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Uncovering an oxide ion substitution for the $OH^- + CH_3F$ reaction

A full-dimensional high-level *ab initio* analytical potential energy surface is developed for the $OH^- + CH_3F$ system allowing efficient dynamics simulations, which provide unprecedented insights into the mechanisms of the title reaction and reveal a novel oxide ion substitution pathway leading to the unexpected $HF + CH_3O^-$ products.

Image credit: Viktor Tajti, Alexandra N. Kovács, Domonkos A. Tasi and Dóra Papp





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Uncovering an oxide ion substitution for the OH⁻ + CH₃F reaction†

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Theoretical investigations on chemical reactions allow us to understand the dynamics of the possible pathways and identify new unexpected routes. Here, we develop a global analytical potential energy surface (PES) for the $OH^- + CH_3F$ reaction in order to perform high-level dynamics simulations. Besides bimolecular nucleophilic substitution (S_N2) and proton abstraction, our quasi-classical trajectory computations reveal a novel oxide ion substitution leading to the $HF + CH_3O^-$ products. This exothermic reaction pathway occurs via the $CH_3OH\cdots F^-$ deep potential well of the S_N2 product channel as a result of a proton abstraction from the hydroxyl group by the fluoride ion. The present detailed dynamics study of the $OH^- + CH_3F$ reaction focusing on the surprising oxide ion substitution demonstrates how incomplete our knowledge is of fundamental chemical reactions.

Introduction

In chemistry, one of the most important reactions is the bimolecular nucleophilic substitution (S_N2). The traditional picture of the S_N2 reactions was described by Ingold and coworkers more than 80 years ago,1,2 and since then these reactions have always been in the focus of scientific interest.3-16 In a schematic $X^- + CH_3Y \rightarrow CH_3X + Y^- S_N2$ reaction, the widelyknown Walden inversion usually occurs as follows: in the entrance channel the reactants form an ion-dipole X⁻···CH₃Y complex, and then a [X···CH₃···Y]⁻ transition state can be found, which finally leads to a XCH3···Y complex in the product channel. The above presented mechanism inverts the configuration of the CH₃Y reactant, due to the umbrella motion of the CH₃ group. Besides inversion, at higher collision energies (E_{coll}) , retention can also be eventuated, and thus the configuration of the CH₃X product remains the same as that of the CH₃Y reactant. There are two mechanisms that result in retention: front-side attack and the recently discovered doubleinversion pathway.17 In the case of the front-side attack, the reaction pathway goes through a high-energy [XYCH₃]

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 \dagger Electronic supplementary information (ESI) available: Detailed computational methods, relative translational energy and internal energy distributions of the products, ratio (%) of the integral cross sections of each pathway [S_N2 (via PostHMIN1, retention and inversion), proton abstraction, oxide anion substitution (with ZPE-constraints) and proton exchange], lifetime of PostHMIN1 in the deep well of the oxide anion substitution at several collision energies and impact parameters, benchmark Cartesian coordinates (Å) and energies (E_h) of the stationary points, and representative trajectories of each reaction pathway. See DOI: 10.1039/d1sc03834f

transition state. The double-inversion mechanism is defined by a proton-abstraction induced inversion followed by a traditional Walden inversion. However, if one considers the reaction conditions and differentiate several direct (rebound and stripping) and indirect (ion-dipole, hydrogen-bond and frontside complex formations, roundabout, barrier recrossing) pathways, the description of the S_N2 reactions becomes more complex. Regarding our previous studies, reactions becomes more complex. Regarding our previous studies, the back-side attack Waldeninversion. It should also be noted that competing with S_N2 , in most cases, proton abstraction can occur leading to the HX + CH_2Y^- products.

S_N2 reactions are mostly studied between halide ions and methyl halides, however, over the past 30 years the variety of the analysed reactions has widened. On one hand, halide ions can be replaced with several other nucleophiles, such as OH⁻, SH⁻, CN-, NH2-, PH2-, etc., 19-25 on the other hand, methyl halides can be substituted with relevant alkyl halides.26-30 For the OH - + CH_3Y [Y = F, Cl, Br, I] gas-phase S_N^2 reactions, previous studies showed that in the product channel, instead of the traditional HOCH3···Y ion-dipole complex, a hydrogenbonded CH₃OH···Y⁻ global minimum can be found.^{21,22,31} In 2002, Hase and co-workers investigated the impact of the $CH_3OH\cdots F^-$ deep well on the $OH^- + CH_3FS_N 2$ reaction by using direct dynamics simulations.6 64 trajectories were initiated from the top of the [HO···CH₃···F]⁻ barrier towards the CH₃OH + F⁻ products, and each trajectory was propagated until the distance of the products reached 17 Å or the lifetime of the trajectory exceeded 3 ps. Among these trajectories, 33 reversed to the entrance channel by forming the OH-...CH3F complex and 27 followed a direct reaction path, where the leaving F avoids the deep minimum along the O-C···F⁻ axis. Only 4 trajectories went through an indirect reaction path and were

trapped in the region of the $CH_3OH\cdots F^-$ well, of which only one trajectory formed the $CH_3OH + F^-$ products in the 3 ps propagation time. Concerning the role of the deep potential well in the dynamics of the title reaction, similar findings were revealed in the studies of Tsutsumi *et al.*^{32,33} and Hare *et al.*³⁴

In the present work, nearly 20 years after the studies of Hase and co-workers,6 we investigate the OH + CH3F reaction performing high-level dynamics simulations. Our main goal is to take into account the effect of the deep post-reaction well on the title S_N2 reaction by computing nearly 1 million full-length quasi-classical trajectories (QCT) and considering a wide range of E_{coll} . To perform these QCT computations, relying on our previous study,35 we construct a global full-dimensional ab initio potential energy surface (PES) for the OH- + CH₃F reaction utilizing the Robosurfer program package recently developed in our group.36 Our dynamics simulations reveal a novel reaction route, observed at lower $E_{\rm coll}$, where a fluoride ion is substituted by an oxide ion leading to the HF + CH₃O⁻ products. How does this oxide ion substitution proceed and how competitive is it with the traditional reaction paths? In the following we answer these questions by providing a detailed dynamical characterization of the title reaction.

Results and discussion

Description of the S_N2 potential energy surface

In order to describe the possible reaction pathways of any given reaction, first a benchmark characterization of the stationary points on its PES (minima and transition states) must be performed. The schematic PES of the $OH^- + CH_3F$ S_N2 reaction is shown in Fig. 1. A detailed description of the applied computations for the stationary-point characterization can be found

in the ESI.† While proton abstraction is endothermic $(21.62 \text{ kcal mol}^{-1})$, $S_N 2$ turns out to be an exothermic reaction with an energy of -19.97 kcal mol⁻¹ relative to the reactants (Fig. 1). As expected, in the entrance channel of the back-side attack, a traditional ion-dipole complex (PreMIN) is situated with a central transition state (WaldenTS). The global minimum of the OH⁻ + CH₃F reaction is in the product channel, where two different types of H-bonded complexes are obtained (PostHMIN1 and PostHMIN2). Note that, as a result of the CH₃ group rotation, PostHMIN1 and PostHMIN1' are conformational isomers, and PostHMIN1 is below the latter by 0.22 kcal mol⁻¹. The back-side attack goes through a submerged WaldenTS with a relative energy of -2.60 kcal mol⁻¹, whilst the transition states of the front-side attack have barrier heights of 42.49 (FSTS1) and 34.22 (FSTS2) kcal mol⁻¹. Concerning the double-inversion mechanism, DITS is found to be energetically lower than FSTS2 by \sim 17 kcal mol⁻¹. Besides S_N 2, proton abstraction is investigated as well, and the resulting two minima are presented together with all benchmark Cartesian coordinates and energies of the stationary points in the ESI.† Based on the defined stationary points, our next step is to build a global, analytical PES for the OH⁻ + CH₃F reaction using the Robosurfer program system.³⁶ The computational details of the PES development are summarised in the ESI.† To underline the accuracy of the PES, the obtained relative energies of the stationary points must be compared with the benchmark values. As Fig. 1 shows, in most cases the PES reproduces the all-electron CCSDT(Q)/completebasis-set-quality benchmark energies within chemical accuracy (1 kcal mol⁻¹), with a slightly higher difference of \sim 1.6 kcal mol⁻¹ for the H₂O + CH₂F⁻ products.

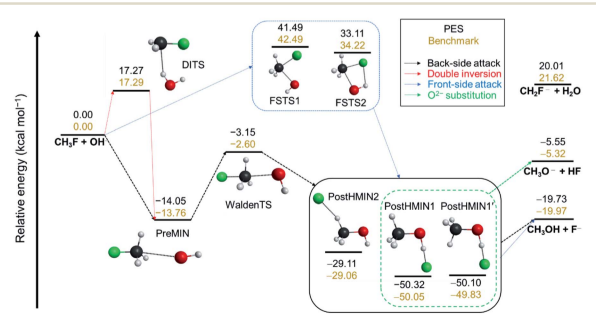


Fig. 1 The schematic potential energy surface of the $OH^- + CH_3F S_N 2$ reaction featuring oxide ion substitution. The corresponding arrows unveil the possible reaction pathways for $S_N 2$ and oxide anion substitution along the stationary points with the accurate all-electron CCSDT(Q)/complete-basis-set-quality benchmark classical energies and PES values (obtained by geometry optimizations on the analytical PES), relative to $OH^-(eq.) + CH_3F(eq.)$. The energy levels of the three product channels ($S_N 2$, proton abstraction and oxide anion substitution) are also presented.

Reaction dynamics simulations: the novel oxide ion substitution

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Building upon an accurate global analytical PES, high-level dynamics simulations can be performed to study the chemical reaction of interest at an atomic level. For the OH⁻ + CH₃F reaction QCT computations are carried out at several E_{coll} . Details of the QCT simulations are provided in the ESI.† Besides the usual products of S_N2 and proton abstraction, the simulation trajectories resulted in the HF and CH₃O⁻ compounds in certain cases, revealing the existence of a new reaction path. We further call this pathway as an oxide ion substitution indicating that the F⁻ of the CH₃F reactant is exchanged by an O^{2-} . As seen in Fig. 1, for the title reaction, oxide ion substitution happens to be exothermic and the reaction energy obtained on the PES (-5.55 kcal mol⁻¹) is in excellent agreement with the benchmark value $(-5.32 \text{ kcal mol}^{-1})$. The snapshots of a representative trajectory are shown in Fig. 2 depicting the oxide ion substitution at a E_{coll} of 10 kcal mol⁻¹. Using time step intervals of 0.0726 fs, dynamics simulations allow us to follow the motion of the atoms step by step. Within the entrance channel of the reaction, the two reactants approach each other for ~ 0.35 ps. Afterwards, within a timeframe of only ~ 0.6 ps, Walden inversion occurs via the above-described pathway (PreMIN → WaldenTS \rightarrow PostHMIN), and the system gets trapped in the postreaction deep well complex for ~2.4 ps. Subsequently, the F removes the proton from the hydroxyl group leading to generation of the products, HF + CH_3O^- in ~ 0.07 ps. The average

lifetime of the PostHMIN1 complex, 1–5 ps depending on $E_{\rm coll}$, is given in Table S1.† Thus, the typical trajectory of oxide ion substitution spends long time in the PostHMIN1 region, however, PreMIN complex formation and fast proton abstraction in the product well can also be found. To confirm the certainty of this novel oxide ion substitution, *ab initio* computations are performed along representative trajectories. As also seen in Fig. 2, the direct *ab initio* energies and the PES values are in good agreement indicating that the given path does exist for the title reaction. In the ESI,† the interested reader can find an actual QCT simulation of the oxide ion substitution in motion along with the other ($S_{\rm N}2$, proton abstraction, proton exchange, *etc.*) pathways.

Following the approach of Hase and co-workers,⁶ our next goal is to identify the S_N2 reactions taking place via the deep PostHMIN1 and to define the probability relation of these pathways with regard to the oxide ion substitution. For this purpose, we follow the relevant atomic distances from the end of the trajectories (see the ESI† for details). The integral cross sections (ICSs) and their ratios as a function of the E_{coll} for the examined reaction routes are shown in Fig. 3 and Table S2,† respectively. Cross sections, which are proportional with the measureable reactivity at a given E_{coll} , are obtained by integrating initial orientation- and vibrational-phase-averaged b-weighted reaction probabilities over b impact parameters. Provided that the back-side attack substitution is a submerged exothermic path, S_N2 has the largest ICS at each E_{coll} with a maximum of ~ 5.2 bohr² at $E_{coll} = 10$ kcal mol⁻¹. The ICS of

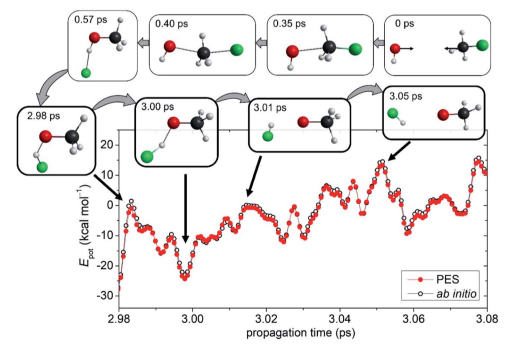


Fig. 2 The novel reaction pathway: snapshots of a dynamics simulation representing the oxide ion substitution for the $OH^- + CH_3F$ reaction at a collision energy of 10 kcal mol^{-1} (b=0). In the course of the exceptional proton abstraction by the fluoride ion, the direct *ab initio* potential energies are compared with the fitted PES values, relative to OH^- (eq.) + CH_3F (eq.), as a function of time. The *ab initio* energies are obtained at the CCSD-F12b/aug-cc-pVTZ + BCCD(T)/aug-cc-pVDZ - BCCD/aug-cc-pVDZ composite level of theory and the PES values correspond to the present analytical fitted composite PES. The time evolution of the potential energies for the complete reaction pathway from reactants to products is shown in Fig. S4.†

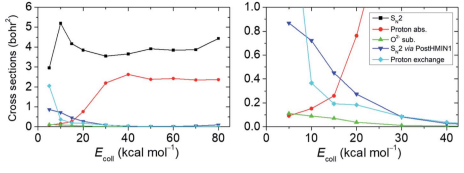


Fig. 3 Integral cross sections of the studied pathways at several collision energies. For the OH $^-$ + CH $_3$ F reaction, cross sections are determined in the range of 5–80 kcal mol $^{-1}$ collision energies for the reaction pathways of S $_N$ 2, proton abstraction, oxide ion substitution, S $_N$ 2 occurring *via* the deep PostHMIN1 and proton exchange.

the proton abstraction suffers a significant increase starting from E_{coll} of 20 kcal mol⁻¹ and plateaus around 2.2–2.6 bohr² at higher energies. (It is important to note that proton abstraction still occurs at E_{coll} below the reaction endothermicity due to the zero-point energy (ZPE) violation of the products, which has not been considered for Fig. 3.) The highest possibility of S_N2 occurring *via* PostHMIN1 is at $E_{\text{coll}} = 5 \text{ kcal mol}^{-1}$ with an ICS of ~ 0.9 bohr². Dynamics simulations also show PreMIN complex formation, but the trajectories usually spend more time in the post-reaction well. As can be expected for a barrierless (WaldenTS is below the reactants) exothermic reaction, the direct path of S_N2 becomes more and more dominant with the gradual increase of E_{coll} avoiding the H-bonded global minimum in the product channel. The finding that the trajectories avoid the Hbonded minimum at high E_{coll} can be explained by the fact that the PostHMIN1 complex is not along the O-C-F reaction coordinate and at higher translational energies F- directly departs along the nearly collinear O-C-F arrangement without roaming around the CH₃OH fragment to form an OH-bonded complex. The above tendency is also reflected by the internal

and the translational energy distributions of the S_N2 and the proton-abstraction products (see Fig. S1 and S2 in the ESI†). Upon increasing E_{coll} , translational energies of the products are clearly higher in contrast to the less impacted internal energies. This indicates that the direct reaction path is favoured in both, S_N2 and proton abstraction as well. Analogously to the S_N2 via PostHMIN1, the ICS of the oxide ion substitution also peaks at an E_{coll} of 5 kcal mol⁻¹ with a maximum value of ~0.1 bohr², and the reaction practically vanishes at higher E_{coll} . Oxide ion substitution is highly indirect, as indicated by the cold product relative translational energy distributions shown in Fig. S3.† The differential labelling of the four H atoms enables us to distinguish between all protons in the course of a dynamics simulation. Hence, it is also possible to detect proton exchange reactions of OH⁻ and CH₃F, although the products are seemingly identical to the reactants. This proton exchange pathway begins with a proton abstraction by OH' from CH₃F, followed by a rotation of the HOH' fragment and a transfer of H'^+ to form CH₂H'F. When the reaction path is more indirect, proton exchange turns out to be a viable possibility as its ICS at E_{coll} =

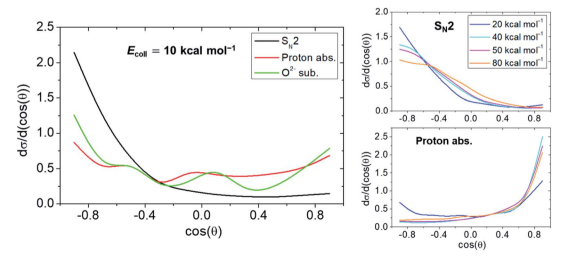


Fig. 4 Normalized scattering angle distributions of the $OH^- + CH_3F$ reaction are presented for S_N2 ($F^- + CH_3OH$), proton abstraction ($H_2O + CH_2F^-$) and oxide anion substitution ($HF + CH_3O^-$) at a collision energy of 10 kcal mol⁻¹. The angular distributions of S_N2 and proton abstraction are also shown at collision energies of 20, 40, 50 and 80 kcal mol⁻¹.

5 kcal mol $^{-1}$ is lower by only \sim 0.9 bohr 2 than the corresponding value of S_N2 . Note that in a few cases we find that proton exchange can be followed by oxide ion substitution leading to the HF + CH $_2$ H'O $^-$ products. Besides the Walden-inversion S_N2 pathway, we also find trajectories with retention of the initial CH $_3$ F configuration (see Table S2 \dagger). At low collision energies retention proceeds with double inversion, whereas at high $E_{\rm coll}$ front-side attack becomes the dominant retention pathway with about 3% probability of the total reactivity at $E_{\rm coll}=80$ kcal mol $^{-1}$.

Scattering angular distributions of selected $OH^- + CH_3F$ reaction pathways determined at various E_{coll} are displayed in Fig. 4. Details of the calculations are provided in the ESI.† Owing to the structure of the WaldenTS, S_N2 results in backward scattered products with the highest proportion being at $E_{coll} = 10$ kcal mol^{-1} . As for the proton abstraction, the increasing E_{coll} supresses backward scattering, and at sufficiently high values, forward scattered products become prominent signifying a direct stripping mechanism. On the other hand, at an E_{coll} of 10 kcal mol^{-1} , the oxide ion substitution displays a largely isotropic angular distribution. Thus, the mechanism is proven indirect, as the reaction can be easily trapped in the deep minimum of the PostHMIN1 complex.

Conclusions

The present work reveals a high-level dynamics investigation of the OH⁻ + CH₃F reaction consisting of nearly 1 million fulllength QCT simulations at several Ecoll on a global ab initio full-dimensional PES developed using the in-house Robosurfer program package.³⁶ At lower E_{coll} , dynamics simulations reveal the existence of a novel exothermic reaction pathway forming the unexpected product pair of HF and CH₃O⁻. This indirect oxide ion substitution is enabled by an energetically trapped region, the deep well of the PostHMIN1 complex, where the proton from the hydroxyl group of CH₃OH can be removed by the fluoride ion. Identified S_N2 reactions from the direction of PostHMIN1 indicate that the relevance of the deep potential well has been previously underestimated by Hase and coworkers.6 This study is aimed to emphasise the importance of theoretical investigations on fundamental reactions exploring new possible pathways, and calls for a further experimental verification of the obtained results. As the masses of the product anions, considering the most abundant isotopes, F (19 u), CH₂F⁻ (33 u), and CH₃O⁻ (31 u), are different, the experimental detection of the oxide ion substitution channel is possible. Note that a recent crossed-beam study could measure reaction channels with branching ratios less than 2%,37 which further supports the future experimental observation of the lowprobability oxide ion substitution. Furthermore, similar postreaction proton-transfer processes revealed in the present work may occur in other chemical reactions as well, providing novel product channels. On the one hand, one may consider different leaving groups for the OH + CH₃Y reaction such as Y = Cl, Br, I, NH₂, CN, etc. In the case of Y = Cl, Br, and I, the HY +CH₃O⁻ formation is less possible due to the higher and higher thermodynamic preference of the S_N2 channel as the atomic

number of Y increases, although, for Y = NH₂ and CN oxide ion substitution may occur due to the high proton affinity of NH₂ and CN $^-$. On the other hand, various nucleophiles such as SH $^-$, PH₂ $^-$, NH₂ $^-$, CCH $^-$, AsH₂ $^-$, CH₃ $^-$, SiH₃ $^-$, GeH₃ $^-$, SeH $^-$, etc. 15,26 may favour post-reaction proton transfer. One of the most promising candidates is the SH $^-$ + CH₃F system, where the exothermic HF + CH₃S $^-$ path is thermodynamically more favoured than the endothermic F $^-$ + CH₃SH S_N2 channel. Obviously, future reaction dynamics investigations will be necessary to determine the probability of post-reaction hydrogen-bonded complex formation which facilitates novel proton-abstraction induced product formation.

Methods

Benchmark ab initio thermochemistry

The stationary points of the OH⁻ + CH₃F reaction are searched using the second-order Møller-Plesset perturbation theory (MP2) with the aug-cc-pVDZ basis set. Then the explicitly-correlated coupled-cluster singles, doubles, and perturbative triples (CCSD(T)-F12b) method was employed with the aug-cc-pVDZ and aug-cc-pVTZ basis sets in order to improve the accuracy of the structures, energies and frequencies of the stationary points. The benchmark classical energies are computed at the CCSD(T)-F12b/aug-cc-pVTZ geometries using the CCSD(T)-F12b method with the aug-cc-pVQZ basis set, as well as considering post-CCSD(T) correlation effects up to CCSDT(Q) and core correlation effects. The benchmark relative energies of the stationary points are adapted from ref. 31, and FSTS2, PostHMIN1' and PostHMIN2 are newly characterized in this work.

Development of the potential energy surface

For the development of the global analytical ab initio PES of the OH + CH₃F reaction the same procedure is utilized as in the case of the OH⁻ + CH₃I reaction described in ref. 35. The PES development is performed with the Robosurfer program package at the MP2/aug-cc-pVDZ level of theory. The first step is to generate an initial dataset by modifying the geometries of the stationary points randomly. Then, this dataset is used to start the Robosurfer program, which adds the remaining points by iterative selection of the energies of the new geometries derived from QCT simulations. Performing 655 iterations overall at 1, 5, 10, 20, 30, 40, 50, 60, 70 and 80 kcal mol^{-1} E_{coll} , the dataset contains 50 434 energy points. The permutationally invariant polynomial approach is used for fitting the PES applying a fifthorder polynomial expansion of Morse-like variables, $\exp(-r_{ii}/a)$, where r_{ij} represents the inter-atomic distances and a = 3 bohr. The 4693 polynomial coefficients of the fit are determined by a weighted linear least-squares fit applying the weight function of $E_0/(E + E_0) \times E_1/(E + E_1)$, where $E_0 = 94 \text{ kcal mol}^{-1}$, $E_1 =$ 314 kcal mol^{-1} , and E is the energy relative to the global minimum. Similar to the OH + CH3I reaction, at certain geometries the gold-standard CCSD(T) breaks down providing too negative energies, therefore to solve this problem, the energy points are recalculated at the CCSD-F12b/aug-cc-pVTZ +

BCCD(T)/aug-cc-pVDZ - BCCD/aug-cc-pVDZ composite level of theory. Moreover, we add about 1000 composite *ab initio* energy points to improve the HF + CH₃O⁻ product region of the PES. The final PES contains 51 431 energy points and the root-mean-squared fitting errors are: 0.83 kcal mol⁻¹, 1.69 kcal mol⁻¹, and 2.54 kcal mol⁻¹ for the energy ranges 0–94 kcal mol⁻¹, 94–188 kcal mol⁻¹ and 188–471 kcal mol⁻¹, respectively.

Quasiclassical trajectory calculations

QCT computations are carried out for the OH⁻ + CH₃F reaction at ten E_{coll} : 5, 10, 15, 20, 30, 40, 50, 60, 70 and 80 kcal mol⁻¹ using a time step of 0.0726 fs. The ground vibrational states of OH and CH₃F are prepared by standard normal-mode sampling and the rotational angular momenta are set to 0. The initial distance of the reactants is 25 bohr with a given impact parameter, b, and the initial spatial orientation of the reactants is randomly sampled. At each b, 5000 trajectories are run and b is scanned with the step size of 0.5 bohr from 0 to $b_{\rm max}$, where the probability of the reaction becomes 0. Each trajectory is propagated until the largest interatomic separation becomes larger by 1 bohr than the largest initial one. The identification of the S_N2 reaction, which does not avoid the CH₃OH···F deep well (PostHMIN1), is based on a similar method used for the numerical separation of the front-side attack and double-inversion mechanisms.38

A more detailed description of the computations is given in the ESI.†

Data availability

The datasets generated and analysed during the current study are not publicly available due to their large size but are available from the authors on reasonable request.

Author contributions

G. C. designed the research, D. A. T. carried out the computations and analysed the data, D. A. T. and G. C. discussed the results, and D. A. T. wrote the paper.

Conflicts of interest

There are no conflicts to declare.

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