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Detailed quasiclassical dynamics of the $\text{F}^- + \text{SiH}_3\text{Cl}$ multi-channel reaction†

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We report a detailed quasiclassical trajectory study on the $\text{F}^- + \text{SiH}_3\text{Cl}$ multi-channel reaction using a full-dimensional *ab initio* analytical potential energy surface. Reaction probabilities, cross sections, initial attack and scattering angle distributions as well as product relative translational, internal, vibrational, and rotational energy distributions are obtained in the collision energy range of 1–40 kcal mol⁻¹ for the following channels: $\text{SiH}_3\text{F} + \text{Cl}^-$, $\text{SiH}_2\text{Cl}^- + \text{HF}$, $\text{SiH}_2\text{F}^- + \text{HCl}$, $\text{SiH}_2\text{FCl} + \text{H}^-$, $\text{SiH}_2 + \text{FHCl}^-$, and $\text{SiHFCl}^- + \text{H}_2$. All the channels are translationally cold indicating indirect mechanisms, except proton transfer ($\text{SiH}_2\text{Cl}^- + \text{HF}$), which shows mixed direct–indirect character. The angular distributions vary depending on collision energy and inversion/retention for $\text{SiH}_3\text{F} + \text{Cl}^-$. In the case of $\text{SiH}_2\text{Cl}^- + \text{HF}$ front-side/back-side attack backward-forward/forward scattering preference is found at low/high collision energy. $\text{SiH}_2\text{F}^- + \text{HCl}$ is formed with isotropic scattering and the preferred angle of attack is similar to the $\text{SiH}_3\text{F} + \text{Cl}^-$ channel. $\text{SiH}_2\text{FCl} + \text{H}^-/\text{SiH}_2 + \text{FHCl}^-$ favors back-side attack and isotropic/backward scattering, whereas $\text{SiHFCl}^- + \text{H}_2$ does not show significant angular preference.

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1. Introduction

The dynamics and mechanisms of bimolecular nucleophilic substitution ($\text{S}_{\text{N}}2$) reactions of halide ions (X^-) with methyl-halide molecules (CH_3Y) have been extensively investigated,^{1–8} however, analogous $\text{S}_{\text{N}}2$ reactions at silicon (Si) centers are less thoroughly studied. Previous theoretical work on the $\text{H}^-/\text{F}^- + \text{SiH}_4$ and $\text{X}^- + \text{SiH}_3\text{Y}$ systems was restricted to stationary-point characterizations^{9–19} and on-the-fly direct dynamics simulations^{20–22} until 2023 when we developed a full-dimensional coupled-cluster-quality analytical potential energy surface (PES) for the $\text{F}^- + \text{SiH}_3\text{Cl}$ reaction,²³ which, to the best of our knowledge, was the first PES for a Si-centered $\text{S}_{\text{N}}2$ reaction. The analytical PES allows the efficient computations of a large number of quasiclassical trajectories (QCTs) providing statistically-converged cross sections and unprecedented insights into the mechanisms of the system of interest. The QCT simulations on the new PES showed that the $\text{F}^- + \text{SiH}_3\text{Cl}$ $\text{S}_{\text{N}}2$ reaction is not stereospecific, because the Walden-inversion and front-side attack retention pathways are competitive, especially at high collision energies.²³ Besides $\text{S}_{\text{N}}2$, our

trajectories uncovered several other product channels such as $\text{SiH}_2\text{Cl}^- + \text{HF}$, $\text{SiH}_2\text{F}^- + \text{HCl}$, $\text{SiH}_2\text{FCl} + \text{H}^-$, $\text{SiH}_2 + \text{FHCl}^-$, $\text{SiHFCl}^- + \text{H}_2$, $\text{SiHF} + \text{H}_2 + \text{Cl}^-$, and $\text{SiH}_2 + \text{HF} + \text{Cl}^-$, showing that the dynamics of Si-centered ion–molecule reactions is even more complex than that of the analogous C-centered systems.²³

In our previous work²³ we focused on the challenges and methodology of the PES development, the evaluation of the accuracy and performance of the analytical PES, and the atomic-level mechanisms of the different reaction channels uncovered by inspecting and analyzing several representative trajectories. In the present study we plan to report a more detailed QCT analysis of the $\text{F}^- + \text{SiH}_3\text{Cl}$ reaction computing differential cross sections such as initial attack angle and scattering angle distributions as well as relative translational, internal, vibrational, and rotational energy distributions for the products. These detailed differential cross section data, for the first time, provide new insights into the dynamics and mechanisms of the different channels of the title reaction, showing the spatial preference of the reactive collisions as well as how the energy transfers from reactants to the various degrees of freedom of the products. In Section 2 we give the methodological and computational details. In Section 3 the results and discussion are given, first introducing the characteristic of the PES, then providing the reaction probabilities and cross sections of the different channels, describing the angular and post-reaction energy distributions, and comparing the present findings to those obtained for the analogous carbon- and phosphorous-centered systems. The paper ends with summary and conclusions in Section 4.

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† Electronic supplementary information (ESI) available: Branching ratios as well as $E_{\text{trans}}/E_{\text{tot}}$, $E_{\text{int}}/E_{\text{tot}}$, $E_{\text{vib}}/E_{\text{tot}}$, and $E_{\text{rot}}/E_{\text{tot}}$ fraction and integration-time distributions for the different channels of the title reaction. See DOI: <https://doi.org/10.1039/d4cp00048j>



2. Methods and computational details

QCT simulations were performed on a full-dimensional analytical composite coupled-cluster-quality PES,²³ which is represented by a permutationally invariant polynomial²⁴ fitted to (CCSD-F12b + BCCD(T)–BCCD)/aug-cc-pVTZ energy points, for the $\text{SiH}_3\text{Cl} + \text{F}^-$ system, across various collision energies (1, 5, and up to 40 kcal mol⁻¹ in 5 kcal mol⁻¹ increments) and impact parameters. The impact parameter, b , ranged from 0 to b_{max} , and the b_{max} values, where the reactive trajectories vanish, ranged from 9.5 to 25 bohr, depending on the collision energy (E_{coll}). At each E_{coll} and b 5000 trajectories were run.

For every simulation, the initial vibrational energy was set to the zero-point energy of SiH_3Cl via normal-mode sampling and its rotational energy was adjusted to zero. We arrange the initial center of mass distance between the reactants by the $\sqrt{x^2 + b^2}$ formula, where x was 40 bohr at $E_{\text{coll}} = 1$ and 5 kcal mol⁻¹, and 25 bohr at higher collision energies.

A more detailed explanation of the fitting method, the creation of the initial fitting set, the iterative PES development and the quality assessment of the final PES is given in our previous article.²³ The computational details of the QCT product analysis are described below.

We define the α initial attack angle as the angle between the center-of-mass velocity vector of the SiH_3Cl reactant, $\mathbf{v}_{\text{SiH}_3\text{Cl}}^{\text{com}}$, and the vector of the spatial displacement between its Cl ligand and the central atom, $\mathbf{x}_{\text{Cl}} - \mathbf{x}_{\text{Si}}$, obtained at the initial trajectory step, as depicted in Fig. 1a. The θ scattering angle is the angle between the center-of-mass velocity vectors of the reactant fluoride ion at the initial trajectory step, $\mathbf{v}_{\text{F}^-}^{\text{com}}$, and one of the resulting products at the final step of the simulation. For each product channel, we employed the velocity coordinates of the product containing the fluorine atom to compute the scattering angles. Fig. 1b illustrates θ for the chloride substitution channel, where we track the SiH_3F product. It is important to note that in the case of a product pair, owing to the law of conservation of momentum and the initial center-of-mass velocities of the reactants being set in parallel, the velocity vectors of the products are always parallel as well. Consequently, the scattering angle defined for one product complements the scattering angle defined for the other product to 180 degrees.

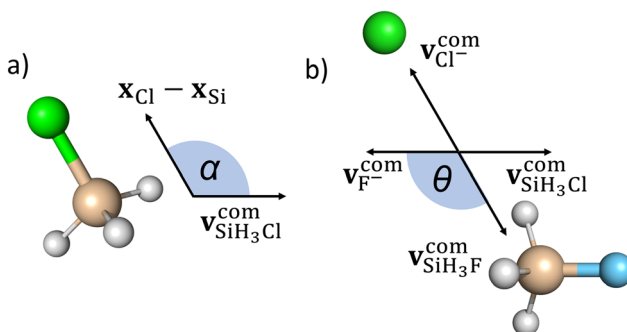


Fig. 1 Illustration of the α initial attack angle (a) and the θ scattering angle (b).

The initial attack angle distribution is obtained by partitioning the cosine of α into equally spaced bins ranging from -1 to 1 . In this scheme, -1 ($\alpha = 180^\circ$) corresponds to back-side attack, where the central atom of SiH_3Cl faces the F^- ion as the reactants approach each other. On the other hand, when $\cos(\alpha) = 1$ ($\alpha = 0^\circ$), it signifies a front-side attack, indicating that the chlorine ligand of SiH_3Cl faces the approaching F^- ion. We follow a similar approach to obtain the scattering angle distributions by dividing $\cos(\theta)$ into evenly spaced bins between -1 and 1 . In this context, a value of -1 ($\theta = 180^\circ$) corresponds to backward scattering, while $\cos(\theta) = 1$ signifies forward scattering.

The number of bins of the distributions of both attack and scattering angles, as well as energy distributions was chosen based on the reaction probabilities. We utilized 20 bins for chloride substitution and proton abstraction, 10 bins for HCl production, and 5 bins for the less likely hydride substitution, molecular hydrogen production, and bihalide ion production channels.

3. Results and discussion

3.1 Potential energy surface

Fig. 2 illustrates the reaction channels of the $\text{SiH}_3\text{Cl} + \text{F}^-$ system, alongside the classical (adiabatic) energies of the stationary points and the products obtained on the PES, as reported in ref. 23. The PES describes six $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ and one $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B} + \text{C}$ type reactions ($\text{SiHF} + \text{H}_2 + \text{Cl}^-$) with reasonable accuracy. Chloride substitution occurs through two primary mechanisms: the barrierless Walden inversion, featuring a minimum (WMIN), and the retention channel with a deep double-well of a transition state (FSTS), a reactant-like minimum (FSPreMIN) and a product-like minimum (FSPostMIN). Stationary points characteristic of chloride substitution can also decay by losing a hydride ion, leading to $\text{SiH}_2\text{FCl} + \text{H}^-$. Proton abstraction by the fluoride ion is a barrierless process, which can be followed by the formation of hydrogen-bonded complexes, *anti*- or *syn*-FBHMIN, preceding the bihalide formation ($\text{SiH}_2 + \text{FHCl}^-$). The anionic products of the substitution reactions, the chloride and hydride ions, can perform proton abstractions, leading to $\text{SiH}_2\text{F}^- + \text{HCl}$ and $\text{SiHFCl}^- + \text{H}_2$, respectively. Molecular hydrogen can be formed through a more complex mechanism as well, featuring the formation of a dihydrogen-bonded complex (CIDHBMIN) originating from SiH_2F^- and HCl , followed by the decomposition of this complex into three fragments, $\text{SiHF} + \text{H}_2 + \text{Cl}^-$, and ultimately, the recombination of SiHF with the chloride ion. There are two additional minima (*anti*- and *syn*-ClBHMIN) and two transition states (FDITS and ClDITS), the geometries of which can be optimized on the energy surface. However, these four stationary points do not play significant roles in the formation of any of the main products.

3.2 Reaction probabilities

Fig. 3 displays the opacity functions, *i.e.*, the reaction probabilities as a function of impact parameter, for the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions obtained at collision energies between 1 and 40 kcal mol⁻¹.



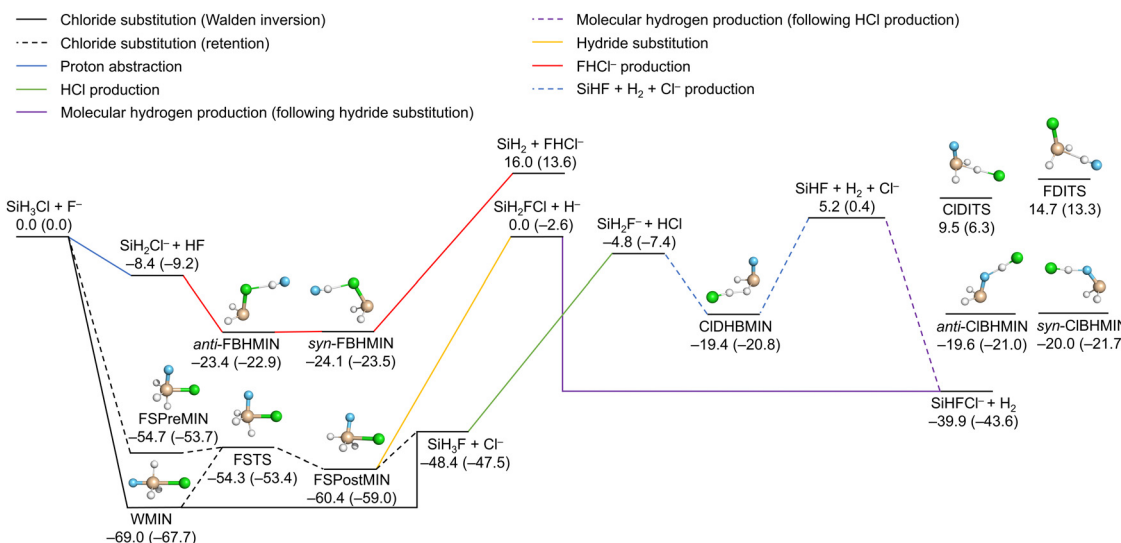


Fig. 2 Schematic potential energy surface of the $\text{SiH}_3\text{Cl} + \text{F}^-$ system featuring the main reaction channels as well as the classical (adiabatic) energies of the minima, transition states and the products relative to the reactants in kcal mol^{-1} .²³

Dividing the chloride substitution into inversion and retention channels, the $b = 0$ reaction probability decreases from 60% to 20% in case of inversion and increases from around 20% to approximately 35% for retention as the collision energy increases. The combined effect leads to a drop in the $b = 0$ reaction probability from 80% to 55%. The maximum reactive impact parameters are also decreasing from $b = 25$ bohr to 8 bohr both for retention and inversion. In the case of proton abstraction, the maximum reactive impact parameters are consistent with those of the chloride substitution across all collision energies. The peaks of these probabilities occur at higher impact parameters, 33% and 26% at $b = 16$ and 6.5 bohr at $E_{\text{coll}} = 1$ and 40 kcal mol^{-1} , respectively. The most pronounced peak is around 45%, observed at $E_{\text{coll}} = 5$ kcal mol^{-1} and $b = 12$ bohr. When considering the hard vibrational energy constraint, which means that both SiH_2Cl^- and HF products must possess a vibrational energy at least equal to their zero-point vibrational energies, reaction probabilities significantly reduce, but the maximum reactive impact parameters remain unaffected. The $b = 0$ reaction probability of proton abstraction with the hard vibrational constraint remains constant 8% at the collision energy range between 10 and 40 kcal mol^{-1} . HCl production displays an almost constant reaction probability across the impact parameter range at $E_{\text{coll}} = 1$ kcal mol^{-1} , slightly below 0.5%. Increasing the collision energy, the maximum reactive impact parameter decreases substantially, while reaction probabilities at smaller impact parameters increase rapidly as the $b = 0$ probability reaches 4.5%. The reaction probability profile for the H_2 production channel is similar to that of the HCl production. At the smallest collision energy this product is observable at high impact parameters, beyond $b = 20$ bohr. As the collision energy increases, the maximum reactive impact parameter drops, and the peak of the reaction probability at low impact parameters rises from 0.2% at the smallest to 2% at the highest collision energy. In the case of

hydride substitution, the maximum reactive impact parameter is roughly $b = 4$ bohr between $E_{\text{coll}} = 25$ and 40 kcal mol^{-1} , and the reaction probability at $b = 0$ grows with increasing collision energy reaching 1%. For the bihalide production channel, between $E_{\text{coll}} = 30$ and 40 kcal mol^{-1} , the maximum reactive impact parameter is between 9.5 and 7.5 bohr. The reaction probability of the bihalide ion production does not exceed 0.2%. Due to the small number of the bihalide production observations its opacity functions are less defined.

As seen above, vibrational energy constraint is only applied for the proton-abstraction channel due to different reasons. In the case of the $\text{S}_{\text{N}}2$ reaction, the zero-point energy violation is negligible partly owing to its high exothermicity. Hydride substitution is also not affected by the vibrational energy constraint. For the other minor channels, the constraint may significantly increase the statistical error because of their low probability.

3.3 Integral cross sections

Integral cross sections of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions are shown by Fig. 4. In the case of chloride substitution, the cross sections decrease with increasing collision energy, for both retention and inversion channels. The combined ICS exceeds 1000 bohr^2 and drops below 70 bohr^2 at the lowest and highest collision energy, respectively. At $E_{\text{coll}} = 1$ kcal mol^{-1} the ICS of the inversion channel is almost 900 bohr^2 , meanwhile the contribution of the retention is only 120 bohr^2 . However, at the highest collision energies their cross sections become comparable. The cross section of proton abstraction also shows a monotonic decrease, from 350 bohr^2 to around 50 bohr^2 in the applied collision energy range, if zero-point energy violating trajectories are also included. By excluding the energy constraint violating trajectories, the cross section is drastically reduced to 50 and 25 bohr^2 at the lowest and highest collision energy, respectively. A maximum appears at around 10 kcal mol^{-1}



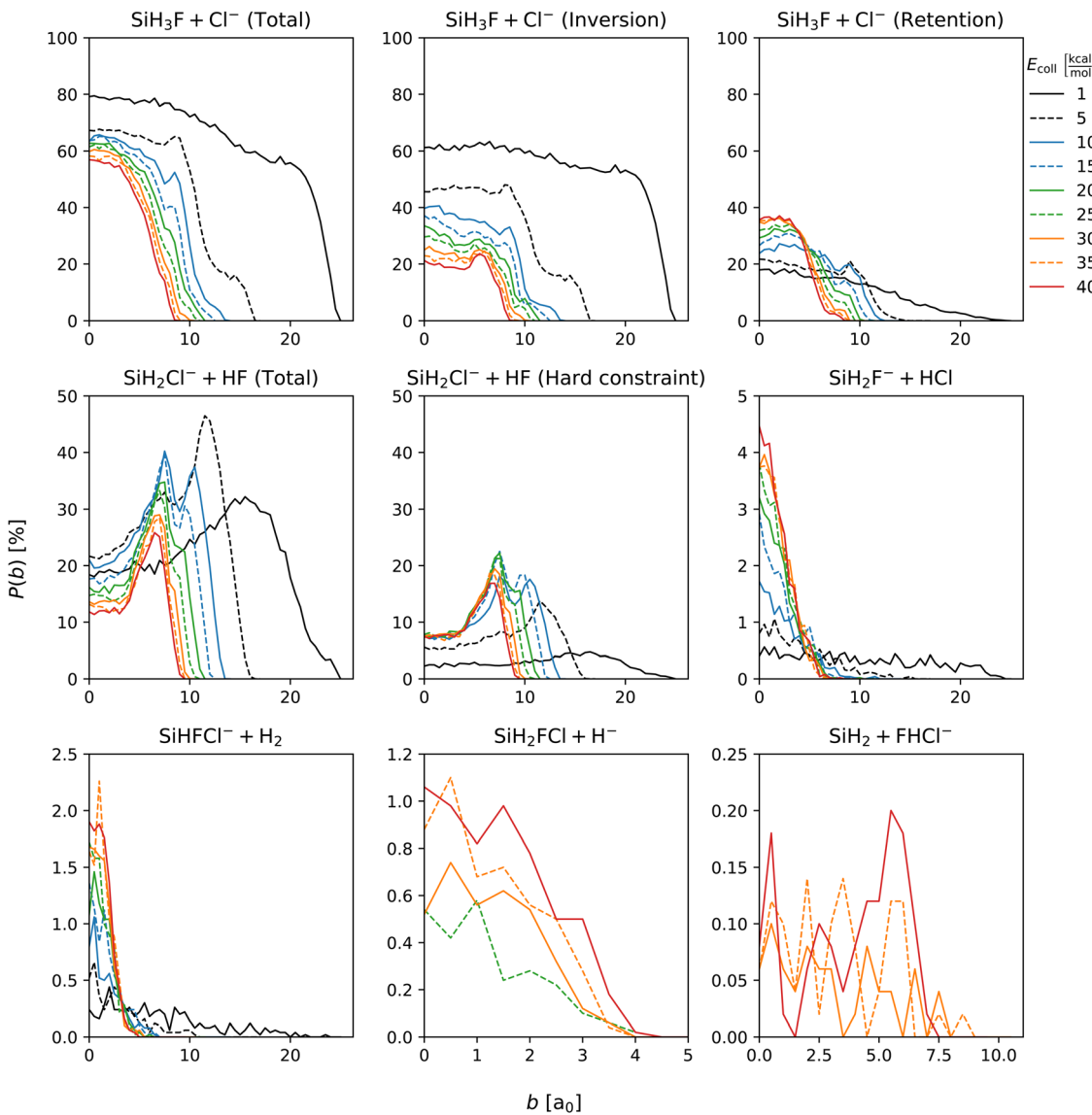


Fig. 3 Reaction probabilities of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions as a function of impact parameter and collision energy.

collision energy, 65 bohr^2 . The integral cross sections of HCl production and H_2 production have similar shapes, maxima at the smallest collision energy, minima at around 10 kcal mol^{-1} and slight increase at higher collision energies. The cross section of the HCl and H_2 productions at the lowest collision energy are 5 and 1.2 bohr^2 , respectively. In case of hydride substitution and the bihalide production channels, their cross sections monotonically increase as the collision energy being increased. However, the cross section of bihalide production is almost negligible at collision energies below 25 kcal mol^{-1} . In case of hydride substitution, a monotonic increase can be observed starting from the smallest collision energy. The cross section of the hydride substitution and bihalide production reaches 0.25 bohr^2 and 0.15 bohr^2 at the highest collision energy, respectively.

Branching ratios as a function of E_{coll} for the different channels and pathways of the title reaction are given in Fig. S1 of the ESI.† It is clearly seen that the $\text{S}_{\text{N}2}$ and proton-

abstraction channels dominate with $\sim 60\%$ and $30\text{--}40\%$, respectively. The fraction of the $\text{S}_{\text{N}2}$ inversion reactivity decreases with E_{coll} , whereas the retention becomes more and more dominant as E_{coll} increases. The branching ratios of the other channels, such as $\text{SiH}_2\text{F}^- + \text{HCl}$, $\text{SiHFCl}^- + \text{H}_2$, $\text{SiH}_2\text{FCl}^- + \text{H}^-$, and $\text{SiH}_2 + \text{FHCl}^-$, are usually small ($< 1\%$) in the E_{coll} range of $1\text{--}40 \text{ kcal mol}^{-1}$ and steeply increase with collision energy.

3.4 Initial attack angle and scattering angle distributions

Fig. 5 shows the initial attack angle distributions. Chloride substitution exhibits a slight preference for back-side attack at low collision energies. However, as the collision energy increases, there is a shift towards a preference for $\cos(\alpha)$ at around -0.5 ($\alpha \approx 120^\circ$). The distribution of α for HCl production has similar shape as that of the chloride substitution channel across all collision energies. In the inversion channel



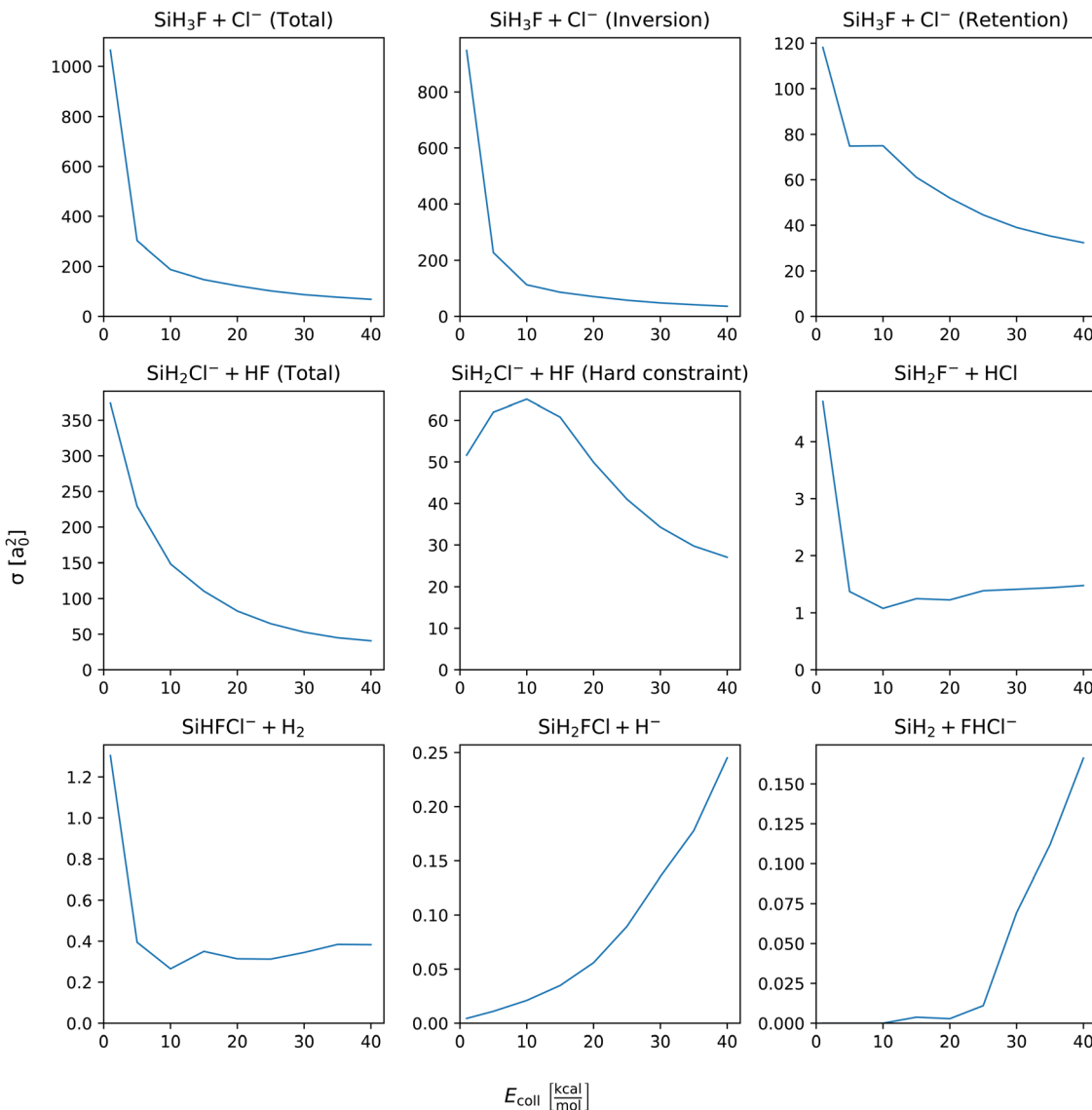


Fig. 4 Integral cross sections of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions as a function of collision energy.

of chloride substitution, there is a pronounced preference for back-side attack at low collision energies, and as the collision energy increases, the distribution becomes isotropic. On the other hand, the retention channel favors the front-side attack at low collision energy, while at high energy, there is a peak in the distribution around $\alpha = 120^\circ$. This significant change in the distribution of the retention channel aligns with our previous observations regarding the collision energy dependence of the mechanism. At the lowest collision energy, the formation of WMIN precedes chloride substitution. However, depending on the impact parameter, energy transfer to the F-Si-Cl bending can occur, leading to the formation of a front-side complex that decomposes into retention products. By examining the $\cos(\alpha)$ distributions, we can conclude that the initial attack angle, in addition to the impact parameter, plays a crucial role in determining the formation of front-side complexes. For proton abstraction, the preference changes from front-side attack at

the lowest collision energy to back-side attack beyond $E_{\text{coll}} = 5 \text{ kcal mol}^{-1}$. The shape of the distributions is not affected by considering the vibrational constraint. The front-side attack preference of proton abstraction at the lowest collision energy can be explained by the extremely competitive nature of WMIN formation at back-side attack which mostly results in chloride substitution. The H_2 production channel exhibits nearly isotropic $\cos(\alpha)$ distribution. The hydride substitution channel shows a preference for back-side attack. The bihalide ion production, which only observed at higher collision energies, also prefers back-side attack. Since this reaction follows the proton abstraction it can be expected that their initial attack angle preference coincides.

Scattering angle distributions are shown at Fig. 6. At low collision energy chloride substitution exhibits backward-forward symmetry. However, as the collision energy increases, forward scattering becomes dominant. Separating the inversion



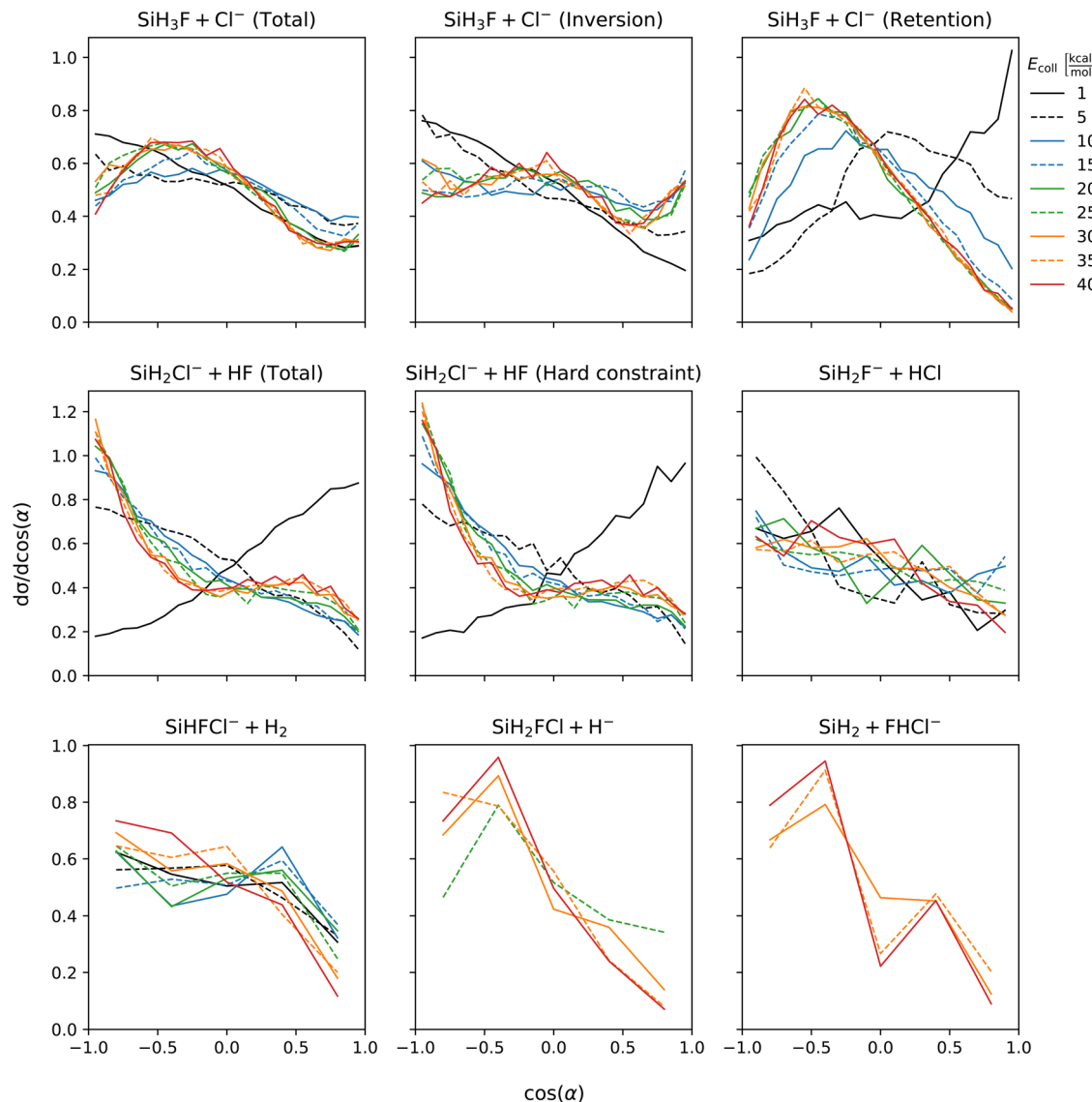


Fig. 5 Initial attack angle distributions of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions obtained at different collision energies.

and retention channels, we find that the preference for forward scattering at high collision energies is primarily attributed to the inversion mechanism. In contrast, the retention channel shows a slight preference for forward scattering at low collision energies, which gradually becomes nearly isotropic at the highest collision energies. Proton abstraction displays symmetric backward-forward scattering at low collision energy. At high collision energies, a preference for forward scattering emerges and the distribution is unaffected by considering vibrational energy constraint. Isotropic scattering angle distributions are observed for both the HCl and H₂ production channels, which is in accordance with the post-reaction-interaction-induced multistep mechanisms, *i.e.*, $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{SiH}_3\text{F} + \text{Cl}^- \rightarrow \text{SiH}_2\text{F}^- + \text{HCl}$ and $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{SiH}_2\text{FCl} + \text{H}^- \rightarrow \text{SiHFCl}^- + \text{H}_2$, respectively, involved in the formation of these products (see Fig. 2). Of course, here the products of the first step are not completely separated and the ion-dipole and/or hydrogen-bond

interactions between them induce the second reaction. Hydride substitution also exhibits an isotropic scattering pattern, possibly due to the ease with which the hydride ion can be deflected due to its extremely small mass. The bihalide production channel, despite its multistep mechanism (see Fig. 2), displays a preferred direction for scattering, which is the backward direction. This contrasts with the forward scattering dominance observed in the proton abstraction reaction at high collision energies. This discrepancy is unexpected since the FHCl^- production follows proton abstraction, and at high collision energies, most of the HF species scatter forward.

3.5 Post-reaction energy distributions

The distribution of both internal and translational energies offers valuable insights into the directness of product formation. For an indirect mechanism, complex formation takes place, leading to the excitation of internal modes of motion:



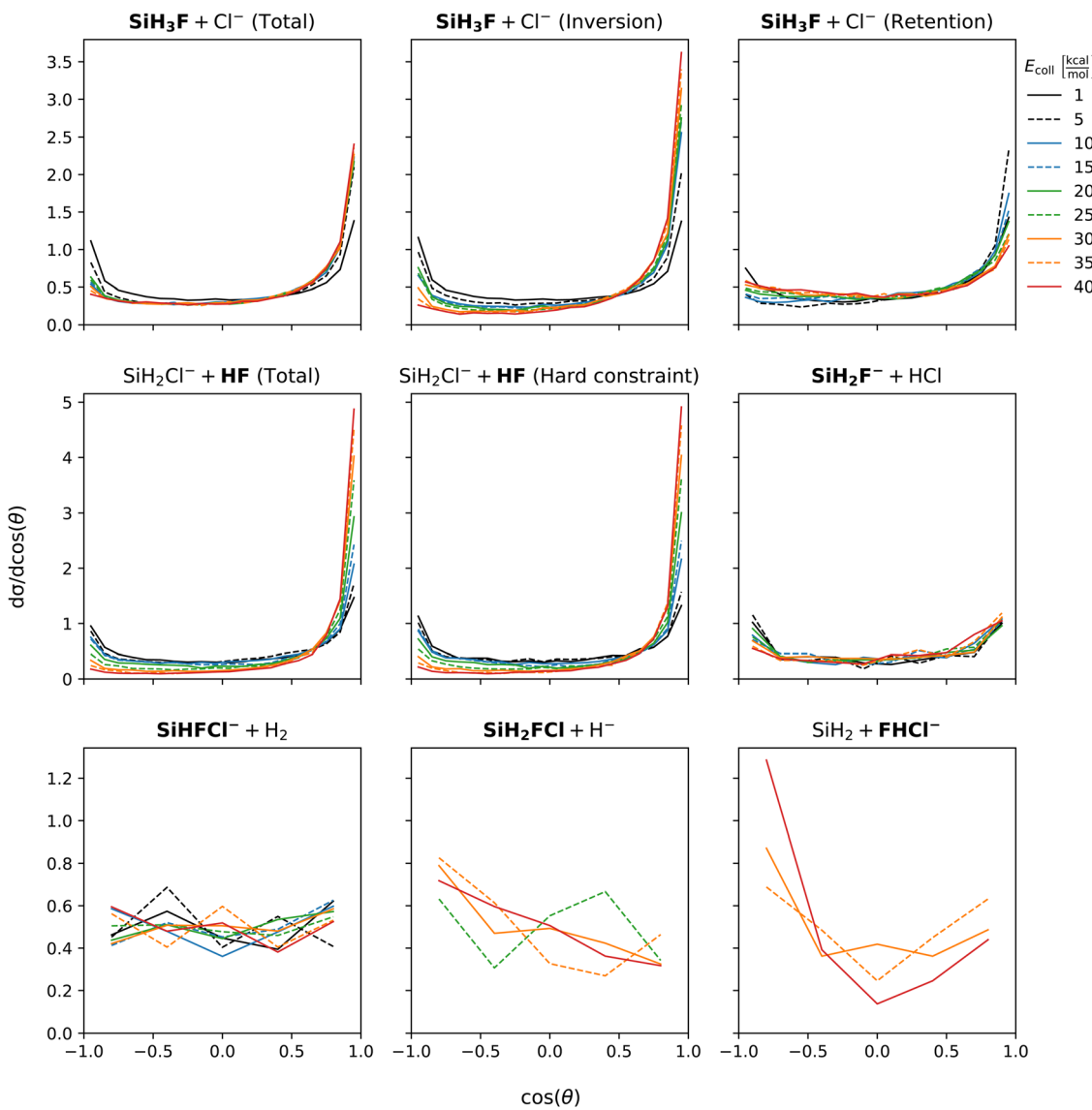


Fig. 6 Scattering angle distributions of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions obtained at different collision energies. Scattering angles are defined as the angle between the center-of-mass velocities of the F^- reactant and the fluorine atom containing product (in bold).

rotation and vibration. In contrast, during a direct reaction, no complexes form and most of the energy from the reactant collision is directly transferred into the translational energy of the resultant products. Throughout our simulations the total energy remains constant. It can be expressed as the sum of the zero-point vibrational energy of SiH_3Cl , the collision energy, and the negative of the reaction energy. The reaction energies are the relative classical energies of the product formations, illustrated in Fig. 2.

Fig. 7–10 show the translational (E_{trans}), internal (E_{int}), vibrational (E_{vib}) and rotational (E_{rot}) energy distributions of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions, respectively. In the ESI,† we provide the distributions of the different energy fractions (E/E_{tot}) as well, where E denotes E_{trans} , E_{int} , E_{vib} , and E_{rot} .

For chloride substitution the total available energy is 66.2 and 105.2 kcal mol⁻¹ at $E_{\text{coll}} = 1$ and 40 kcal mol⁻¹, respectively.

The translational energy distribution of chloride substitution, including both inversion and retention channels, peaks at low energies, specifically below 10 kcal mol⁻¹, across all collision energy ranges. As collision energy increases, this distribution only broadens. Only a minor fraction of the chloride substitution products possesses more than half of the available energy as translation. Conversely, the internal energy distribution, as depicted in Fig. 8, has peaks near the maximum available energy. These observations are consistent with indirect mechanisms characterized by complex formation. According to Fig. 9, the vibrational contribution consistently hovers around 60 kcal mol⁻¹ across all collision energies and the most probable contribution from rotation is less than 20 kcal mol⁻¹ (Fig. 10). At high collision energies the peak of the $E_{\text{vib}}/E_{\text{tot}}$ distribution is around 50%. The $E_{\text{trans}}/E_{\text{tot}}$ and $E_{\text{rot}}/E_{\text{tot}}$ distributions are comparable, meaning that at high collision energy



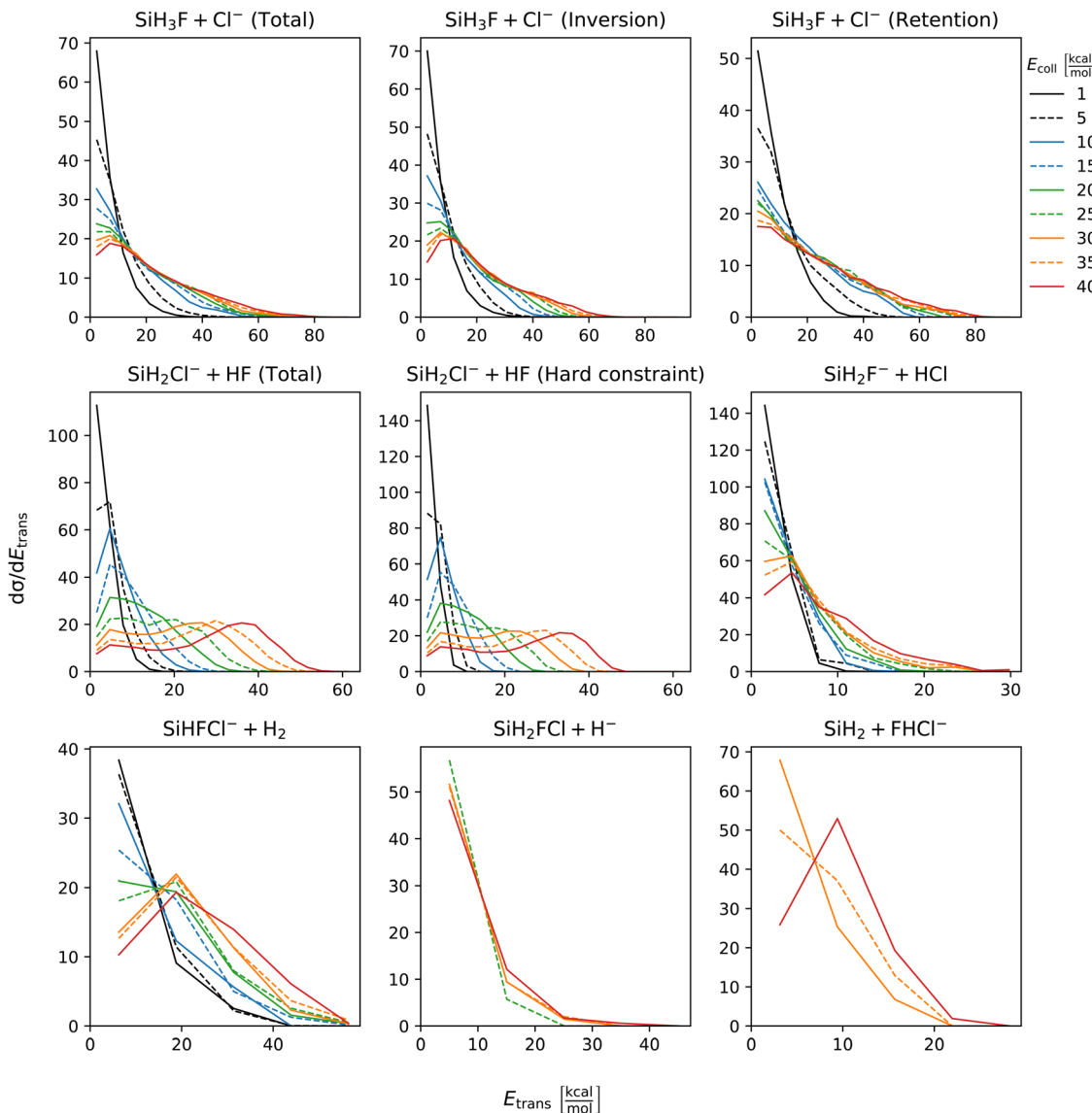


Fig. 7 Translational energy distributions of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions as a function of collision energy.

half of the available energy is approximately evenly shared between rotation and translation (Fig. S2, S6 and S9, ESI[†]).

Analogous to chloride substitution, the peak of the E_{trans} distribution in the hydride substitution channel remains consistent across varying collision energies, peaking at the lowest energy region, implying complex formation. The total available energy for hydride substitution at $E_{\text{coll}} = 40 \text{ kcal mol}^{-1}$ is $56.8 \text{ kcal mol}^{-1}$. Fig. S3 (ESI[†]) shows that almost all hydride substitution products allocate more than half of this energy in internal motion, predominantly as vibration (Fig. S6, ESI[†]).

In the case of proton abstraction, $E_{\text{tot}} = 65.2 \text{ kcal mol}^{-1}$ at the highest collision energy. This reaction shows a compound translational energy distribution consistent with a mixture of direct and indirect mechanisms, as there is a peak near the smallest translational energy across all collision energies, and a second peak which shifts according to the collision energy increase. At low collision energies, most of the total available

energy goes into vibration as the $E_{\text{vib}}/E_{\text{tot}}$ distribution shows a maximum at around 70%. However, at the highest collision energy the peak shifts towards 40%.

The formation of HCl is marked by a low translational energy. The peak of this distribution remains static, even with increasing collision energy, indicating indirect mechanism. During HCl production, the proton abstraction by the leaving ion is more favored when the preceding substitution products move slowly. Interestingly, the partial energy distributions remain consistent during the collision energy increase. The $E_{\text{vib}}/E_{\text{tot}}$ and $E_{\text{rot}}/E_{\text{tot}}$ distributions peak at 75% and around 15–20%, respectively. A significant portion of the vibrational energy is localized at SiH_2F^- , as shown in Fig. S7 (ESI[†]).

The peak of the translational energy distribution of the H_2 production shifts with increasing collision energies, approximately with half the increase. As depicted in Fig. S7 (ESI[†]), at lower collision energies, most of the available energy is

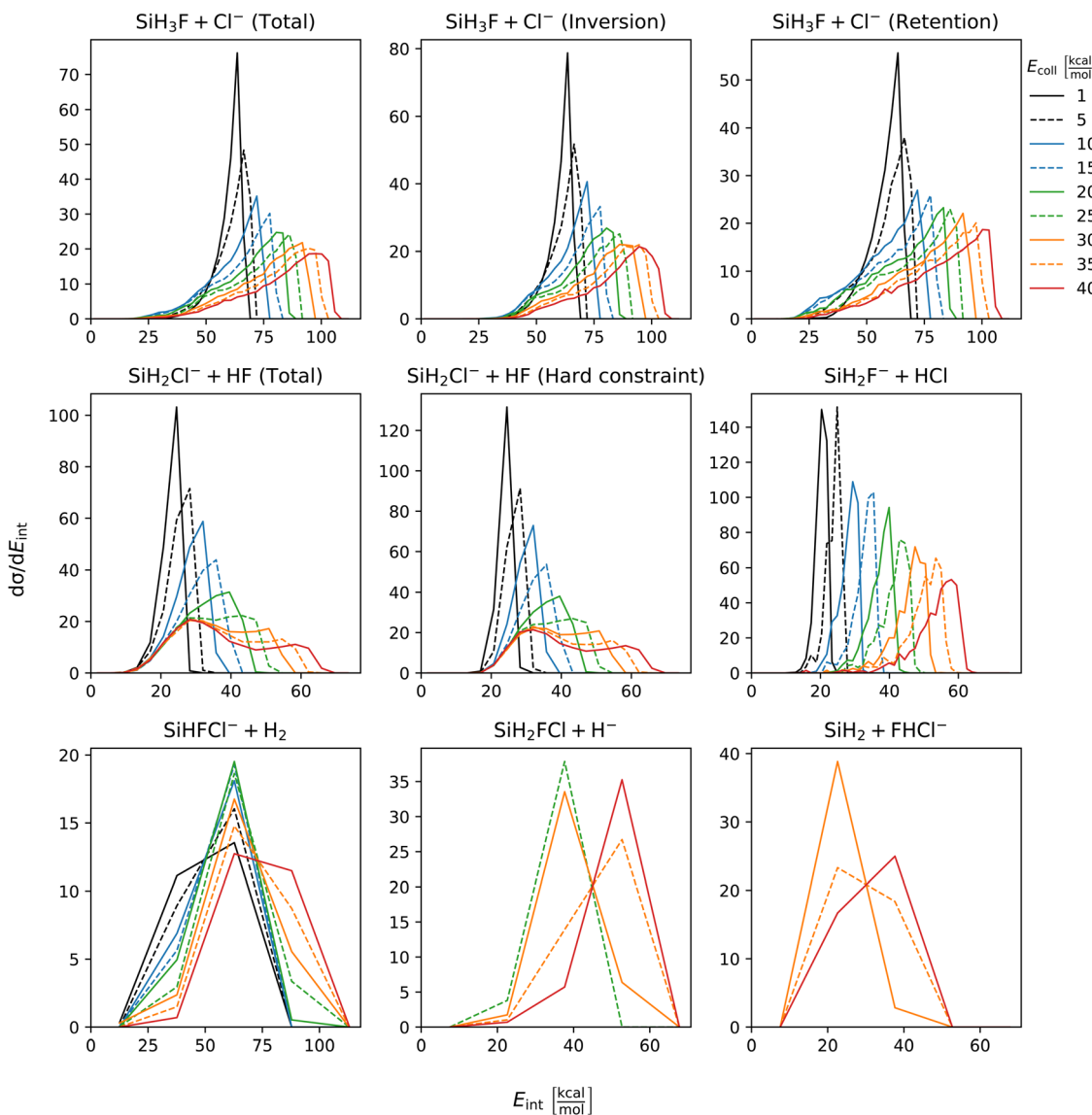


Fig. 8 Internal energy distributions of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions as a function of collision energy.

channeled into the vibration of SiHFCl^- . However, at higher collision energies, this proportion reduces to around half.

In case of bihalide production, the maximum of the translational energy distribution shifts by the collision energy increase. According to the $E_{\text{trans}}/E_{\text{tot}}$, $E_{\text{rot}}/E_{\text{tot}}$ and $E_{\text{vib}}/E_{\text{tot}}$ distributions, about half of the available energy goes into vibration and the remaining half is shared equally between rotation and translation. About two thirds of the vibrational energy goes into the FHCl^- ion.

In order to obtain additional insight into the directness of the title reaction, we have plotted the integration time distributions of the different channels in Fig. S12 of the ESI.† As E_{coll} increases all the distributions shift toward shorter times, however, at the highest E_{coll} range of 30–40 kcal mol⁻¹ the collision energy dependence is not significant. The minor channels, especially $\text{SiH}_2\text{F}^- + \text{HCl}$, $\text{SiHFCl}^- + \text{H}_2$, and $\text{SiH}_2 + \text{FHCl}^-$, are the most indirect, *i.e.*, their distributions peak at the longest

time values at a given E_{coll} , but the indirect character (time scale) of the $\text{S}_{\text{N}}2$ and proton-abstraction channels is also substantial.

3.6 The effect of replacing the central atom

In a theoretical study of gas-phase ion–molecule reactions, it is worth comparing analogous systems to place our observations in a larger context. Our title system, $\text{SiH}_3\text{Cl} + \text{F}^-$, can be compared to the carbon- and the phosphorous-analogues: $\text{CH}_3\text{Cl} + \text{F}^-$ and $\text{PH}_2\text{Cl} + \text{F}^-$, respectively. These comparisons are justified by the fact that carbon and silicon are in the same main group, while phosphorus and silicon are in neighboring groups within the same period. The $\text{CH}_3\text{Cl} + \text{F}^-$ system has been widely studied,^{1,3,6–8,25} meanwhile for $\text{PH}_2\text{Cl} + \text{F}^-$, our research group recently published a QCT study.²⁶

In these three systems, the most important reactions are the $\text{S}_{\text{N}}2$ chloride ion substitutions and the proton abstractions.



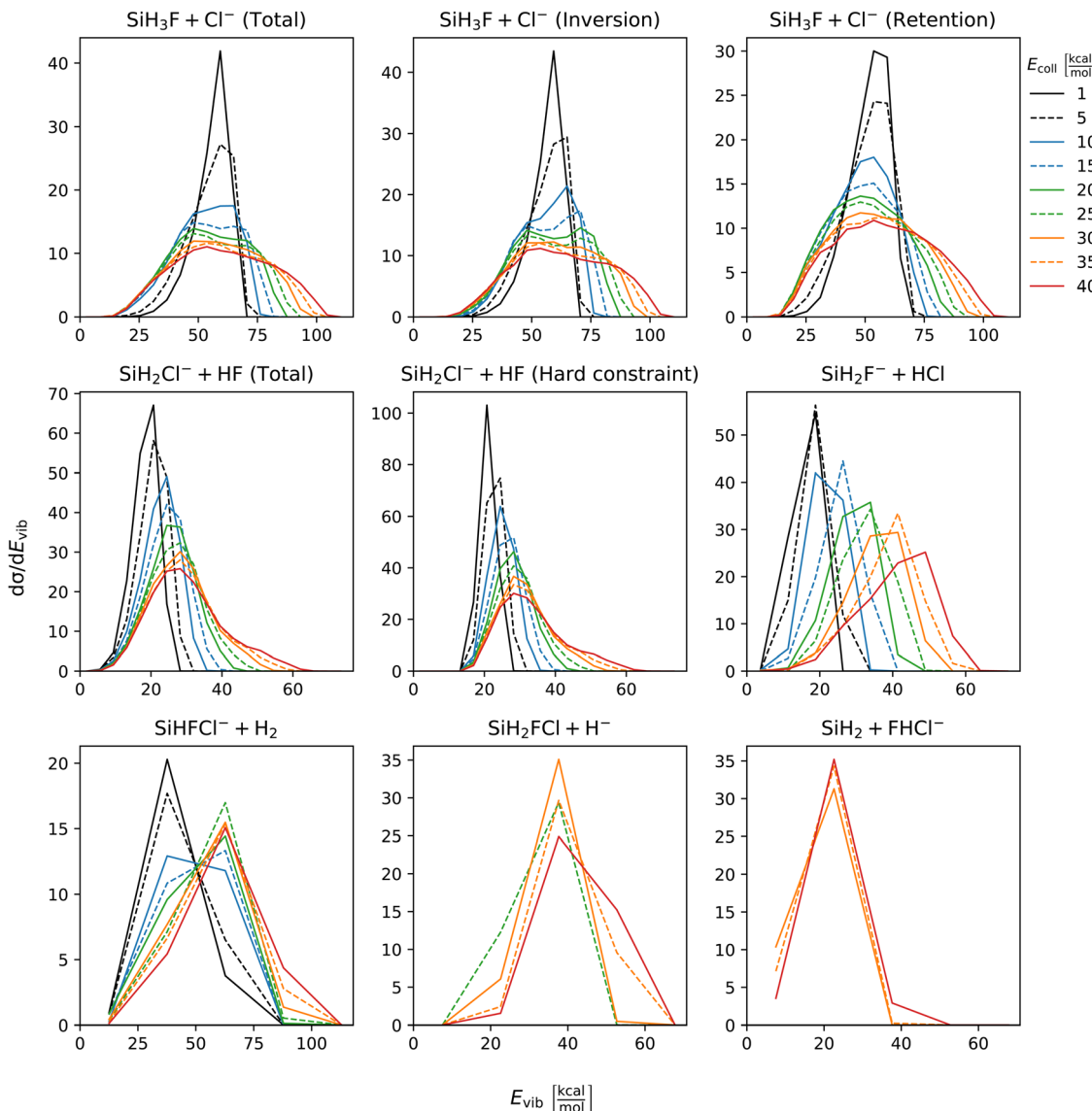


Fig. 9 Vibrational energy distributions of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions as a function of collision energy.

The analogous chloride substitutions are all exothermic reactions.^{25,26} However, from a stereochemical perspective, we observe significant differences in the substitution reactions, largely attributable to the relative energies of the stationary points.

$\text{S}_{\text{N}}2$ by Walden inversion is favorable in all systems, as evidenced by the submerged stationary points. A notable difference between the carbon analogous inversion and the others is that, in case of carbon, the potential shape is a double well, whereas for $\text{SiH}_3\text{Cl} + \text{F}^-$ and $\text{PH}_2\text{Cl} + \text{F}^-$, single well potentials can be found. Front-side attack resulting in retention $\text{S}_{\text{N}}2$ products is characteristic of both $\text{SiH}_3\text{Cl} + \text{F}^-$ and $\text{PH}_2\text{Cl} + \text{F}^-$, as the stationary points of the corresponding reactions all possess negative energies. The result is a stereochemical mixture of inversion and retention $\text{S}_{\text{N}}2$ products at all collision energies. Meanwhile, for $\text{CH}_3\text{Cl} + \text{F}^-$, the front-side attack transition state leading to retention has high positive energy, limiting retention product formation at low collision energy.

However, retention $\text{S}_{\text{N}}2$ becomes possible at collision energies lower than the relative energy of the corresponding front-side attack transition state, thanks to a recently-discovered compound mechanism known as double inversion.²⁵ In the double-inversion chloride substitution of $\text{CH}_3\text{Cl} + \text{F}^-$, the attacking nucleophile fails to detach the proton from the methyl chloride molecule. Instead, it creates a transition state with an inverted configuration by moving the proton to the opposite side of the carbon atom. This transition state then rearranges into an inverted Walden transition state, which decomposes *via* a second inversion into the retention $\text{S}_{\text{N}}2$ products. The discovery of the double-inversion mechanism was significant in explaining the unexpectedly high probability of $\text{S}_{\text{N}}2@C$ retention at low collision energy.²⁵ For systems with phosphorus and silicon central atoms, investigating both the possibility and significance of double inversion is worthwhile.

In the $\text{SiH}_3\text{Cl} + \text{F}^-$ system, chloride substitution with retention makes a significant contribution even at the lowest

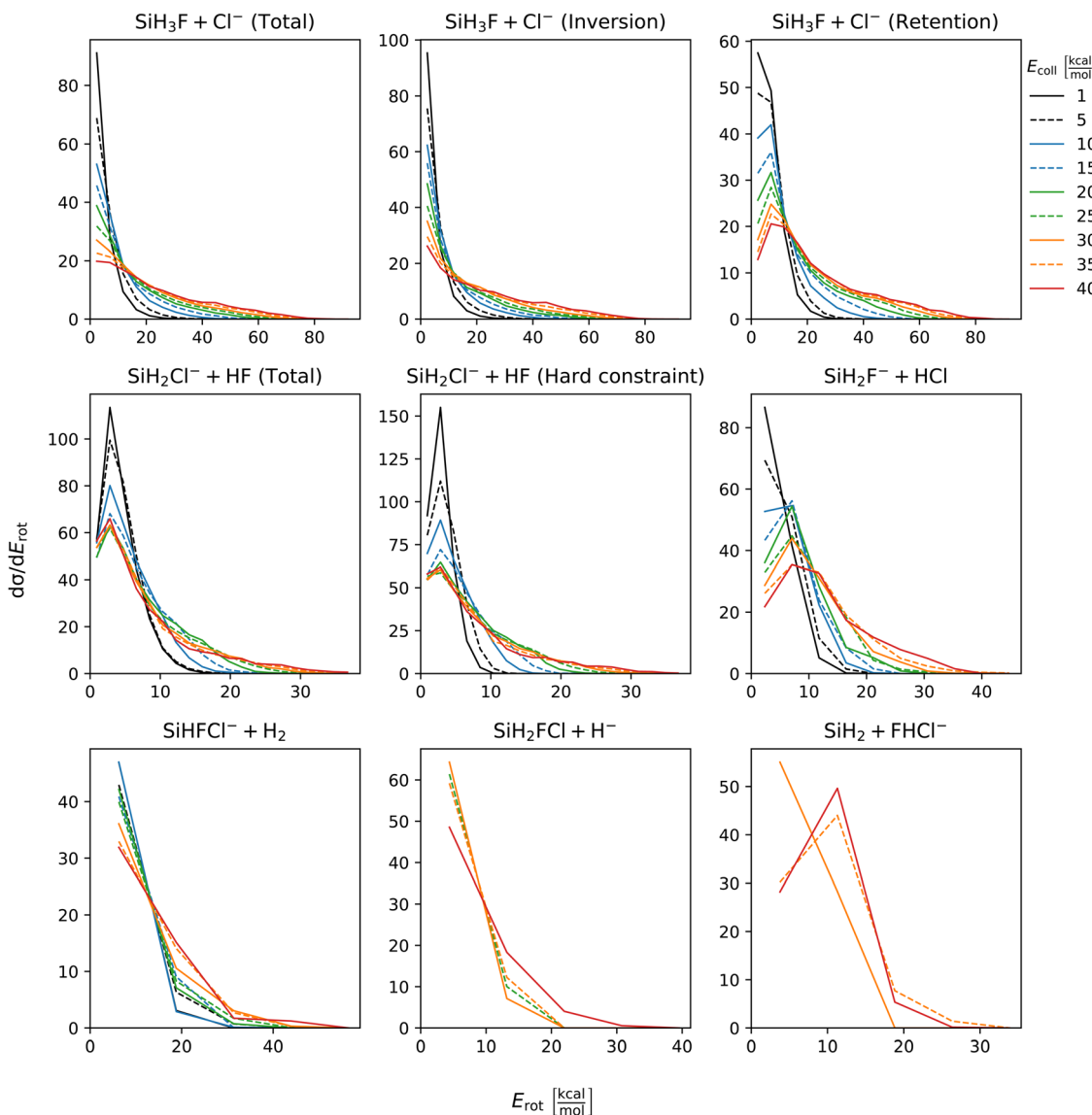


Fig. 10 Rotational energy distributions of the $\text{SiH}_3\text{Cl} + \text{F}^- \rightarrow \text{A} + \text{B}$ reactions as a function of collision energy.

collision energies, as predicted by the schematic PES and evidenced by QCT simulations. Therefore, the retention $\text{S}_{\text{N}}2@\text{Si}$ can be explained by the simple, straightforward front-side attack mechanism. On the other hand, it is of great theoretical interest to explore if the analogue of double inversion is even possible in silicon-centered systems, as it elucidates the atomistic dynamics of this mechanism type.

After examining several $\text{S}_{\text{N}}2$ trajectory animations in the $\text{SiH}_3\text{Cl} + \text{F}^-$ system, in which at least the Si–H–F bond is present or proton abstraction is one of the partial steps of the entire mechanism, we have observed that double inversion described by Szabó and Czakó²⁵ is atypical here because of the following reasons. (1) Unlike at the analogous C-centered reaction, in the present case the attacking nucleophile easily detaches the proton due to the exothermicity of the HF formation channel. (2) Substitution *via* retro-proton-abstraction leads to a stereochemical mixture because the formation of both inversion and

retention $\text{S}_{\text{N}}2$ complexes are allowed at low energies. It appears that in double inversion, not only the existence of the DI transition state and its energy being lower than that of the front-side attack transition state are relevant. It is favorable for the mechanism if the reaction energy of proton abstraction is also higher than the relative energy of DITS; otherwise, proton abstraction occurs instead of the partial abstraction and stereochemical rearrangement.

It is also noteworthy that vibrationally highly-excited SiH_3F can invert due to its flexibility. In carbon-centered systems, post-reaction $\text{S}_{\text{N}}2$ product inversion is usually not observed, though in the case of the $\text{NH}_2^- + \text{CH}_3\text{I}$ reaction retention can occur *via* the formation of a $[\text{CH}_3 \cdots \text{NH}_2]$ complex in which a rotation of the CH_3 group may cause a second inversion.²⁷

In the analogous proton abstraction reactions, the process is endothermic in the $\text{CH}_3\text{Cl} + \text{F}^-$ system, accessible only at high collision energies. Conversely, proton abstraction in both



$\text{SiH}_3\text{Cl} + \text{F}^-$ and $\text{PH}_2\text{Cl} + \text{F}^-$ is exothermic and a high-probability reaction at all collision energies.

In addition to chloride substitution and proton abstraction in $\text{PH}_2\text{Cl} + \text{F}^-$, various low-probability channels analogous to reactions in the silicon-centered system are observed.²⁶ $\text{PHF}^- + \text{HCl}$ is the third most probable reaction, similar to how HCl production is the third most dominant in $\text{SiH}_3\text{Cl} + \text{F}^-$. Molecular hydrogen, bihalide ion, as well as hydride ion production reactions are observed with low probability as well, which are also characteristic of the silicon-containing system. None of these product formations are significant in $\text{CH}_3\text{Cl} + \text{F}^-$.

4. Summary and conclusions

We have computed the first differential cross sections such as various angular and product energy distributions, for a Si-centered $\text{S}_{\text{N}2}$ reaction, $\text{F}^- + \text{SiH}_3\text{Cl}$. The QCT simulations have utilized our recently-developed full-dimensional analytical *ab initio* PES,²³ which allows uncovering even the low-probability channels such as $\text{SiH}_2\text{F}^- + \text{HCl}$, $\text{SiH}_2\text{FCl}^- + \text{H}^-$, $\text{SiH}_2^- + \text{FHCl}^-$, $\text{SiHFCl}^- + \text{H}_2$, $\text{SiHF}^- + \text{H}_2 + \text{Cl}^-$, and $\text{SiH}_2^- + \text{HF} + \text{Cl}^-$, besides the major $\text{S}_{\text{N}2}$ and proton-transfer channels leading to $\text{SiH}_3\text{F} + \text{Cl}^-$ and $\text{SiH}_2\text{Cl}^- + \text{HF}$, respectively. The following conclusions can be summarized based on the differential cross sections obtained for the different two-fragment product channels:

(a) The cold translational and hot internal energy distributions indicate that the $\text{S}_{\text{N}2}$ reaction, $\text{SiH}_3\text{F} + \text{Cl}^-$ channel, is mainly indirect, however, the angular distributions show spatial preferences depending on collision energy as well as inversion/retention.

(b) The angular distributions of the proton-abstraction channel, $\text{SiH}_2\text{Cl}^- + \text{HF}$, show front-side attack backward-forward scattering preference at low E_{coll} , which shifts toward back-side attack and forward scattering as E_{coll} increases. The energy distributions indicate a mixed direct-indirect proton-transfer mechanism.

(c) The attack angle distributions of the $\text{SiH}_2\text{F}^- + \text{HCl}$ channel are similar to those of the $\text{S}_{\text{N}2}$ reaction. The $\text{SiH}_2\text{F}^- + \text{HCl}$ products are formed with isotropic scattering and translationally cold due to the indirect mechanism of this channel.

(d) The $\text{SiH}_2\text{FCl}^- + \text{H}^-$ channel prefers back-side attack and scatters isotropically with low translational energy indicating an indirect pathway.

(e) Even if $\text{SiH}_2^- + \text{FHCl}^-$ has a multistep mechanism, its angular distributions show back-side attack and backward scattering preference.

(f) The angular distributions of the $\text{SiHFCl}^- + \text{H}_2$ channel are mostly isotropic and the product translational energy is cold, in consistent with its indirect mechanism.

Finally, we note that the above-described two-fragment channels produce different ionic products, thus, their selective experimental detection may be straightforward. We hope that the present work motivates such experimental investigations as well as further theoretical studies.

Conflicts of interest

There are no conflicts of interest to declare.

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