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Atomistic dynamics of elimination and nucleophilic substitution disentangled for the $F^- + CH_3CH_2CI$ reaction

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Chemical reaction dynamics are studied to monitor and understand the concerted motion of several atoms while they rearrange from reactants to products. When the number of atoms involved increases, the number of pathways, transition states and product channels also increases and rapidly presents a challenge to experiment and theory. Here we disentangle the dynamics of the competition between bimolecular nucleophilic substitution (S_N 2) and base-induced elimination (E2) in the polyatomic reaction $F^- + CH_3CH_2CI$. We find quantitative agreement for the energy- and angle-differential reactive scattering cross-sections between ion-imaging experiments and quasi-classical trajectory simulations on a 21-dimensional potential energy hypersurface. The *anti*-E2 pathway is most important, but the S_N 2 pathway becomes more relevant as the collision energy is increased. In both cases the reaction is dominated by direct dynamics. Our study presents atomic-level dynamics of a major benchmark reaction in physical organic chemistry, thereby pushing the number of atoms for detailed reaction dynamics studies to a size that allows applications in many areas of complex chemical networks and environments.

hemical reactions of small molecules can today be understood with an impressive level of detail¹⁻³. However, once more than four atoms are involved in a chemical reaction, the atomistic dynamics of bond cleavage and bond formation become challenging to experiment and theory, as the number of involved vibrational modes, reaction pathways, transition states and product channels increases rapidly^{4,5}. The fundamental process to clarify is the motion and rearrangement of all involved atoms during a reaction, for all posssible pathways⁵⁻¹². A ubiquitous example is the competition of bimolecular nucleophilic substitution (S_N2) versus base-induced elimination (E2). Both reactions are part of numerous synthesis protocols in organic chemistry due to their stereospecificity. The dynamics of these reactions therefore constitute major benchmark systems in physical organic chemistry¹³⁻¹⁸. Separating S_N2 and E2 reactions is complicated by the fact that several important aspects affect both reactions similarly: the attacking nucleophile/base, the type of leaving group and the polarity of the solvent. Only the steric environment at the reaction centre allows one to directly manipulate the branching ratio. Studies under single-collision conditions, which allow focusing on the intrinsic dynamics while excluding solvation effects, are complicated by the fact that both reactions usually form the same product ion and cannot be distinguished by standard mass spectrometry. Several elaborate methods have therefore been employed to distinguish between the two reactions in the gas phase^{13,14,17,19}. Interestingly, $S_N 2$ leads to the thermodynamically more stable product while E2 features the lower transition state barrier, that is E2 forms the kinetically preferred product. S_N2 and E2 reaction coordinates in the gas phase commonly follow a double-well

path with submerged barriers accompanied by entrance and exit channel complexes^{18,20} as opposed to a single barrier in solution. Energy partitioning for gas-phase $S_N 2$ reactions was found to often occur non-statistically²¹. In certain cases the reaction may even proceed by avoiding its minimum energy structures altogether²².

A model reactant that allows for both elimination (reaction (1)) and nucleophilic substitution (reaction (2)) is CH_3CH_2Cl with its primary carbon centre (exothermicities are taken from ref.¹⁵ and ref.²³). The ethyl residue is not so sterically congested at the α -carbon as to fully suppress the S_N2 reaction (Fig. 1a):

$$F^- + CH_3CH_2Cl \rightarrow Cl^- + HF + C_2H_4 \quad \Delta E(E2) = -0.96 \text{ eV}$$
(1)

$$\rightarrow \text{Cl}^- + \text{CH}_3\text{CH}_2\text{F} \quad \Delta E(\text{S}_{\text{N}}2) = -1.44 \text{ eV}$$
(2)

 $S_N 2$ reactions in the gas phase are known to have rich and complex atomistic dynamics²⁴. Recently, we have also identified fingerprints for atomistic dynamics of E2 reactions, using measurements of their differential scattering cross-sections^{17,25}. Trajectory simulations for elimination have so far been limited to on-the-fly dynamics^{16,26} with statistics not sufficient for a direct comparison with the experiment. The reaction coordinates for the three most important pathways, $S_N 2$, *anti*-E2 and *syn*-E2, feature submerged barriers with *anti*-E2 having the least congested transition state (Fig. 1a)^{15,23}. $S_N 2$ and *anti*-E2 are connected by a shared ion-dipole-bound

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Fig. 1 Reaction of $F^- + CH_3CH_2Cl. a$, Minimum energy pathway illustrating the three most important reaction channels: *anti*-E2 (dashed red), *syn*-E2 (dotted orange) and $S_N 2$ (solid blue). Structures show the atomic arrangement in the transition states (TSs) and pre-reaction minima. For $S_N 2$, the classic Walden inversion TS is depicted³². The *syn*- and *anti*-E2 configurations are termed after the relative orientation of the attacking base (F⁻), the carbon-carbon bond and the leaving group (Cl) in the TS. The E2 TS requires a planar arrangement of the three involved bonds of CH₃CH₂Cl: the C_p-H bond, the C-C bond and the C_a-Cl bond (minimum energy pathway and structures adapted from refs.^{15,23}). **b**, Velocity distributions of Cl⁻ product ions at 0.35 eV collision energy are shown from experiment and simulation together with a sketch of a Newton diagram depicting the relative orientations of reactant and product beams. The dashed arrows indicate the velocities of the neutral products with the red arrow illustrating the recoil of HF and C₂H₄ from each other. Forward (fw) and backward (bw) are defined along the original direction of the CH₃CH₂Cl beam.

pre-reaction minimum^{15,27}, whereas in *syn*-E2 a hydrogen-bonded complex is found¹⁵.

It can be misleading to only use stationary properties to predict atomistic mechanisms, direct versus indirect dynamics and energy partitioning²⁸. To investigate $S_N 2$ and E2 reactions at this level of detail, experiments and theory need to directly probe the atomic-level dynamics. In the case of elimination, tracing the internal excitation of the molecular products can be obscured by the three-body break-up (Fig. 1b). For experiments under single collision conditions that probe one ionic product, only the combined internal excitation of both molecular products, together with their translational recoil, can be extracted. Trajectory simulations with sufficiently high statistics can be used to disentangle the three contributions.

Results

We use experimental differential cross-sections from ion-molecule crossed-beam 3D velocity map imaging experiments and quasi-classical trajectory (QCT) simulations on a new fulldimensional (21-dimensional) potential energy surface to study the F⁻+CH₃CH₂Cl reaction. The combination of both methods allows for the kinematically complete characterization of the $S_N 2$ and E2 reactions as well as their competition. Experimental angleand energy-differential cross-sections for Cl⁻ product ions are presented in Fig. 2a-e. The Cl- velocity distributions are shown in the centre-of-mass frame for five relative collision energies E_{rel} in the range from 0.35 to 1.96 eV. At the lowest collision energy a clear forwardbackward symmetry is visible together with widespread isotropic scattering and isotropic scattering into low product ion velocities around the centre of mass (Fig. 2a). This scattering feature is typical for indirect S_N2 reactions²⁹. The forward-backward symmetric events are due to a large impact parameter indirect, complex-mediated mechanism that was recently identified in another E2 reaction²⁵. With increasing collision energy the forward-backward symmetry is lifted. The images become dominantly isotropic with some scattering into the forward hemisphere ($\theta = 0-90^{\circ}$) at small angles (Fig. 2b-e). The integral angular distributions (Fig. 2f-j, black) confirm the trend towards forward scattering with some minor backward scattering appearing at the two highest collision energies (Fig. 2i,j). Forward reactive scattering is a fingerprint of direct E2 dynamics^{17,25}. Backward scattering is a signature of the direct rebound mechanism known for S_N^2 (refs. ^{28,29}).

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Fig. 2 | **Differential scattering cross-sections. a-e**, Two-dimensional representation of the experimental 3D scattering distribution of ${}^{35}Cl^-$ product ions at collision energies of 0.35 eV (**a**), 0.83 eV (**b**), 1.14 eV (**c**), 1.61 eV (**d**) and 1.96 eV (**e**). The colour scales are shown on the right of the images. Superimposed circles represent the kinematic cutoff for S_N2 (orange) and E2 (red). The Newton diagram is given in Fig. 1b. Forward scattering is defined for a scattering angle $\theta = 0-90^{\circ}$ and backward scattering for $\theta = 90-180^{\circ}$. **f-j**, Velocity integrated angular distributions corresponding to **a-e**, respectively. **k-o**, Product ion kinetic energy distributions E_{kin} corresponding to **a-e**, respectively. In the middle and bottom panels, the total distributions, including all product channels forming Cl⁻, are given for experiment and theory in black and green, respectively. Individual contributions from E2 (red, combined *anti*-E2 and *syn*-E2) and S_N2 (orange) are scaled according to their respective integral cross-sections (Supplementary Table 3). Dashed lines in the bottom panels represent the kinematic (kin.) cutoffs for E2 (red) and S_N2 (orange). Experimental data for E_{rel} =1.96 eV taken from ref. ¹⁷.

Cuts in velocity space confirm the presence of the direct rebound by the characteristic umbrella of events in the backward hemisphere at high Cl- velocities (Supplementary Fig. 2). Integral angular distributions from the QCT simulations (Fig. 2f-j, green) quantitatively reproduce the experiment at all five collision energies. The remaining differences are well accounted for by the statistical and systematic uncertainties of both simulation and experiment. The simulations capture the forward-backward symmetry and isotropic contribution at the lowest collision energy and the shift towards forward scattering at higher collision energies. The experimental and simulated product ion kinetic energy distributions (Fig. 2k-o; experimental (Exp.): black, simulated (Sim.): green) also show very good agreement. The distributions reveal a broadening with increasing collision energy, while the maxima do not markedly shift. This decoupling of product ion kinetic energy from reactant collision energy has been identified as a signature for indirect dynamics in the case of pure E2 reactions²⁵.

Discussion

The excellent agreement between experiment and simulation allows us to disentangle the competition between $S_N 2$ and E2, plotted in orange and red in Fig. 2f–o. At 0.35 eV both reactions scatter into all angles (Fig. 2f) but with different dynamic signatures: E2 leads to forward–backward symmetric scattering and $S_N 2$ to almost isotropic scattering. This follows patterns known from $S_N 2$ reactions of CH₃Cl and E2 reactions of (CH₃)₃CCl with F⁻ (refs. ^{25,28}) and is ascribed to complex-mediated indirect dynamics. With increasing collision energy, simulations reveal more anisotropic angular distributions (Fig. 2h–j). This is attributed to a growing contribution of direct scattering. Whereas scattering into the forward hemisphere is almost pure elimination at all collision energies (>80%), S_N^2 contributes up to half of the events scattered into the backward hemisphere at the two highest collision energies. This superposition causes the seemingly wholly isotropic indirect character of the experimental velocity images. The prevalence of E2 over S_N^2 dynamics is also seen in the much larger range of attack angles that lead to the elimination pathway (Supplementary Fig. 5).

To test the contribution of direct reaction dynamics to the overall scattering, the fraction of simulated trajectories that swiftly form products without forming a transient intermediate complex is determined. This confirms that E2 and $S_N 2$ shift from highly indirect to direct character with increasing collision energy (Fig. 3a). Comparable direct fractions have been found for the pure $S_N 2$ reaction of $F^- + CH_3CI$ (ref. ²⁸). By contrast, E2 reactions with the bulkier reactant (CH₃)₃CCI are much more indirect²⁵. Reactions of ethyl halides with different bases and leaving groups show similar direct fractions^{16,17,26,30}.

Besides the Cl⁻ product ion, $FH\cdots Cl^-$ and trace amounts of CH_3CHCl^- are also found in both experiment and simulation, following reaction (3) and reaction (4):

$$F^{-} + CH_{3}CH_{2}Cl \rightarrow FH \cdots Cl^{-} + C_{2}H_{4}$$
$$\Delta E(FH \cdots Cl^{-}) = -1.93 \text{ eV}$$
(3)

$$\rightarrow$$
 CH₃CHCl⁻ + HF $\Delta E(PT) = 1.15 \text{ eV}$
(4)

Experimental product branching ratios, which are extracted by time-of-flight traces (Supplementary Fig. 3), compare well with

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Fig. 3 | Direct fractions and product branching ratios. a, Fraction of direct trajectories for *anti*-E2 (red), $S_N 2$ (orange) and their sum (green) as a function of relative collision energy. As a comparison, direct fractions for $F^- + CH_3Cl$ (ref. ²⁸; blue stars, experiment; blue open circles, simulation) and $F^- + (CH_3)_3CCl$ (ref. ²⁵; grey stars, experiment) are given. **b**, Product branching for Cl⁻ (green) and FH····Cl⁻ ions (black) (experiment, open symbols; simulation, filled symbols). The contributions to Cl⁻ of *anti*-E2, *syn*-E2 and $S_N 2$ are also displayed separately. FH····Cl⁻ stems solely from *syn*-E2. Contributions of CH₃CHCl⁻ product ions are not displayed for clarity. Supplementary Table 1 gives the numerical values and their accuracies for all three product ions.

the relative integral cross-sections for the two major product ions (Supplementary Table 3), as shown in Fig. 3b. The simulated $FH\cdots Cl^{-}$ fraction lies somewhat above the experimental fraction, which may indicate late dissociation of this product that is not

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captured within the time frame of the QCT simulations. $CH_3CHCl^$ is formed in an endothermic proton transfer (PT) (reaction (4)). It is most likely a transient species that can undergo 1,2-H-shift and subsequently form elimination products. Similar branching ratios for the three product ion species have been found for $F^- + CH_3CH_2I$ (ref. ¹⁷).

Figure 3b also shows the relative contributions for the $S_N 2$, anti-E2 and syn-E2 reaction pathways that have been extracted from the QCT results (see Methods for details). The anti-E2 reaction is by far the dominant reaction at all collision energies. While its contribution drops with increasing collision energy, that of the syn-E2 reaction rises. The S_N2 reaction becomes more important at higher collision energies accounting for 30% of the total reactivity at 1.96 eV. The reason for this behaviour is found in the reaction probability as a function of impact parameter (Supplementary Fig. 4). In the case of the anti-E2 reaction, these probability functions retain a significant contribution of large reactive impact parameters. This remains the case even at the highest collision energy, whereas at small impact parameters the S_N2 reaction has a higher probability. The anti-E2 and syn-E2 reactions give comparable angular distributions (Supplementary Fig. 7). Only at the lowest collision energy, is the forward-backward peaked signature of the anti-E2 reaction not present for the syn-E2 reaction. The syn-E2 reaction branches into two product ion channels: Cl- (reaction (1)) and FH…Cl- (reaction (3)). Both channels are formed with a comparable probability at low collisions energies, whereas Cl- formation dominates at high collision energies (Fig. 3b). This is likely to be a consequence of an increased dissociation probability of FH---Cl- into HF and Cl⁻, which requires ~1 eV energy (Fig. 1a). Experimental scattering images and QCT results for FH…Cl- product ions show mostly forward scattering (Supplementary Fig. 8).

Due to the three-body break-up in E2, the experiment cannot directly extract the amount of energy partitioned into internal excitation. Instead, the experimental data reveal the sum of the rovibrational excitation E_{int} of HF and C_2H_4 and the relative kinetic energy E_{recoil} from the recoil between the two neutral products (black line in Fig. 4). However, the trajectory simulations for the E2 reaction (red line in Fig. 4) allow one to separate the energy partitioning between HF and C_2H_4 internal excitation and their relative recoil energy. The QCT result deviates slightly from the experimental distribution because the contribution from S_N^2 reactions, which is part of the experimental data, has been filtered out. The total internal energy, the sum of HF and C_2H_4 internal excitation, is shown as grey shaded areas in Fig. 4. It remains essentially constant with increasing collision energy. However, the full experimental or simulated distribution, which include the recoil energy (black and red traces), extend



Fig. 4 | Internal energy E_{int} **distributions for the E2 reaction. a**-e, Distributions at collision energies of 0.35 eV (**a**), 0.83 eV (**b**), 1.14 eV (**c**), 1.61 eV (**d**) and 1.96 eV (**e**). In addition to the internal energy of HF and C_2H_4 , the experimental curve contains the relative recoil energy E_{recoil} between HF and C_2H_4 (experiment, solid black; simulation, solid red). Dashed curves show the individual internal energy distributions for HF (dashed blue) and C_2H_4 (dashed green) with the sum of both distributions given by the grey shaded areas. The difference between the red and grey distributions indicates the energy channelled into recoil of both neutral products. The blue bars illustrate the vibrational quanta in the HF stretching mode. Distributions are area normalized and simulations weighted by the branching ratio of E2 versus $S_N 2$ (blue/green normalized to 1; red/grey normalized to 2). Experimental data for $E_{rel} = 1.96$ eV taken from ref.¹⁷.

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to larger energies. This is direct evidence that most of the additional collision energy is partitioned into recoil energy between the two neutral products. The dashed curves show the separate internal energy distributions for HF (blue) and C_2H_4 (green). HF is predominantly formed in its vibrational ground state, irrespective of whether it is formed by *syn*-E2 or *anti*-E2 reaction (Supplementary Fig. 9). This is seen from the blue bars spaced by the vibrational quantum of the HF stretch. Hase and co-workers found a similar behaviour for the proton transfer forming ICH₂⁻ and HF in reactions of $F^- + CH_3I$ (ref. ³¹). The internal excitation of C_2H_4 increases only

slightly with collision energy, which is evidenced by a growing tail of the distribution. For the $S_N 2$ reaction, the comparison between the simulated internal excitation distribution for the CH_3CH_2F product and the experimental data is shown in Supplementary Fig. 10. With the reaction becoming increasingly direct when the collision energy is increased, the internal excitation decreases slightly.

Conclusion

The present study investigates a benchmark reaction for complex polyatomic reaction dynamics in a combined approach of experiment and theory. We have probed the differential cross-section to disentangle the energy dependent atomistic dynamics and competition of $S_N 2$ versus E2 reactions. The reaction of $F^- + CH_3 CH_2 Cl$ is dominated by direct dynamics despite the numerous internal degrees of freedom, except at the lowest collision energy. The energy partitioning in the three-body break-up of E2 reactions is complex, with a significant amount of energy channelled into the recoil of the two neutral products. It is remarkable that the reaction dynamics of this complex reaction can be disentangled into distinct atomic-level mechanisms that have been identified through their fingerprints on the differential cross-sections in systems that undergo either pure S_{N2} or only E2 reactions. This clearly supports the notion of a bottom-up approach to understanding multidimensional reaction dynamics on the way to real-life chemical reactions. Furthermore, with the intrinsic reactivity and dynamics on the ground-state potential energy surface fully characterized it becomes possible to track down the effects of manipulating such reactions, for example by functionalization, micro-hydration, or selective internal excitation of the reactants.

Online content

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Methods

Experiment. Reactive crossed-beam scattering in combination with 3D velocity map imaging³³ was used to record differential cross-sections. The experimental setup and experimental procedure have been described previously^{34,35}. In brief, ions are produced by a pulsed plasma discharge ignited in a supersonic beam (10% NF₃ in argon). F⁻ ions are extracted perpendicularly to the initial direction of the expansion by a Wiley–McLaren type electrode arrangement and loaded into an octupole radio frequency ion trap operated at room temperature. Ions are stored for 40 ms and collisions with N₂ buffer gas reduce the translational energy spread of the ion ensemble. Ions are ejected from the trap and guided into the interaction region of the velocity map imaging spectrometer. There, the ion beam is intersected by a skimmed molecular beam of CH₃CH₂Cl seeded at a couple of percent in helium. Densities in the interaction roulme are such that the experiment operates under single collision conditions.

Cl⁻ product ions are extracted perpendicular to the scattering plane and mapped onto a position and time-sensitive detector consisting of a multichannel plate/phosphor screen combination with a CCD (charge-coupled device) camera recording the position of impact. Simultaneously, the arrival time is recorded by a photomultiplier tube which detects the photon signal emitted by the phosphor screen. Cylindrical symmetry allows us to display the 3D velocity distributions in a 2D representation to better compare them with sliced images.

The energy and angular spreads of both reactant beams are determined using 2D velocity mapping and fitting of the resulting distributions. The translational temperature of the neutral beam (CH₃CH₂Cl in He) in the co-moving frame is ~150–200 K, which means that internal rotation of the methyl groups along the C-C axis is frozen out³⁶. The energy spread of the relative collision energy for the presented crossed-beam experiments is in the range 20–50 meV (1 σ of the energy distribution). A graphical representation of the experimental resolution at the collision energy of 0.35 eV is shown in Supplementary Fig. 1. The experimental uncertainty for the resulting product kinetic and internal energy distributions is determined by error propagation of the reactant beams³⁴ and ranges from 150 to 250 meV.

Theory. We have developed a full-dimensional (21-dimensional) analytical ab initio potential energy surface (PES) for the F⁻ + CH₃CH₂Cl reaction using the ROBOSURFER program system37 to automate the construction of the PES and the permutationally invariant monomial symmetrization approach (MSA)³⁸ to fit the energy points. The initial dataset of 5,319 structures was obtained by randomly modifying the Cartesian coordinates of the previously known¹⁵ stationary-point geometries. The ab initio energies were computed using the density-fitted explicitly correlated second-order Møller-Plesset perturbation method (DF-MP2-F12) with the aug-cc-pVDZ basis set. Then, ROBOSURFER iteratively developed the PES by (1) running quasi-classical trajectories at different collision energies and impact parameters, (2) selecting configurations along the trajectories that are most likely to improve the quality of the PES, (3) performing ab initio computations at the DF-MP2-F12/aug-cc-pVDZ level with MOLPRO (ref. 39) and (4) fitting the energy points using MSA. At the final stage of the PES development the energy points were recomputed with a higher, composite ab initio level of theory: CCSD(T)-F12b/aug-cc-pVDZ+MP2-F12/aug-cc-pVTZ-MP2-F12/ aug-cc-pVDZ. The gold-standard explicitly correlated CCSD(T)-F12b method was employed to provide an accurate description of electron correlation while the basis-set effects beyond aug-cc-pVDZ were described with the computationally less-expensive MP2-F12 method. The final PES was fitted to 35,474 energy points by a weighted linear least-squares method using a 5th-order polynomial expansion of Morse-like variables, $\exp(-r_{ij}/a)$, a = 3 bohr, of r_{ij} interatomic distances, resulting in 11,241 terms (coefficients).

The analytical PES developed in this study allows efficient and accurate reaction dynamics (QCT) simulations for the nine-atom $F^- + CH_3CH_2CI$ system. Standard normal-mode sampling⁴⁰ is used to prepare the quasi-classical vibrational ground state of the CH_3CH_2CI reactant molecule, thereby incorporating the quantum-mechanical zero-point energy effects into the classical simulations. The initial distance of the reactants was 25 bohr, their relative orientation was randomly sampled and impact parameters were scanned from zero to their maximum value with a step size of 0.5 bohr. We computed 1,000 trajectories at each impact parameter at each of the following collision energies: 0.35, 0.83, 1.15,

1.60 and 1.98 eV. Each trajectory was propagated until the largest interatomic separation became 1 bohr larger than the largest initial one. Cross-sections were computed using an impact-parameter-weighted numerical integration of the reaction probabilities over impact parameters.

More details of the experimental and theoretical methods are given in the Supplementary Information.

Data availability

Data are provided in the online material accompanying this article. Source data are provided with this paper.

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Author contributions

E.C., J.M. and M.S. carried out the experiment. J.M. and E.C. analysed the data. B.B. extended the data analysis program suite. T.M. and B.B. contributed to data discussion and interpretation. V.T. and T.G. developed the potential energy surface. V.T. performed the trajectory simulations and analysed the data. G.C. and R.W. supervised the project. J.M. and R.W. prepared the manuscript. V.T. and G.C. contributed the theoretical sections.

Competing interests

The authors declare no competing interests.

Additional information

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