. Reed, Acc. Chem. Res., 1998, 31, 133-139; (b) C. A. Reed, Acc. Chem. Res., 2010, 43, 121-128.
- 14 For the synthesis and crystallographic characterization of related trimethylsilvlium salts, see: (a) Me₃Si⁺[CRB₁₁F₁₁] (R = H, Et): T. Küppers, E. Bernhardt, R. Eujen, H. Willner and C. W. Lehmann, Angew. Chem., Int. Ed., 2007, 46, 6346-6349; (b) $Me_3Si(arene)^+[B(C_6F_5)_4]^-$: M. F. Ibad, P. Langer, A. Schulz and A. Villinger, J. Am. Chem. Soc., 2011, 133, 21016-21027.
- 15 CCDC 1818576 for Me₃Si⁺[CHB₁₁H₅Br₆]⁻, CCDC 1818582 for CCDC $MePh_2Si^+[CHB_{11}H_5Br_6]^-$, and 1818581 iPr₂PhSi⁺[CHB₁₁H₅Br₆] contain the supplementary crystallographic data for this paper.†
- 16 Z. Xie, R. Bau, A. Benesi and C. A. Reed, Organometallics, 1995, 14, 3933-3941.
- 17 The decomposition of the $[B(C_6F_5)_4]^-$ counteranion is likely to proceed via an SEAr reaction of the formed silylium ions with the borate. The formation of $B(C_6F_5)_3$ was verified by ¹⁹F NMR spectroscopic analysis, and GC-MS analysis revealed formation of several silanes containing a C₆F₅ unit.

- 18 The generated silylium ions were converted to the corresponding fluorosilanes using $(C_6F_5)_3PF_2$ (1.0 equiv.), thereby facilitating product characterization by both NMR spectroscopic and GC-MS analysis. For the preparation of $(C_6F_5)_3PF_2$, see: C. B. Caputo, L. J. Hounjet, R. Dobrovetsky and D. W. Stephan, Science, 2013, 341, 1374-1377.
- 19 Small amounts of the triarvlsilvlium ion $(C_6Me_5)_3Si^+[CHB_{11}H_5Br_6]^-$ were also detected (cf. ref. 10).
- 20 The mechanism of intermolecular substituent exchange reactions at related ferrocene-stabilized silylium ions had already been studied by quantum-chemical analyses (cf. ref. 8d). However, the calculated barriers for the transition states were relatively high. For the calculated mechanism of an intramolecular substituent exchange reaction at silicon cation with a rigid naphthalene-1,8-divl backbone, see: ref. 8c.
- 21 (a) S. P. Hoffmann, T. Kato, F. S. Tham and C. A. Reed, Chem. Commun., 2006, 767-769; (b) M. Nava and C. A. Reed, Organometallics, 2011, 30, 4798-4800.
- 22 R. Meyer, K. Werner and T. Müller, Chem.-Eur. J., 2002, 8, 1163-1171.
- 23 We were not able to locate a four-centered transition state from 8A to directly arrive at 10A. See Fig. S67 in the ESI† for geometric scan calculations.
- 24 Me₃Si(toluene)⁺[CHB₁₁H₅Br₆]⁻ was calculated to be only mol^{-1} energy 0.7 kcal lower in than $MePh_2Si(toluene)^{+}[CHB_{11}H_5Br_6]^{-}$ (see the ESI† for details).
- 25 Although our calculations predict formation of small amounts of Ph₃Si⁺[CHB₁₁H₅Br₆] in the reaction of MePh₂SiH with Ph₃C⁺[CHB₁₁H₅Br₆]⁻, we were not able to detect this silylium ion by ¹H/²⁹Si HMQC NMR spectroscopy.