Fundamental Aspects of Gas Phase Ion Chemistry
Studied Using the Selected Ion Flow Tube Technique

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Gas phase ion chemistry using the selected ion flow tube (SIFT) technique is discussed with the focus on fundamental studies on ion structure, energetics, reactivity, and dynamics: Laser-induced fluorescence (LIF) kinetics/dynamics of $N_2^+$ ions (vibrationally state-selected ion-molecule reactions; energy transfer processes), mechanisms of prototypical organic ion reactions ($S_{82}$ nucleophilic substitution and kinetic isotope effects; hydrogen/deuterium exchange; isotope effects in termolecular association), and ion-molecule reactions with biological implications ($S_{82}$ reactions at a nitrogen center; $E_{82}$ elimination reactions of alkyl hydroperoxides). Anion chemistry of small nitroalkanes is also presented. A powerful combination of SIFT and negative ion photoelectron spectroscopy is used to study thermochemistry and structure of energetic and exotic ions (alkyl peroxides; anionic derivatives of diazo and azole compounds, cyclooctatetraene, and cyclopentanone). Finally, some of the new directions in SIFT research are discussed, i.e., reactions of ions with polyatomic organic radicals, and chemical ionization mass spectrometry with new detection schemes.

1. Introduction

The selected ion flow tube (SIFT) is an MS-MS technique to investigate chemical reactions of mass-selected ions in a flowing afterflow environment. Mass-selected ions are thermalized by collision with buffer gas molecules (e.g., helium) prior to reaction with a neutral reagent added. Compared to conventional ion beam techniques, low collision energies with the SIFT technique allow studies of ion–molecule reactions under truly thermal conditions (300 K or $E_{kin} = 0.04$ eV). The technique was originally developed by Smith and Adams at Birmingham. A later review described some additional features to the original design, e.g., the VT–SIFDT (variable temperature-selected ion flow drift tube) apparatus to explore different collision-energy regimes. Graul and Squires thoroughly and comprehensively reviewed flow reactor techniques in general, including SIFT. The technique later incorporated laser-induced fluorescence and triple quadrupole detections for extended capabilities.

This article discusses the SIFT ion chemistry work at Colorado, with the focus on fundamental studies which the author has conducted in recent years. Following a brief description of the instrument, studies of vibrationally state-selected ion-molecule reactions and energy transfer processes are presented as examples of the most fundamental aspects of gas phase ion chemistry. Nucleophilic substitution ($S_{82}$) reactions and associated kinetic isotope effects, hydrogen/deuterium exchange reactions, and isotope effects in termolecular association reactions are subsequently discussed, with an emphasis on how isotope labeling can be used to probe the structure and dynamics of ion–neutral intermediate complexes. These sections are followed by discussions of individual ion chemistry in two categories. First, there are ion–molecule reactions with strong biological implications: $S_{82}$ reactions at a nitrogen center and a novel $E_{82}$ type of elimination reactions. Secondly, the technique has been used extensively to determine the thermochemistry and structure of energetic and exotic ions. The article concludes by presenting some of the most recent projects and new directions that are being pursued with the SIFT technique, i.e., reactions of ions with polyatomic organic radicals and new schemes for chemical ionization mass spectrometry.

2. SIFT Instrument

The selected ion flow tube apparatus used at Colorado is schematically illustrated in Fig. 1. Details of the design and operation have been given elsewhere. Ions are produced by using simple electron impact on neutral precursors or sequential ion-molecule reactions in the source flow tube containing helium ($\sim 0.3$ Torr). They are selected by the SIFT quadrupole mass filter and injected through an injection orifice into the second flow tube containing the helium buffer gas ($0.5$ Torr, $\sim 300$ K). The SIFT injection energy ($E_{inj}$), which is defined by the potential difference between the source flow tube and the injection orifice, is typically set to $20$ eV in order to cleanly inject the reagent ions with negligible fragmentation. Injected ions are quickly thermally equilibrated to $300$ K by collisions with the helium buffer gas. A laminar flow of He is established after the initial turbulent flow regime (30–40 cm in length). The ions entrained in He are then allowed to react with neutral gases added through multiple reactant inlets placed along the reaction flow tube. The length of the laminar region ($\approx 1$ m) and the flow velocity ($\approx 100$ ms$^{-1}$) gives a maximum reaction time of about 10 ms. Parent and product ions are analyzed.

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with a detection quadrupole mass filter placed at the end of the flow tube. Absolute reaction rate constants are measured by monitoring the kinetic decay of the parent ions while varying the reaction distance, i.e., the neutral inlet positions. The detector has a triple quadrupole capability that allows studies of collision-induced dissociation (CID) of ions. Alternatively, unimolecular dissociation can be effected by SIFT injection of ions at higher \( E_{\text{inj}} \); in this case ion dissociation takes place near the injection orifice in the high-pressure He environment. This mode of experiment ("SIFT-CID") involves multiple ion-He collisions, thus producing qualitative and yet informative results with high sensitivity. An additional feature with the SIFT instrument is the drift section in the second flow tube (Fig. 1). Ion mobilities and reactions at higher kinetic energies (up to \( E_{\text{cm}} \sim 1 \) eV) are examined in the electric field applied along the flow tube.\(^9\), \(^10\)

Compared to polyatomic ions, vibrational relaxation of small diatomic ions is extremely slow in He. Thus, ion chemistry of vibrationally excited ions can also be explored. The capability of the SIFT technique has been extended by incorporating high-resolution spectroscopic detection with laser induced fluorescence (LIF, Fig. 1)\(^6\), \(^11\). The SIFT–LIF technique allows studies of vibrationally state selected ion chemistry, e.g., vibrational specificity and energy transfer in ion-molecule reactions, at thermal energy (\( E_{\text{cm}} \approx 0.04 \) eV). The optical detection has a significant advantage over the conventional monitor ion technique,\(^12\) which chemically probes for vibrationally excited states using reactions that are fortuitously specific to certain excited states, e.g., \( \text{N}_2^+(v > 0) + \text{Ar} \rightarrow \text{N}_2 + \text{Ar}^+ \) without isotope specificity. Reactant ions are generated by electron impact in various vibrational states, and immediately extracted into the SIFT quadrupole chamber before collisional relaxation takes place. The ions are then mass selected and injected into the second flow tube, where they are rapidly thermalized translationally and rotationally while retaining vibrational excitation. Following a reaction with a reagent added through the sample inlet (Fig. 1), the reactant and product ion vibrational states are monitored using LIF. There are requirements for SIFT-LIF experiments to be feasible.\(^6\), \(^11\) Vibrationally excited reagent ions need to be readily generated and stably injected at high densities. Collisional quenching of vibrationally excited states must be sufficiently slow with the He buffer gas (although slow quenching with He can also be measured; see below) and the radiative quenching be also negligible within the reaction time scale of about 150 \( \mu \text{s} \) before the LIF detection.\(^6\) Small, non-polar \( \text{N}_2^+(v) \) ions have thus been studied extensively with this technique using the B–X electronic transition for LIF.

3. Vibrationally State-Selected Ion Chemistry

3.1 Vibrational energy transfer and relaxation

Vibrational relaxation of ions in collision with rare gas atoms (He, Ne, Ar, Kr, Xe) is slow. The ion–neutral interaction is relatively weak because the rare gas atoms have no permanent dipoles or chemical interactions with ions. Relaxation via vibration-to-vibration (V-V) or vibration-to-rotation (V-R) energy transfer does not occur with monatomic rare gas molecules, and only the vibration-to-translation (V-T) process is available for the disposal of ion vibrational energy. Compared to intermolecular potential wells for neutral–neutral systems, however, charge–induced dipole interactions facilitate deep potential wells for ion–neutral systems (typically \( \sim 0.2 \) eV). As the consequence, ion vibrational relaxation is generally more efficient than the corresponding neutral relaxation.\(^12\) The V-T energy transfer/relaxation with rare gas atoms is the simplest case to examine the effects of intermolecular potentials during ion–neutral versus neutral–neutral collisions. Because of the inherent experimental difficulties, ion vibrational relaxation has been considerably less explored than is relaxation of neutral molecules.\(^13\)-\(^15\)

We used the SIFT–LIF technique to measure vibrational relaxation of \( \text{N}_2^+(v) \) ions with rare gas atoms.\(^16\)
$N_2^+(v) + M \xrightarrow{k_{ct}(v)} N_2^+(v' < v) + M$ \hspace{1cm} (1)

$N_2^+(v) \xrightarrow{k_{ct}(v)} N_2 + M^+$ \hspace{1cm} (2)

The optical detection allows individual vibrational levels of $N_2^+(v)$ to be monitored. At an injection energy of $E_{inj} = 49$ eV, initial vibrational population distributions of $N_2^+(v)$ were determined by LIF to be $46\% (v=0)$, $23\% (v=1)$, $15\% (v=2)$, $12\% (v=3)$, $4\% (v=4)$, and less than $1\% (v=5)$. Following addition of reagent rare gases, evolution of these states can be measured and analyzed, and vibrationally specific quenching rate constants $k_{ct}(v)$ were determined (Eq. (1)). In practice, measuring slow rate constants is challenging because the measurement can easily be biased by experimental artifacts arising from trace impurities and ion mobility changes upon addition of large amounts of reagents. Also, there are concurrent charge transfer reactions (Eq. (2)) with Ar (for $v' > 0$), Kr, and Xe. Conventional mass spectrometry, on the other hand, is blind to the quenching process in Eq. (1) but can detect the charge transfer products in Eq. (2) as well as impurity reactions. We thus employed a powerful combination of SIFT-LIF and mass spectrometry to unambiguously determine the ion-rare gas relaxation rate constants.\(^{15}\)

The measured rate constants $k_{ct}$ for vibrational relaxation of $N_2^+(v=1) \rightarrow N_2^+(v=0)$ are very small: 9.8 $\pm$ 0.7 $\times 10^{-16}$, 3.6 $\pm$ 0.5 $\times 10^{-14}$, and 1.6 $\pm$ 0.2 $\times 10^{-12}$ cm$^3$ s$^{-1}$ for He, Ne, and Xe, respectively.\(^{16}\) Upper limits were obtained for Ar ($< 2 \times 10^{-11}$ cm$^3$ s$^{-1}$)\(^{15}\) and Kr ($< 2 \times 10^{-12}$ cm$^3$ s$^{-1}$).\(^{16}\) These values are one to two orders of magnitude smaller than previous measurements, which presumably were affected by impurities (reviewed in Kato et al.).\(^{16}\) Specifically, vibrational relaxation with Ar was measured for the first time using LIF. This process cannot be studied with the conventional monitor technique because Ar itself is the only monitor species available to distinguish between $N_2^+(v=0)$ and $N^+ (v > 0)$. It is confirmed that $N^+ (v > 0) +$ Ar charge transfer is an excellent monitor reaction; vibrational relaxation is negligibly slow compared to the fast charge transfer ($k_{ct}(v=0)$ = $4 \times 10^{-10}$ cm$^3$ s$^{-1}$).

For comparison with neutral-neutral vibration relaxation, the rate constant $k_{ct}$ is converted to the deactivation probability per collision, $\langle P \rangle = k_{ct} = k_{col}$, where $k_{col}$ is the collision rate constant. The values of $\langle P \rangle$ are extremely small, ranging from 1.7 $\times 10^{-6}$ (He) to 8.4 $\times 10^{-5}$ (Ne) and 1.6 $\times 10^{-4}$ (Xe) for collisions with $N_2^+(v=0)$. The inefficient relaxation is reasonable because the $N_2^+$ ion ($\omega_m = 2207$ cm$^{-1}$) has to dissipate the large vibrational energy solely into translation. The small probabilities imply that $N_2^+$ relaxation proceeds via transfer of a single vibrational quantum per collision, i.e., $v' = v - 1$, in accord with Landau-Teller theory for V-T energy transfer established for neutral-neutral collisions at thermal energy.\(^{13,14}\) There is a marked difference observed between relaxations of $N_2^+(v)$ and $N_2^+$ (however). Measured or estimated values of $\langle P \rangle$ for neutral $N_2(v)$ are 1.6 $\times 10^{-8}$ (He), 6 $\times 10^{-13}$ (Ne), and 1 $\times 10^{-14}$ (Ar)\(^{16}\) relaxation is more efficient for lighter collision partners. This trend is consistent with Landau-Teller theory for V-T energy transfer, which assumes the relaxation to occur at the repulsive part of the intermolecular potential. In contrast, observed $N_2^+(v)$ ion relaxation is more efficient for heavier collision partners and has a positive correlation with the rare gas polarizability. The magnitudes of the deactivation probability are also greater for $N_2^+(v)$ than for $N_2^+(v)$. The attractive, charge-induced dipole interaction must play a significant role in the $N_2^+(v)$ ion vibrational relaxation. We discussed the results in terms of direct versus complex-formation mechanisms.\(^{16}\)

The SIFT-LIF technique was extended to studies of $N_2^+(v)$ vibrational relaxation/energy transfer with diatomic molecules $N_2$,\(^{19-21}\) O$_2$,\(^{19,21}\) CO and NO,\(^{22}\) and H$_2$.\(^{23}\) In addition to V-T energy transfer, relaxation via V-V and V-R processes is possible for these diatomics; the vibrational constants $\omega_v$ are 2359 cm$^{-1}$ ($N_2$), 1580 cm$^{-1}$ ($O_2$), 2170 cm$^{-1}$ ($CO$), 1904 cm$^{-1}$ ($NO$), and 4395 cm$^{-1}$ ($H_2$) in comparison to 2207 cm$^{-1}$ ($N_2^+$). The attractive potential wells are especially deeper ($\sim$ 1 eV) with $N_2$, $O_2$, and NO due to the electron-exchange interactions.\(^{24,25}\) Efficient relaxation of $N_2^+(v)$ is thus expected for these reactants.

Rate constants $k_{ct}$ and probabilities $\langle P \rangle$ were determined for $N_2^+(v)$ vibrational relaxation, in competition with concurrent charge transfer ($N_2^+, O_2^+, CO^+$, and NO) and hydrogen atom transfer ($N_2^+ + H_2 \rightarrow N_2H^+ + H$). As expected, vibrational relaxation is generally more efficient with the diatomic molecules by orders of magnitude than with rare gas atoms, but there are interesting variations found among the different diatomics. Vibrational relaxation with H$_2$ is noticeably slow and only an upper limit of 0.06 was obtained for $\langle P \rangle$, on average for all vibrational levels of $N_2^+(v=0)$\(^{23}\). This may be reasonable in light of the poor vibrational energy match between $N_2^+$ and H$_2$ and also the small polarizability of H$_2$.

An intriguing comparison is made between $N_2^+(v) + NO$ and $N_2^+(v) + CO$.\(^{22}\) On average for $v > 0$, vibrational quenching is efficient for NO with $\langle P \rangle$ of 0.22 whereas the deactivation probability is only 0.04 for CO, even though CO and $N_2^+$ has a nearly perfect vibrational energy match for resonant V-V energy transfer. This suggests that ion vibrational relaxation does not always follow the resonance principles that are well known for neutrals.\(^{15}\) Instead, the lifetime of energized adducts is a more important factor than the V-V energy gap. The $N_2^+(v) + NO$ system is likely to have a deep attractive well due to the near energy-resonant, electron-exchange interaction between $N_2^+$ and electronically excited NO$^+$($\omega_S^+ \Sigma^+$).\(^{25}\) Efficient vibrational energy transfer may occur during the long complex lifetime. Relaxation of $N_2^+(v)$ with O$_2$ is also efficient and vibrational-level dependent, the value of $\langle P \rangle$ increasing from 0.15 ($v=1$) to 0.30 ($v=4$)\(^{21}\) A similar electron-exchange interaction with the excited O$_2^+$($\omega_{1\Pi}$) state has been suggested.\(^{25}\) Tachibana et al.\(^{26}\) proposed vibrational relaxation within the stable quartet $[N_2 \cdot O_2^+]$ complex, which is computationally shown to be formed from $N_2^+$ and O$_2$ and correlate with $N_2 + O_2^+$($\omega_{1\Pi}$).

The SIFT-LIF technique revealed another novel aspect in ion-molecule interactions: multiquantum vibrational transfer/deactivation in collision of $N_2^+(v)$
with N2 and O2.\textsuperscript{11,19-21} The [N2 • N2\textsuperscript{+}] adduct formed from N2\textsuperscript{+} + N2 is strongly bound by about 1.1 eV.\textsuperscript{21,28} In collision of vibrationally excited \[^{15}\text{N}_2\textsuperscript{+}(v=2)\] ions with \(^{14}\text{N}_2\), the reactant and product ion vibrational states were distinguished by high-resolution LIF with isotope specificity. We observed simultaneous transfer of charge and vibration in this near-energy resonant charge transfer reaction (Eq. (3)).\textsuperscript{19} Most notably, two vibrational quanta are transferred efficiently in a single collision.

\[^{15}\text{N}_2\textsuperscript{+}(v=2)+^{14}\text{N}_2(v'=0) \rightarrow ^{15}\text{N}_2(v=0)+^{14}\text{N}_2\textsuperscript{+}(v'=2) \tag{3}\]

A detailed kinetic analysis including all reactant vibrational states \((v=0-4)\) indicates that approximately 10% of the initial \[^{15}\text{N}_2\textsuperscript{+}(v=2)\] forms \[^{14}\text{N}_2\textsuperscript{+}(v'=2)\] (Eq. (3)) and, by symmetry, another 10% forms \[^{14}\text{N}_2\textsuperscript{+}(v'=2)\].\textsuperscript{21} The observation that the \[^{15}\text{N}_2\textsuperscript{+}(v=2)\] ion relaxes via \(|\Delta v|=2\) transitions with such a high efficiency \((P \approx 0.2)\) is a marked result. For neutral-neutral collisions, only the \(|\Delta v|=1\) transition is allowed at thermal energy to first order within the framework of Landau-Teller theory.\textsuperscript{13,14}

A possible mechanism is direct V-V energy transfer between the high-frequency moieties in the long-lived \[^{15}\text{N}_2-^{14}\text{N}_2\textsuperscript{+}\] complex, as inferred from the similar vibrational constants for \[^{15}\text{N}_2\textsuperscript{+}\] and \[^{14}\text{N}_2\textsuperscript{+}\]. Alternatively, multiquantum vibrational transfer may result from near-resonant, intracomplex charge-exchange transitions \[^{15}\text{N}_2\rightarrow^{14}\text{N}_2\textsuperscript{+} \rightarrow^{15}\text{N}_2\textsuperscript{+} \rightarrow^{14}\text{N}_2\textsuperscript{+}\]. Extensive quantum mechanical calculations of the state-to-state cross sections\textsuperscript{9,29} support the importance of \(|\Delta v|=2\) transitions in thermal energy \[^{15}\text{N}_2\textsuperscript{+}(v)+^{14}\text{N}_2\] collisions, along with facile single-quantum V-V and V-T processes which were also confirmed from analysis of the LIF kinetics.\textsuperscript{21}

Multiquantum vibrational transfer/relaxation was also indicated in the \[^{15}\text{N}_2\textsuperscript{+}(v)+^{14}\text{N}_2\textsuperscript{+}\] reaction from SIFT-LIF rate kinetics for \[^{15}\text{N}_2\textsuperscript{+}(v=0-4)\].\textsuperscript{21} As noted above, a deep potential well facilitated by the intermediacy of electronically excited \(^{12}\text{O}_2\textsuperscript{+}(\pi\text{II})\) must be responsible for the observation. By comparison, ion deactivation proceeds exclusively \(via\) the \(|\Delta v|=1\) transition for NO\textsuperscript{+}(v) + M systems,\textsuperscript{30} which have shallow potential wells. In this vein, deactivation of \[^{15}\text{N}_2\textsuperscript{+}(v)+\text{NO}\textsuperscript{+}\] may also proceed \(via\) multiquantum vibrational energy transfer. Unfortunately, for this system the fast charge transfer (Eq. (2)) precludes detailed analysis of the \(v=2 \rightarrow v=0\) versus \(v=2 \rightarrow v=1\) components in vibrational deactivation (Eq. (1)).

### 3.2 Vibrational specificity in chemical reactions

A dramatic enhancement in charge-transfer rates was observed by vibrational excitation of the \[^{15}\text{N}_2\textsuperscript{+}\] ion in the \[^{15}\text{N}_2\textsuperscript{+}(v=0-4)\] + Kr reactions at thermal energy;\textsuperscript{17} the rate constant increases by a factor of \(\approx 80\) in going from \(v=0\) to \(v=4\). The observed increase is almost entirely due to enhanced charge transfer since the vibrational relaxation has been shown to be negligibly slow.\textsuperscript{16} This is a particularly interesting system because charge transfer is highly exothermic for vibrational ground state \[^{15}\text{N}_2\textsuperscript{+}\] and yet the \(v=0\) charge transfer is surprisingly slow \((h=1.0 \times 10^{-12} \text{ cm s}^{-1})\), occurring once in 800 collision events.
Condon factors within the transient (\(N_2 - Kr\)) \(^+\) pair, the vibrational wave functions of \(N_2^+\) and \(N_2\) are displaced relative to each other. In this instance the \(N_2^+\) wave function is simply shifted from the equilibrium inter-nuclear distance \(r_e\) of 1.0977 Å by a short displacement of \(d = 0.02\) Å (Eq. (6)). While the \(N_2^+\) wave function is fixed at \(r_e = 1.1164\) Å. The Franck–Condon overlap is consequently altered.

\[
q_{\nu\nu'} = \left| \Psi_{N_2^+}(r) \Psi_{N_2}(r+\delta \nu) \right|^2
\]  
(6)

The rate constants for \(N_2^+(v)+Kr\) were calculated from the most energy-resonant product channels \((\Delta E < 100 \text{ meV})\), in this case those producing \(N_2(v+3)+Kr^+(P_{1/2})\) (Eqs. (7)–(11)). This is a reasonable approximation in the spirit of Rapp-Francis theory, as discussed in Kato et al.,[37] that charge transfer processes become extremely sensitive to the energy gap at low collision energies.

\[
\begin{align*}
N_2^+(v=0) + Kr &\rightarrow N_2(v=3) + Kr^+(P_{1/2}) + 59 \text{ meV} \\
N_2^+(v=1) + Kr &\rightarrow N_2(v=4) + Kr^+(P_{1/2}) + 51 \text{ meV} \\
N_2^+(v=2) + Kr &\rightarrow N_2(v=5) + Kr^+(P_{1/2}) + 41 \text{ meV} \\
N_2^+(v=3) + Kr &\rightarrow N_2(v=6) + Kr^+(P_{1/2}) + 32 \text{ meV} \\
N_2^+(v=4) + Kr &\rightarrow N_2(v=7) + Kr^+(P_{1/2}) + 22 \text{ meV}
\end{align*}
\]  
(7–11)

The “distorted Franck–Condon” model was originally developed by Lipeles to explain anomalous product vibrational state distributions in slow ion-molecule collisions.[34] We extended his approach to predict state-selective reaction cross sections at low \(E_{\text{cm}}\), where the Rapp–Frances treatment starts to fail. We also applied this model to successfully reproduce the \(N_2^+(v=1 \rightarrow 4) + Ar\) charge transfer rates.[35] The measured rate constants are nearly unchanged over the exothermic channels \(N_2^+(v=1 \rightarrow 4) + Ar \rightarrow N_2 + Ar^+\), a result unexplained by standard Franck–Condon arguments.

Vibrational specificity in reactions of \(N_2^+(v)\) with diatomic reagents was also studied. Upon vibrational excitation, the charge-transfer component for the \(N_2^+(v) + CO\) reaction[32] increases considerably whereas the \(N_2^+(v) + NO\) and \(N_2^+(v) + O_2\) charge transfer rates[18,21] show only moderate increases. These observations are intriguing since, for both the NO and O₂ reactions, electronically excited states of the ion products become energetically accessible, \(\text{i.e., } NO^+(a^3Σ^+)^+\) from \(N_2^+(v \geq 1)\) and \(O_2^+(a^3Π\text{L})\) from \(N_2^+(v \geq 2)\). In contrast, for the \(N_2^+(v) + CO\) reaction, the lowest electronically excited product ion state \((2A^1Π\text{L})\) is not thermodynamically accessible. The moderate increase in the \(N_2^+(v \geq 2) + O_2\) charge transfer rate has been explained theoretically by state coupling \(via\) the quartet \([N_2 \cdot O_2]^+\) complex[26] and also by wavepacket dynamics.[35] A guided ion beam experiment reported an anomalously small charge-transfer reactivity for \(N_2^+(v=1) + O_2\).[26] However, such a decrease is not observed in other studies using mass spectrometric kinetics (reviewed in Kato et al.[36,37] and photoelectron–photoion coincidence,[37]) and the discrepancy remains unresolved.

We also studied vibrational specificity in the hydrogen-atom transfer reaction \(N_2^+(v) + H_2 \rightarrow N_2H^+ + H\).[38] The reaction rates are large and nearly unchanged with respect to the reactant vibrational excitation. The fact that the rates significantly exceed the Langevin collision limit suggests that the reaction proceeds \(via\) initial charge transfer at large intermolecular distances producing \(N_2 + H_2^\circ\), followed by proton transfer at a shorter range. The Langevin rate is greater for \(N_2 + H_2^\circ\) than for \(N_2^+ + H_2\). Vibrational specificity may well be smeared out during these sequential processes in a single collision event.

4. Dynamics of Organic Ion-Molecule Reactions

4.1 Nucleophilic substitution (SN2) reactions

Nucleophilic substitution is one of the most fundamental chemical transformations in solution, and there have been numerous gas phase studies as well.[39] In the absence of solvent molecules, gas phase SN2 studies reveal intrinsic features of this reaction (mechanism and energetics, transition state structure, energy transfer/randomization, product dynamics) as well as provide key calibration for theory. This part discusses mechanisms and dynamic effects in SN2 reactions in some detail to provide a background for later sections. SN2 reactions of methyl halides with halide anions, \(X^- + CH_3Y \rightarrow CH_3X + Y^-\), represent the simplest system in this category. The SIFT technique was used to detect the occurrence of a slow, symmetric SN2 reaction \((X = Y = Cl)\) in the gas phase.[39]

\[
^{35}Cl^- + CH_3Cl \rightarrow CH_3^35Cl + ^{35}Cl^-
\]  
(12)

The reaction is extremely slow at room temperature with the measured rate coefficient of \(k \approx 3.5 \times 10^{-14} \text{ cm}^3 \text{ s}^{-1}\) (one reaction in \( \approx 50000\) collisions). The remarkable slowness of the thermoneutral SN2 reaction is in sharp contrast to the general rapidity of proton-transfer (PT) reactions, which occur at nearly every collision for many systems that are thermoneutral or near-thermoneutral.[40]

Figure 3 schematically shows potential energy curves for SN2 and PT processes, for which the reactions are assumed to be similarly exothermic by \(\Delta E_{\text{exot}}\). The transition state (TS) for proton transfer is located deep in the ion-molecule attractive potential well (Fig. 3a); the TS barrier is as low as a few kcal mol \(^{-1}\) or virtually nonexistent for typical PT systems, \textit{e.g.,} those involving oxo acids.[41] In contrast, for SN2 transformation involving Walden inversion (Fig. 3b), the central energy barrier \((\Delta E_{\text{exot}})\) is considerably high and comparable with the well depth (complexation energy \(\Delta E_{\text{comp}} \approx 10 - 25 \text{ kcal mol}^{-1}\)).[42] The double-well scheme was first proposed by Olmstead and Brauman[43] for solvent-free SN2 reactions in the gas phase. Even when the transition state is somewhat lower in energy than the reactants \(\text{i.e.,} \) the overall energy barrier \(\Delta E_{\text{exot}} < 0\), back-dissociation of the reactant complex \([X^+ + CH_3Y^-]\) is favored over Walden inversion to form the product complex \([XCH_3\cdots Y^-]\), based on a simple density-of-states argument. The reaction efficiency can thus be
considerably small. The slow SN2 reaction (Eq. (12)) is dramatically enhanced as the ion–molecule collision energy is increased in the SIFT-drift field, stimulating fundamental interest in the mechanism of gas phase nucleophilic substitution. A high-level ab initio theoretical study suggested this behavior as evidence for the occurrence of a novel “front-side” SN2 mechanism, in which 37Cl approaches the carbon atom from the same side as 35Cl leaves (Fig. 4b). At the G2(+) level of theory the energy threshold for the front-side attack was predicted to be 46.3 kcal mol⁻¹. However, guided ion beam experiments under single collision conditions measured the threshold for the translational activation to be 11 ± 4 kcal mol⁻¹, significantly lower than the theoretical front-side threshold. Therefore, the front-side mechanism must not be reasonable. The observed enhancement can then be attributed to translational activation of the conventional back-side SN2 reaction (Fig. 4a), which must be dynamically restricted (see below). A more recent, ab initio direct dynamics trajectory simulation supports the back-side SN2 mechanism, substantial vibrational excitation in the CH₃Cl reactant is required to access the front-side transition state, and even so, the front-side pathway will still be dominated by the back-side reaction.

For the thermoneutral Cl⁻ + CH₃Cl system, the transition state for the back-side SN2 reaction is actually calculated to be slightly higher in energy than the reactants at the G2(+) level of theory (ΔEᵣₓᵤₓ = +2.7 kcal mol⁻¹). This is consistent with the fact that the reaction is very slow at 300 K and accelerated by translational excitation. However, why is the translational threshold (≈11 kcal mol⁻¹) significantly greater than the overall energy barrier? As it turned out, the “simple” SN2 reaction (Eq. (12)) is far from simple. The Cl⁻ + CH₃Cl reaction is a member of methyl halide SN2 reactions that display nonstatistical kinetics and dynamics (see Sun, Song, and Hase and references therein). The short lifetime of the initial [Cl⁻⋯CH₃Cl] complex (τ ≈ 4 ps) hampers efficient coupling of the Cl⁻⋯CH₃Cl intermolecular mode (i.e., translational energy) and CH₃Cl intramolecular mode, resulting in the failure to utilize the translational energy to surmount the central SN2 barrier. Orientational and angular-momentum constraints are also suggested to explain the inefficient translational activation observed. While statistical behaviors have been demonstrated for large SN2 systems with complex lifetimes τ > 10 ns, SN2 reactions of methyl halides are nonstatistical in many cases primarily because of the inefficient intramolecular vibrational relaxation (IVR) within the complex lifetimes of ~10 ps or shorter. Such incom-

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Fig. 3. Schematic potential energy curves for (a) proton transfer reaction of base anion B⁻ with acid HA, and (b) SN2 reaction of nucleophile X⁻ with methyl halide CH₃Y, in the gas phase. The diagram (a) represents typical proton-transfer processes involving oxo- and sulfo-acids. Zero-point energies have been omitted for simplicity.
plete energy randomization violates a basic assumption in statistical rate theories, *e.g.*, canonical, transition state theory (TST) and microcanonical, Rice–Ramsperger–Kassel–Marcus (RRKM) theory.\(^5^{2,53}\)

In discussing "nonstatistical" behaviors, it should be clarified in what region of the potential energy surface IVR is impeded and how it affects the experimental observables. Statistical rate theories assume complete randomization of energy (i.e., statistical population of phase space) all the way from the reactants to the transition state (A in Fig. 3b). Thus, nonstatistical behaviors show up as deviations in rate kinetics (i.e., kinetic isotope effects, see below) and as differential effects of reactant excitation (translation, rotation, vibration) on the reaction rates.\(^5^{4,55}\)

When energy randomization is hampered *after* the transition state all the way down to the products (C in Fig. 3b), nonstatistical dynamics are observed as anomalous *product* energy distributions,\(^5^{6,57}\) while the rate kinetics may behave normally. It has also been predicted that the $\text{S}_2\text{N}_2$ reaction of $\text{HO}^- + \text{CH}_3\text{F}$ avoids the deep potential energy minimum in the exit channel\(^5^{19}\): it is caused by inefficient IVR in the post TS region within the time scale of the product complex dissociation.

A more subtle case is inefficient translation-to-vibration (T-V) energy coupling in the TS region (B in Fig. 3b). As shown by simulations for the $\text{Cl}^- + \text{CH}_3\text{Br}$ and $\text{Cl}^- + \text{CH}_3\text{Cl}$ systems,\(^4^{8,57}\) this traps the complex in the central barrier region and allows multiple TS crossings to occur, another violation of a basic assumption in statistical rate theories.\(^5^{2,53}\) The reaction probability per ion-molecule collision will consequently decrease. Direct techniques such as transition-state spectroscopy may be used to probe the transient species trapped in this region. Recently, femtosecond time-resolved photoelectron spectroscopy has been employed\(^5^{56}\) to monitor the dissociation dynamics of an energized $\text{S}_2\text{N}_2$ complex $\text{I}^- \cdot \text{CH}_3\text{I}$. Because of the experimental constraints\(^5^{56}\) the complex cannot have sufficient energy to surmount the $\text{S}_2\text{N}_2$ barrier. In this symmetric system the $\text{S}_2\text{N}_2$ process is not distinguishable from complex dissociation, either. Nonetheless, a future breakthrough may be expected with these approaches.

### 4.2 Kinetic isotope effects in $\text{S}_2\text{N}_2$ reactions

Kinetic isotope effects (KIEs)\(^5^{56}\) have been used as a somewhat indirect but sensitive probe for the mechanism and transition state structure of $\text{S}_2\text{N}_2$ reactions. For methyl halides reacting with nucleophile anions ($\text{Nu}^-$), the secondary $\alpha$-deuteron KIE is defined as the ratio of the rate constants for unlabeled ($k_\text{u}$) to deuterated ($k_\text{d}$) $\text{S}_2\text{N}_2$ substrates ($\text{KIE} = k_\text{u}/k_\text{d}$).

$$\text{CH}_3\text{X} + \text{Nu}^- \rightarrow \text{CH}_3\text{Nu} + \text{X}^- \quad (13)$$

$$\text{CD}_3\text{X} + \text{Nu}^- \rightarrow \text{CD}_3\text{Nu} + \text{X}^- \quad (14)$$

The inverse KIE ($k_\text{u}/k_\text{d} < 1$) is generally consistent with a *tight* $\text{S}_2\text{N}_2$ transition state, for which the overall energy barrier $\Delta E^{\ast}_{\text{TS}}$ is slightly higher for the protonated reactant than for the deuterated reactant.\(^5^{52}\) A working hypothesis in quantitatively discussing KIEs is that the reaction is statistical, i.e., there is complete energy randomization all the way from the reactants up to the transition state. The assumption allows for prediction of KIEs based on conventional TST and the associated partition function relationship between the reactants and transition state. Deviations from statistical dynamics are noticed as the difference between experimental and theoretical KIEs, either in the magnitude for a specific reaction, or in the trend for a series of systematic $\text{S}_2\text{N}_2$ reactions.

To illustrate the latter, Poirier *et al.*\(^\text{60}\) devised a transition-state "looseness" parameter, $R_{\text{TS}}$ (Fig. 5a). For a given $\text{S}_2\text{N}_2$ substrate reacting with different nucleophiles, they demonstrated computationally that KIE values are greater for nucleophiles that have larger $R_{\text{TS}}$. As $R_{\text{TS}}$ increases and the transition state becomes looser, more "space" becomes available around the C–H(D) bonds at the transition state. Thus, at the transition state, the out-of-plane C–H bending frequencies are lowered more dramatically than are the corresponding C–D frequencies, leading to an increase in KIEs from inverse toward normal ($k_\text{u}/k_\text{d} > 1$). This effect is contrasted by that of methyl C–H stretching frequencies, which are the major contributor to the overall inverse KIE but are nearly unchanged across different nucleophiles.\(^\text{60}\)

Under well-defined thermal energy conditions with the SIFT technique, we determined $\alpha$-deuteron KIEs\(^\text{51}\) for $\text{S}_2\text{N}_2$ reactions of methyl halides CH$_3$Cl, CH$_3$Br, and CH$_3$I and compared them with theoretical predictions (Kato *et al.*\(^\text{61}\) and references cited therein). Experimental KIEs for methyl chloride are found to increase from inverse toward normal as $R_{\text{TS}}$ increases (Fig. 6), in excellent agreement with TST predictions in both the trend and magnitude. This suggests that $\text{S}_2\text{N}_2$ reactions of CH$_3$Cl proceed statistically. There is a caveat, however, that consistent KIEs do not necessarily warrant

![Fig. 5](image-url)
Interestingly, computed partition functions for CH$_3$I is independent of the reactant internal energy. A calculation shows a nonstatistical behavior such that the rate
energies and by the opposite magnitudes of computational KIE values for individual KIE value and the looseness parameter. Although still predict a normal, positive correlation between the
cal. Other kinetics observations also suggest incom-
dynamics of the heavier methyl halides is nonstatisti-
cal. Experimental KIE values for CH$_3$Br and CH$_3$I
also suggest by polyatomic nucleophiles. However, the observed discrepancy in the KIE trend cannot be ex-
plained by these adjustments.
SIFT studies have been extended to microsolvated SN$_2$ reactions, which have attracted considerable attention recently. Reactions of F$^-$ (H$_2$O) + CH$_3$X (X
Cl, Br, I) and their deuterated analogs display sub-
stantial inverse KIEs by deuteration of the water sol-
vent as well as of the methyl group in methyl halides.
For unlabeled reactants, the reaction proceeds via two
channels (Eqs. (15) and (16)).

$$F^-(H_2O) + CH_3X \rightarrow X^- + CH,F + H_2O \quad (15)$$

$$\rightarrow X^- (H_2O) + CH,F \quad (16)$$

The measured KIE values are in excellent agreement
with TST predictions, which have been made for F$^-
(H_2O) + CH_3Cl.$ This may not be surprising since IRV
is expected to be very efficient for these large systems.
The product branching, however, shows intriguing dy-
namical effects. For this reaction, the measured isotope e-
ffects are consistent with TST, a factor analysis dissecting the overall KIEs
provides useful clues to the structure and molecular
interaction at the SN$_2$ transition state. The tran-
sition-state structure for the F$^-(H_2O) + CH_3F
reaction is schematically shown in Fig. 5c, in comparison to the
bare ion system F$^- + CH_3F$ (Fig. 5b). In addition to the inverse methyl halide KIE as explained by a mech-
anism similar to the bare ion case, the strong inverse KIE upon deuteration of the water solvent is due pri-
marily to the stretching vibration between the oxygen atom and the hydrogen atom H(2) that is hydrogen bonded to fluorine (Fig. 5c). The O-H(2) stretching frequency increases at the transition state compared to
that in free F$^-(H_2O),$ giving rise to the strong inverse isotope effect.

For SN$_2$ reactions of heavier methyl halides, ob-
served $\alpha$-deuterium KIEs are distinctly anomalous. Experimental KIE values for CH$_3$Br and CH$_3$I decrease (i.e., become more inverse) as $R_{TS}$ increases (Fig. 6). Interestingly, computed partition functions for CH$_3$I still predict a normal, positive correlation between the KIE value and the looseness parameter. Although
magnitudes of computational KIE values for individual reactions may vary depending on the theoretical
methods employed, the opposite trend between experi-
ments and statistical theory clearly indicates that SN$_2$
dynamics of the heavier methyl halides is nonstatisti-
cal. Other kinetics observations also suggest incom-
plete IVR for these systems.

The transition-state looseness model has been modified by incorporating steric effects in more refined manners, e.g., considera-
tion of atomic radii for the nucleophile and leaving
groups and three dimensional “crowdedness” im-
posed by polyatomic nucleophiles. However, the ob-
served discrepancy in the KIE trend cannot be ex-
plained by these adjustments.
observed.\textsuperscript{71} We speculate that a similar mechanism is operative for the KIE upon the deuteration of H(3) in methanol.\textsuperscript{74}

### 4.3 Hydrogen/deuterium exchange and scrambling

In highly exothermic ion-molecule reactions, lifetime of the intermediate complex can be substantially short that rapid complex dissociation may effectively compete with other chemical processes that are to occur therein. The reaction of CH\textsubscript{3}\textsuperscript{+} with CH\textsubscript{4} is exothermic by nearly 30 kcal mol\textsuperscript{-1} and produces the ethyl cation and H\textsubscript{2} via the protonated ethane intermediate complex [C\textsubscript{2}H\textsubscript{7}\textsuperscript{+}]. Hiraoka and Kebarle\textsuperscript{75} found evidence for the existence of two structures for the C\textsubscript{2}H\textsubscript{7}\textsuperscript{+} intermediate, both classical and non-classical. The SIFT technique was used to probe the hydrogen/deuterium (H/D) scrambling process during this reaction,\textsuperscript{76} by starting from differently labeled reactants, e.g., CD\textsubscript{3}\textsuperscript{+} and CH\textsubscript{4} (Scheme 1), CHD\textsubscript{3}\textsuperscript{+} and CH\textsubscript{3}D, and CH\textsubscript{3}\textsuperscript{+} and CHD\textsubscript{3} and observing variations in the losses of H\textsubscript{2}, HD, and D\textsubscript{2}. Isotope scrambling occurs via interconversion between the non-classical 1 and classical 2 structures. The reaction is all the way downhill relative to the reactants, since the relative energies of those intermediates as well as of the connecting transition states are significantly lower than the reactant energy by at least 20 kcal mol\textsuperscript{-1}.\textsuperscript{75, 77} If the complex is sufficiently long-lived, equilibrium between the two structures is established and the losses of hydrogen will be statistical and the same, regardless of the different deuteration in the reactants. The experiments indicate that the scrambling is incomplete and there are subtle differences among the individual reactant systems. This is caused by rapid dissociation of the complex; a modeling shows that approximately seven H/D scramblings occur before the complex dissociates to products.\textsuperscript{76}

Hydrogen/deuterium exchange of ions with neutral reagents is a thermoneutral process besides the small isotope effects. In anion H/D exchange, the reaction is considered to proceed via a key intermediate complex 3, in which all hydrogen atoms are identical (Scheme 2).\textsuperscript{78} Abstraction of a proton by B\textsuperscript{-} followed by complex dissociation results in one H/D exchange. By repeating collisions with BD, all hydrogen atoms in AH\textsubscript{n}\textsuperscript{-} may exchange. We observed anomalous H/D exchange for mono- and difluorophenyl anions reacting with D\textsubscript{2}O and CH\textsubscript{3}OD.\textsuperscript{79} The observed behavior reveals the dynamical motion within the ion–dipole complex. For example in the p-difluorophenyl anion, all three hydrogen atoms are thermodynamically identical after intracomplex deuteron transfer to form 5 or 6, neglecting any small isotope effects (Scheme 3). Thus one would expect that the three hydrogen atoms in the anion will exchange with either reagent. In fact, CH\textsubscript{3}OD does exchange all three, but D\textsubscript{2}O exchanges only one. This indicates that DO\textsuperscript{-} is trapped on one side of the benzene ring and cannot access the other side bearing two hydrogen atoms, whereas CH\textsubscript{3}O\textsuperscript{-} can migrate freely within the complex.

The kinetically restricted exchange with D\textsubscript{2}O can be understood in terms of the calculated energy diagram.\textsuperscript{79} Ab initio computations indicate that the DO\textsuperscript{-} species can migrate along the side of the benzene ring where it is formed (5); however, it cannot migrate across the barrier caused by the fluorine atoms or by...
the π electrons on the top and bottom of the aromatic ring. For H/D exchange with methanol the CH₃O⁻ species can cross over to the other side of the ring by surmounting the π electron barrier (6). The asterisks in 6 indicate the location of the methoxide anion at the “crossover” transition state. A similar transition state exists for hydroxide but the energy is higher than the reactants so the crossover does not take place. The number of exchangeable hydrogens thus provides information about the dynamics of ion–dipole intermediate complexes. On the other hand, these results caution against a simple interpretation of ion structure based solely on the extent of H/D exchange. 7,8

4.4 Termolecular association

Mechanisms of gas phase ion-molecule termolecular association reactions can be probed by species- and site-specific deuterium of the reagents. For example, in the association reaction of methoxide and water molecules that is assisted by collisional stabilization with the helium buffer gas, how are the association rates affected by deuterium substitution of the bridging hydrogen (Eq. (18)) or the methyl group (Eq. (19)) relative to the perprotio system (Eq. (17))?  

\[
\begin{align*}
\text{CH}_3\text{O}^- + \text{H}_2\text{O} & \xrightleftharpoons[\text{k}_b]{\text{k}_a} \text{CH}_3\text{O}^- \cdot \text{H}_2\text{O} \quad (17) \\
\text{CH}_3\text{O}^- + \text{D}_2\text{O} & \xrightarrow[\text{k}_a]{\text{k}_b} \text{CH}_3\text{O}^- \cdot \text{D}_2\text{O} \quad (18) \\
\text{CD}_3\text{O}^- + \text{H}_2\text{O} & \xrightleftharpoons[\text{k}_b]{\text{k}_a} \text{CD}_3\text{O}^- \cdot \text{H}_2\text{O} \quad (19)
\end{align*}
\]

We systematically studied 14 association reactions of hydroxide–water, methoxide–water, methoxide–methanol, and ethoxide–ethanol and their deuterated analogs. 8,9 Apparent bimolecular rates are measured as a function of helium pressure to derive the rate constants for termolecular association. Association rates are significantly enhanced by deuterium of the alkyl groups (by factors of 1.4–3.0) whereas deuteration of the bridging hydrogen does not affect the association rates.

The observed isotope effects can be understood in terms of the structure and vibrational modes for the association complex. 8,9 For example, in association of methoxide with water, the intermediate complex 7 formed from CH₃O⁻ and H₂O (rate constant kₚ) is likely to have a hydrogen-bonded structure (Scheme 4). 8,9 The energized complex either back-dissociates to the reactants (rate constant kₜ) or is collisionally stabilized to the adduct CH₃O⁻ • H₂O (rate constant kₛ, neglecting slow radiative stabilization). If the increase in the complex density of states overwhelms the increase in the reactant density of states upon deuteration substitution, the complex lifetime increases (i.e., kₛ decreases) and more efficient adduct formation results. The observed lifetime increase upon deuteration of the methyl group is dominated by the increase of the density of states provided by the low frequency modes (<200 cm⁻¹), which are due to methyl torsions and internal rotation. 8,9 On the other hand, deuteration substitution on the bridging position does not affect the complex lifetime because the high-frequency O–H and O–D vibrations do not substantially increase the density of states. Detailed analysis of the magnitude and pressure dependence of the association rates has also provided insights into the potential energy surface (single- versus double-well) and dynamical bottleneck for ion-molecule association. 8,9

5. Fundamental Ion Chemistry of Biological Interest

5.1 S₈₂ reaction at nitrogen center

Nucleophilic substitution at a nitrogen center is a convenient synthetic path to the formation of C–N bonds. 8,9 The reactions have also been recognized to play a significant role in carcinogenesis. 8,9 The existence of a classical S₈₂ transition state at a nitrogen substrate has been inferred using double labeling experiments in the liquid phase and theoretically supported for a simple system X⁻ + H₂NY → XH₂ + Y⁻. 8,9 However, no experimental data were available for the gas-phase kinetics of S₈₂ reactions at nitrogen centers.

We first investigated the S₈₂ reactions of chloramine (NH₂Cl) with several anionic nucleophiles (Fig. 4c). 9,10 The unstable chloramine substrate was synthesized in situ and allowed to react with several nucleophiles in the flow tube (Nu⁻ = HO⁻, CH₃O⁻, C₆H₅O⁻, C₃H₇O⁻, F⁻, Cl⁻, CH₃CH₂O⁻, CF₃CH₂O⁻, HS⁻, and Cl⁻), in the decreasing order of gas phase basicity). The most stable ion-molecule complex has the structure 8, which directly correlates with the proton–transfer products NuH⁻ + NHCl⁻. In fact, proton transfer was observed for CH₃H₂O⁻ and more basic anions, leading to the assignment of the deprotonation enthalpy of chloramine as 37.4 ± 3.0 kcal mol⁻¹. Nucleophilic substitution was observed for all reactant anions via the transition state 9 producing NuNH₂ + Cl⁻. The S₈₂ reactions are faster than the corresponding S₈₂ reactions of methyl chloride; the chloramine reactions take place at nearly every collision when the reaction is exothermic. Most dramatically, the symmetric S₈₂ reaction of 37Cl⁻ + H₂N₃5Cl is found to be more than two orders of magnitude faster than the carbon-center analog of 37Cl⁻ + CH₃Cl (Eq. (12)). There has been controversy on the S₈₂ barrier height ΔF°ₓₓ at nitrogen relative to that at carbon. The significantly enhanced S₈₂ reactivity of

![Scheme 4. Termolecular association of methoxide and water assisted by collision with He.](image)
NH₂Cl supports a high-level ab initio theoretical prediction\(^{93}\) that overall barrier heights for identity Sn2 reactions X⁻ + NH₂X are negative and lower than for carbon-center Sn2 reactions X⁻ + CH₂X.

### 5.2 ECO₂ elimination reactions

For larger reactants such as propyl halides reacting with nucleophile anions, elimination at the β-carbon (E2) can compete with nucleophilic substitution at the α-carbon (Eqs. (20) versus (21)).

\[
\begin{align*}
\text{S}_{n2}^2 & \rightarrow \text{CH}_3\text{CH}_2\text{CH}_2\text{Nu} + X^- \quad (20) \\
\text{E}_{2} & \rightarrow \text{CH}_3\text{CH} = \text{CH}_2 + \text{NuH} + X^- \quad (21)
\end{align*}
\]

These two pathways yield the same halide anion X⁻ and hence cannot be distinguished directly by product ion analysis. Relative importance of Sn2 versus E2 channels was deduced from systematic measurements of rate constants for α-branched alkyl halides.\(^{92}\) It was concluded that the Sn2 channel dominates with sulfur nucleophiles, whereas the reaction proceeds mainly via the E2 channel with oxo anions.\(^{92}\) Deuterium KIEs have also been measured using the SIFT technique to differentiate the two mechanisms.\(^{93}\) More recently an ingenious method has been devised that uses dianions as attacking nucleophiles, thereby leaving a negative charge on the (otherwise) neutral product and allowing for its mass spectrometric detection.

We recently used the SIFT technique to study a different type of elimination reactions ("ECO₂"), in which alkyl hydroperoxides react with gas phase anions to form C=O double bonds in the product.\(^{95}\) In the rapid reaction of methyl hydroperoxide with fluoride (49%) efficiency with respect to the collision rate, Scheme 6, formation of hydroxide via complex 11 is the major pathway (≈85%) whereas slightly endothermic proton transfer via complex 10 is minor (≈10%). Nucleophilic substitution at the methyl group to form HO⁻ is endothermic and is not observed. The ECO₂ reaction proceeds via anionic attack on an α-hydrogen followed by concerted elimination of formaldehyde and HO⁻. The mechanism is substantiated by observing HO⁻ as the major product from deuterated reactant CD₃OOH. The importance of the α-hydrogen is demonstrated by the formation of hydroxide from CH₃OOH and CH₃CH₂OOH and not from (CH₃)₃COOH.\(^{95}\)

Energy and structure calculations support the mechanism and high efficiency for the ECO₂ reaction (Fig. 7).

The CH₃OOH + F⁻ reaction proceeds via initial formation of complex 11 with the fluoride associated with one of the α-hydrogens, then surmounting the key transition state TS1 which is nearly 15 kcal mol⁻¹ lower in energy than the reactants. Consistent with the concerted nature of the reaction, the ECO₂ transition state is an antiperiplanar structure with C–H and O–O bonds elongated by about 0.2 Å relative to free CH₃OOH while the C–O bond is contracting.\(^{95}\) The reverse reaction, CH₃OOH + HF, is found to proceed at nearly every collision and yields HO⁻ as the major primary reaction product (≈80%), despite the possibility for exothermic proton transfer. This is also totally consistent with the computation that the reactants can follow the same pathway as above after the initial complex 10 has surmounted the transition state TS2 (Fig. 7).

The gas phase ion study has unambiguously revealed the mechanism and importance of base-mediated decomposition of alkyl hydroperoxides, which has been controversial in solution.\(^{96,97}\) These results also bear biological implications associated with degradation of lipid hydroperoxides producing genotoxic aldehydes and ketones.\(^{98,99}\) In contrast to fluoride, a weaker base such as NC⁻ cannot induce the ECO₂ reaction with CH₃OOH.\(^{95}\) However, our preliminary calculations suggest that as the acidity of the α-hydrogen increases, weak bases such as cyanide and chloride may facilitate ECO₂ decomposition. Those peroxides include allyl hydroperoxide (CH₂=CHCH₂OOH) and conjugated diene hydroperoxides, good models for endogenous biological lipid hydroperoxides.

---
We also studied collision-induced dissociation of the \([\text{M}^-/\text{c8142H}]\) anions of alkyl hydroperoxides, \(\text{CH}_3\text{OO}^\cdot/\text{c8140}, \text{CH}_3\text{CH}_2\text{OO}^\cdot, \text{and (CH}_3)_3\text{COO}^\cdot/\text{c8140}\). Facile decomposition of ethyl- and \(t\)-butyl peroxide anions to produce \(\text{HOO}^\cdot/\text{c8140}\) was observed. Dehydration of peroxide anions was also observed but it is a minor process occurring only at high collision energies.100) This is in contrast to the CID of the \([\text{M}^-/\text{c8142H}]\) anions of lipid hydroperoxides, in which loss of water is the key initial step in the decomposition.101) Combining the results of deuterium labeling,101) we propose that decomposition of lipid peroxide anions actually proceeds via an intramolecular \(\text{ECO}_2\) mechanism.100) Incorporation of a powerful electrospray ionization (ESI) source in the flow tube 7) is expected to allow studies of large peroxides with the SIFT setup. We are currently developing those ESI sources.

6. Structure and Thermochemistry of Energetic and Unstable Anions

6.1 Peroxide anions

Hydroperoxyl (HOO) and alkyl peroxy (ROO) radicals are key intermediates in combustion. HOO radicals also play a vital role in oxidation of volatile organic compounds in atmosphere. ROO radicals are usually the first oxidation products of hydrocarbons in troposphere, resulting in ozone formation through photochemical cycles. Heats of formation of the peroxy radicals are crucial in modeling these processes, and yet the experimental values have been controversial and largely unestablished (these aspects reviewed and discussed in Blanksby et al.102) and Ramond et al.103).

To accurately determine the peroxy radical heats of formation, we used the acidity/electron affinity thermochemical cycle104) which combines the gas phase acidity measurements with SIFT and electron affinity measurements with negative ion photoelectron spectroscopy (NIPES).102,103) The gas phase acidity of alkyl hydroperoxide, \(\Delta_{\text{acid}}G_{298}(\text{ROOH})\), is determined at room temperature by measuring the forward \((k_f)\) and reverse \((k_r)\) rate constants for the proton transfer equilibrium between the peroxide anion and a reference acid (HA) (Eq. (22)).

\[
\text{ROO}^\cdot/\text{c8142H} + \text{HA} \overset{k_f}{\underset{k_r}{\rightleftharpoons}} \text{ROOH} + \text{A}^\cdot
\]

The equilibrium constant \(K_{\text{equil}} (= k_f/k_r)\) is connected to \(\Delta_{\text{acid}}G_{298}(\text{ROOH})\) through a known acidity value for HA (Eq. (23)). The enthalpy of deprotonation, \(\Delta_{\text{acid}}H_{298}(\text{ROOH})\), is then derived using computational entropies.

\[
\Delta_{\text{rxn}}G_{298} = \Delta_{\text{acid}}G_{298}(\text{HA}) - \Delta_{\text{acid}}G_{298}(\text{ROO}^-/\text{c8142H}) = -RT \ln K_{\text{equil}}
\]

The electron affinity of the peroxy radical, \(\text{EA}(\text{ROO})\), is measured by NIPES of the ROO\(^-\) anion. The measured deprotonation enthalpy and electron affinity, along with the ionization energy of hydrogen \(\text{IE}(\text{H})\), afford the bond dissociation enthalpy, \(\text{DH}_{298}\) (ROO\(^-\)–H) (Eq. (24)). The thermal correction term arises from the fact that \(\text{EA}(\text{ROO})\) and \(\text{IE}(\text{H})\) are measured at 0 K. The correction is usually very small (< 0.3 kcal mol\(^{-1}\)) and can be neglected,104) or computed from the relevant heat capacities. The 0 K bond dissociation energy, \(\text{DH}_0\) (ROO\(^-\)–H), is also calculated from the 298 K bond enthalpy following similar thermal corrections.

\[
\text{DH}_{298}(\text{ROO}^-/\text{c8142H}) = \Delta_{\text{acid}}H_{298}(\text{ROO}^-/\text{c8142H}) - \text{IE}(\text{H}) + \text{EA}(\text{ROO}) + \text{thermal correction}
\]

Finally, the heat of formation for ROO is obtained from the bond enthalpy and known heats of formation for hydrogen and ROOH (Eq. (25)), without the need for relying on thermochemistry of other intermediate species.
Table 1. Experimental Deprotonation Enthalpies, Electron Affinities, Bond Dissociation Energies, and Heats of Formation for Alkyl Peroxy Radical, ROO, and Hydroperoxides, ROOH

<table>
<thead>
<tr>
<th>R</th>
<th>$\Delta_{\text{expt}} H_{298}(\text{ROOH})$ (kcal mol$^{-1}$)</th>
<th>$EA(\text{ROO})$ (eV)</th>
<th>$DH(\text{ROO-H})$ (kcal mol$^{-1}$)</th>
<th>$\Delta H_{298}(\text{ROO})$ (kcal mol$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H</td>
<td>376.5 $\pm$ 0.4</td>
<td>1.078 $\pm$ 0.006</td>
<td>86.6 $\pm$ 0.5</td>
<td>3.2 $\pm$ 0.5</td>
</tr>
<tr>
<td>CH$_3$</td>
<td>374.6 $\pm$ 1.0</td>
<td>1.161 $\pm$ 0.005</td>
<td>86.9 $\pm$ 1.0</td>
<td>4.8 $\pm$ 1.2</td>
</tr>
<tr>
<td>CH$_3$CH$_2$</td>
<td>371.0 $\pm$ 2.2</td>
<td>1.186 $\pm$ 0.004</td>
<td>83.9 $\pm$ 2.2</td>
<td>$-$ 6.8 $\pm$ 2.3</td>
</tr>
<tr>
<td>(CH$_3$)$_3$C</td>
<td>370.2 $\pm$ 2.1</td>
<td>1.196 $\pm$ 0.011</td>
<td>83.3 $\pm$ 2.1</td>
<td>$-$ 25.2 $\pm$ 2.5</td>
</tr>
</tbody>
</table>

$^{a}$ Ramond et al.$^{(103)}$ for $R = H$ and Blanksby et al.$^{(102)}$ for $R = \text{CH}_3$, \(\text{CH}_3\text{CH}_2\), and \((\text{CH}_3)_3\text{C}\).

$$
\Delta H_{298}(\text{ROO}^-) = DH_{298}(\text{ROO-H}) - \Delta H_{298}(\text{H}) + \Delta H_{298}(\text{ROOH})
$$

Table 1 summarizes the thermochemical parameters for ROO, CH$_3$OO, CH$_3$CH$_2$OO, and \((\text{CH}_3)_3\text{COO}\) radicals determined to within an accuracy of a few kcal mol$^{-1}$ or better. The uncertainties in bond dissociation energies and heats of formation arise primarily from those in the measured heats of deprotonation. It is noted that the SIFT measurement of the relative acidity ($\Delta_{\text{expt}} G_{298}$) is very accurate to within a factor of kcal mol$^{-1}$, and the uncertainty in $\Delta_{\text{expt}} H_{298}(\text{ROOH})$ is almost entirely due to the error bars in the reported gas phase acidities for the reference acids (Eq. (23)). The measured heats of formation for the alkyl peroxy radicals$^{(102)}$ also agree with several previous reports, settling the controversy in the thermochemistry of these reactive species.

### 6.2 Nitrogen-containing anions

Compounds containing many nitrogen atoms or nitrogen substituents (e.g., $\text{NO}_2$ and $\text{N}_2$ groups) are energetic species. Anions and radicals of those species are also considered important intermediates in combustion. The SIFT technique has been employed to study, through an anionic approach, the structure and thermochemistry of diazocarbene (CNN) and the diazomethyl radical (HCCN)$^{(106)}$ small nitroalkanes,$^{(107)}$ and a series of five-membered azole compounds.$^{(108)}$

HCNN$^-$ and CNN$^-$ anions can be generated, for example, by HO$^-$ deprotonation and O$^-$ dehydrogenation of diazomethane, respectively (Scheme 7). Proton affinities of these anions were determined by proton-transfer equilibrium (for HCCN$^-$) and acidity bracketing (for CNN$^-$) measurements, to be 372 $\pm$ 2 and 352 $\pm$ 4 kcal mol$^{-1}$, respectively.$^{(109)}$ It is noted that only acidity bracketing is feasible with CNN$^-$ because the reverse reaction requires the HCCN radical reactant. Photodetachment of HCCN$^-$ and CNN$^-$ gives access to [HCCN] and [CNN], which are conjectured to be combustion intermediates that provide low energy paths to cleave N$_2$ to produce N atoms (Scheme 7). The nitrogen atoms are rapidly oxidized to nitric oxide, a major source driving urban air pollution. Electron affinities of HCCN and CNN were measured by NIPES of these anions. The heat of formation of diazomethane, heats of formation for the HCCN radical and CNN carbene were accurately determined. It turns out that the HCCN intermediate provides a deep well of $-$31 $\pm$ 3 kcal mol$^{-1}$ relative to CH + N$_2$ reactants in

$$
\text{HCNN} + \text{CH}_3 \rightarrow \text{HCNN}^- + \text{CH}_3^+ \rightarrow \text{HCNN}^- + \text{H} + \text{CH}_3^+ \rightarrow \text{HCNN}^- + \text{H} + \text{N}_2
$$

Scheme 7. Formation of HCCN and CNN intermediates using gas phase ion schemes (solid arrows), along with formation and subsequent dissociation in combustion (broken arrows).

![Scheme 8](image1)

Scheme 8. Competitive proton transfer and E2 elimination reactions of nitroalkane, and the chemical reactivity of the deprotonated nitroalkane anion.
the spin-forbidden CH(2II) + N2 → HCN+N(4S) reaction, which is slightly endothermic, $\Delta_{Rxn}H^0=1.6\pm0.7$ kcal mol$^{-1}$. The detailed mechanism of how the doublet/quartet crossing occurs within the deep potential well remains an intriguing question.

Nitro compounds are prototypical propellants and explosives, as exemplified by high-performance explosives trinitrotoluene (TNT) and 1,3,5-trinitro-1,3,5-triazacyclohexane (RDX). Apart from acidity determinations, reactivity studies of nitroalkanes and their anions are relatively few in the gas phase. We are especially interested in the stability and reactivity of deprotonated nitroalkanes (aci-anions), which have been considered a key intermediate in the condensed phase, base-sensitized detonation of nitroalkanes.

Small nitroalkanes (nitromethane, nitroethane, 2-nitropropane, 2-methyl-2-nitropropane, and nitrocyclopropane) were allowed to react with gas phase bases HO$^-$, CH$_3$O$^-$, and HOO$^-$. We find the reactions to proceed very rapidly, $k = 2.5-4.3\times10^{-3}$ cm$^3$ s$^{-1}$, primarily producing [M $\rightarrow$ H]$^-$ anions via proton transfer (where an α-hydrogen is available) as well as NO$_2^-$ via the E2 pathway (where a β-hydrogen is available).

S$_n$2 formation of NO$_2^-$ is ruled out because NO$_2^-$ is not observed from CH$_3$NO$_2$. The acyclic nitroalkanes are similarly very acidic because of resonance stabilization involving the aci form for the deprotonated anions, in addition to the strong inductive effect of the nitro group.

The anomalously low acidity for nitrocyclopropane, which has the R−R bonding, is explained by the destabilizing strain present in the aci form.

Reactivity studies revealed the importance of the aci form in the [M $\rightarrow$ H]$^-$ anions of acyclic nitroalkanes. In the S$_n$2 reactions of [M $\rightarrow$ H]$^-$ anions with CH$_3$I, the efficiencies do not decrease but rather gradually increase with larger substituents R and R (Scheme 8).

This is an interesting result because the larger attacking anions have greater crowdedness around the deprotonated carbon atom and hence are expected to display a reduced S$_n$2 reactivity. The observation is rationalized if the nuclophilic attack is actually made by the oxo anion in the aci form (Scheme 8). The major products CS$_2$O$^-$ and SO$_2^-$ observed from the reactions with CS$_2$ and SO$_2$, respectively, are also consistent with the aci form being the important resonant structure. The [M $\rightarrow$ H]$^-$ anion of nitrocyclopropane displays different S$_n$2 and bimolecular reactivities than do acyclic anions, which is explained by the reduced contribution of the aci form in the cyclic anion.

Polynitrogen compounds have been of both fundamental and practical interests. In theory those species can derive from isoelectronic substitution of N atoms for CH units; however, polynitrogen compounds are considerably less stable because of the nitrogen lone pair repulsion that is introduced. Only a handful of homonuclear polynitrogen species has been discovered, e.g., azide (N$_3^-$). This, in turn, indicates that these species are promising high energy density materials (HEDM). For example, five-membered rings pentazole (HN$_5$) and the pentazolide anion (N$_5^-$) have been predicted to be metastable. In fact, the N$_5^-$ anion ($\Delta H^0_f=-62 \pm 4$ kcal mol$^{-1}$) has recently been generated in the gas phase by CID and laser ablation of appropriate precursor molecules containing N$_5$ units. N$_5^-$ has also been formed as a zinc salt in a $-40^\circ$C solution.

Despite the observation of N$_5^-$ anions, experimental characterization of N$_5^-$ and its derivatives has been extremely difficult. We have taken a different approach in which we study deprotonated anions of lower azoles in the gas phase by measuring the proton affinity (PA) and electron binding energy (EBE), then

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Fig. 8. Pentazole (HN$_5$), pentazolide anion (N$_5^-$), pentazolyl radical (N$_5^+$), and lower azolide anions (N$_1^-$N$_3$).
extrapolate the results to the pentazole family (Fig. 8). CID threshold energies and fragmentation patterns can also be examined for these lower azolide anions in order to evaluate the pentazole stability against metastable decomposition: N5→N3+N2. We have generated all azolide anions in Fig. 8. While this is a relatively new project with most of the work currently in progress, our first experiments on the pyrrolyl anion (C4H4N−)106 have determined its proton affinity and electron binding energy. The anion is very stable (EBE = 2.145 ± 0.010 eV), and strong vibronic coupling is indicated between the ground and first excited states for the detached pyrrolyl radical.106 The SIFT instrument is also being used for CID studies of these azolide anions. Preliminary experiments indicate an increasing trend for EBE and decreasing trends for ΔSIFT/ΔS and the CID threshold as the heterocyclic ring incorporates more nitrogen atoms.

6.3 Cyclooctatetraene and cyclopentanone

Conformational changes in 1,3,5,7-cyclooctatetraene (COT, C8H8) and its substituted derivatives have been a subject of considerable research interest for several decades; tub-like COT [D2d] undergoes ring inversion via a planar D8h structure.119,120 Relatively little is known, however, about the thermochemical properties of COT and, in particular, its radical and anionic derivatives. We generated anionic derivatives of COT using electron impact, HO2− and electron binding energy. The anion is very stable (EBE = 2.145 ± 0.010 eV), and strong vibronic coupling is indicated between the ground and first excited states for the detached pyrrolyl radical.106 The SIFT instrument is also being used for CID studies of these azolide anions. Preliminary experiments indicate an increasing trend for EBE and decreasing trends for ΔSIFT/ΔS and the CID threshold as the heterocyclic ring incorporates more nitrogen atoms.

The radical anion of cyclooctatetraene [COT−(1)] is one of the first open-shell annulenes observed experimentally. The gas phase proton affinity for COT−(1), along with those for C8H7−(2a) and C8H6−(3), was also measured. Combining the values of proton affinities with electron binding energies measured using photoelectron spectroscopy, we determined the C−H bond dissociation energies of COT and its derivatives, C8H8 (n = 7−10),122,126 Photoelectron spectroscopy and molecular orbital calculations revealed that the C8H7−(2a) anion is a novel structure incorporating a π-electronic system and a bent allenic moiety within the eight-membered ring.122

The large C8 ring systems undergo unexpected transformations upon collisional excitation. When SIFT-injected at high collision energies, COT−(1) and C8H6−(3) radical anions appear to detach an electron for signal loss. Remarkably, the C8H7−(2a) anion undergoes a collision-induced isomerization to C8H−(2b) via transannular bridging (Scheme 9),126 instead of usual ring opening or fragmentation. Measured chemical reactivity and molecular orbital calculations characterized C8H−(2b) as a fairly stable bicyclic species (ΔHf = 33 kcal mol−1) with EBE and PA very similar to those for cyclopentadienide anion. Duplication of this transannular bond formation process in solution would provide entry into bicyclic ring systems that pose significant synthetic challenge.

At even higher SIFT injection energies the C8H7−(2b) product is replaced by a C8H6−(γ) species, apparently following loss of a hydrogen atom. This species exhibits distinctly different chemical reactivities from the isomeric C8H6−(3). If it retains the skeletal structure of C8H7−(2b), this species may be the radical anion of pentalene, a prototypical Hückel antiaromatic molecule which eluded chemists for several decades.127 With appropriate tuning of the injection energy, the

![Scheme 9. Cyclooctatetraene, its anionic derivatives, and their collision-induced transformations.](image)
C₈H₇⁻(29) anion is completely converted to C₈H₆O⁻(28) with minimal fragmentation to C₈H₆⁻(γ). The C₈H₇⁻(28) anion can then be subjected to CID under a single collision condition with the triple quadrupole (Fig. 1). Interestingly, formation of C₈H₆O⁻(γ) has been very difficult to observe under experimental conditions where similar m/z ions are easily being detected. This result may suggest that C₈H₆⁻(γ) can only be stabilized in the high-pressure collisional environment in the flow tube. Pressure-dependent product branching has been observed by a separate group,120.

As illustrated above, the atomic oxygen radical anion (O⁻) reacts with many organic compounds, formally abstracting H₂ and producing a variety of carbene and distonic radical anions129,130 as well as anions with bent triple bonds.131 We have shown that O⁻ reacts with cyclopentanone to form a mixture of three C₅H₆O⁻ isomers (Scheme 10).132 The major product cyclopentanone-2,5-diyl radical anion (16) is thermodynamically most stable. Photodetachment of this species will determine the singlet-triplet splitting of the 2,5-diyl radical, an ethano-bridged analog of oxyallyl which has been postulated as an intermediate in many organic reactions. NIPES experiments are under way in the laboratory of Prof. W. C. Lineberger.

7. New Directions

7.1 Reactions of ions with organic radicals

Thousands of ion reactions have been studied with stable neutrals and with some atomic radicals in the gas phase.140 In contrast, reactions of ions with polyatomic organic radicals have rarely been studied because of the inherent experimental difficulties. The fundamental chemistry of ion–polyatomic radical reactions, where the charge and radical sites are split between the collision partners, is thus largely unexplored. Ion–organic radical reactions are expected to be important in several environments, including in combustion processes where both positive and negative ions have been identified133 at densities of 10⁸–10¹⁵ cm⁻³. In the interstellar medium more than a hundred organic species have been detected, among them about one-third are radicals or carbones and several are ions.134 Under very low temperature conditions in space, barrierless ion–radical association and reactions may play a significant role for the formation and evolution of those interstellar species.

Investigations of ion–organic radical reactions are challenging primarily because of the difficulty in generating the radicals cleanly and in high densities. This contrasts with generation of atomic radicals, which can be achieved relatively easily by dissociation of their precursors in a microwave discharge or over a heated metal surface. These atomic sources have been combined with the SIFT135,136 and Fourier transform ion cyclotron resonance (FT-ICR)137 instruments to study reactions of H, N, and O atoms.

For polyatomic radicals, chemical ionization mass spectrometry (CIMS) has begun to use some ion–radical reactions to probe the atmospheric chemistry of allyl peroxyl (ROO⁻)138–141 and peroxycetyl (CH₃C(O)OO⁻) radicals.142 For example, when ROO⁻ radicals are undergoing neutral radical chemistry with NO, their concentration changes are monitored by using chemical ionization with O₂⁻, O₃⁻, or H₂O⁻(H₂O)⁻. The allyl peroxyl radicals are prepared by addition of O₂ to the corresponding allyl radicals, which have been generated from suitable precursor molecules by pyrolysis, low-power radio-frequency discharge, or atom reactions.138–142

Besides those CIMS applications, a versatile method for polyatomic radical generation is expected to open a fertile research field of ion–radical chemistry. We have employed a new approach that combines a supersonic nozzle with the SIFT instrument.143,144 Thermolysis of a variety of organic compounds in a pulsed heated nozzle (300–1800 K) produces clean and intense beams of organic radicals or diradicals.145 The high temperature and short residence time in the nozzle (=30 μs) ensures clean pyrolysis of the precursor molecules while retaining high conversion efficiencies (60–90%). The produced radicals are pulsed (~1.3 ms in duration, 20–40 Hz) into the high-pressure SIFT flow tube, where both the ions and radicals are quickly thermally equilibrated before the reaction initiates (~300 K). The ions and radicals are then contained for a moderately long reaction time of about 10 ms. While the ion–radical reaction products can easily be detected and quantified, depletion of the reactant ion is too small to be measured directly under the experimental conditions. However, a simple treatment of pulsed kinetics allows evaluation of the reaction rate constants based on the product yields.144,145

A simple radical allyl (C₃H₅) and a diradical ortho-benzene (α-C₆H₄) have been generated by pyrolysis of allyl iodide (C₃H₅I + Δ → C₃H₅I⁻) and benzoyl chloride (C₆H₅COCl + Δ → α-C₆H₄+CO+Cl⁻), respectively, where Δ represents heat. The strained triple bond in α-benzene carries a triplet diradical character, which renders the bond highly reactive. These radicals encounter hydronium (H₃O⁺) or hydroxide ions (HO⁻) in the flow tube.146,147 The allyl radical and α-benzene react with H₂O⁺ via exothermic proton transfer (Eqs. (26) and (28)) at nearly 100% efficiencies relative to the ion-radical collision rates. No charged products were observed from allyl + HO⁻ within detection limits (Eq. (27)). The ion–radical encounter might be essentially nonreactive; the heat of deprotonation for the allyl radical is unknown. Alternatively, this finding could result from associative/reactive detachment pathways...
(e.g., \( \text{CH}_2=\text{CHCH}_2\bullet +\text{HO}^- \rightarrow \text{CH}_2=\text{CHCH}_2\text{OH} + e^- \)) which are highly exothermic. The reaction is then analogous to associative detachment of anions with atomic radicals, which is commonly very rapid.\(^{40, 136, 150}\)

\[
\text{CH}_2=\text{CH}^- + \text{H}_2\text{O}^+ \rightarrow \text{C}_2\text{H}_5^+ + \text{H}_2\text{O} \quad (26)
\]

\[
\text{CH}_2=\text{CH}^- + \text{H}_2\text{O}^- \rightarrow \text{no product ions} \quad (27)
\]

The reaction of \( \sigma \)-benzyne with \( \text{HO}^- \) shows an intriguing result. Although abstraction of the most acidic proton is exothermic by more than 10 kcal mol\(^{-1}\), the observed product \( \text{C}_6\text{H}_5^- \) (Eq. (29)) accounts for only about 15% of the collision. This is an unexpected result because exothermic proton abstraction is usually a barrierless process (Fig. 3a) that takes place at every collision. The observation may be rationalized by competing associative detachment; the \( \text{HO}^- \) ion could add to the triple bond to form \( \sigma \)-hydroxy phenyl anion \( (\sigma-\text{C}_3\text{H}_4\text{O}^-) \), which could subsequently lose the electron because of the large reaction exothermicity far exceeding the binding energy.\(^{147, 148}\) Thus, this is a novel fundamental reaction channel for \( \text{HO}^- \) ions that is facilitated by the high energy content of \( \sigma \)-benzyne. Future experiments include trapping of detached electrons with electron scavengers such as SF\(_6\). Other radicals methyl and phenyl as well as diradical cyclopropenylidene \( (\sigma-\text{C}_2\text{H}_4) \) are among the reactants that can be generated and studied using the SIFT-supersonic pyrolysis technique.

### 7.2 Chemical ionization mass spectrometry

Compared to other ionization methods such as electron impact or hollow cathode discharge, chemical ionization is a soft ionization that bestows charge to neutral molecules with minimal fragmentation.\(^{151}\) Chemical ionization mass spectrometry, which conveniently uses proton transfer/abstraction reactions of \( \text{H}_2\text{O}^+ \) or \( \text{HO}^- \) reagent ions in flow-drift tube reactors,\(^{152, 153}\) has become increasingly popular for highly sensitive, on-line detection of a variety of volatile organic compounds (VOCs). We utilized \( \text{H}_2\text{O}^+ \) proton transfer reactions to measure aldehydes from human cancer cells\(^{154-156}\) and VOCs from fermenting bacteria.\(^{157}\) We also demonstrated that negative ion chemistry with \( \text{HO}^- \) is an alternative tool to monitor plant emission of VOCs\(^{158-160}\) and to detect small, volatile nitroalkane explosives.\(^{161}\) These analytical applications and bio/medico-chemical implications, as well as ion–molecule chemistry of interstellar relevance, will be discussed elsewhere.\(^{161}\)

A new reaction scheme of CIMS interest is given here. While reactions with \( \text{H}_2\text{O}^+ \) and \( \text{HO}^- \) are facile and versatile in converting a majority of organic species to respective \([\text{M} + \text{H}]^+\) and \([\text{M} - \text{H}]^-\) product ions, identification of isobaric products is usually a difficult task. CID and \( \text{H/D} \) exchange reactions may be used to differentiate some product isomers, for example, deprotonated acetone \( \text{versus} \) propanal.\(^{157}\) Another approach to structure identification is to devise a CI scheme with chemical specificity. In analogy to acid-catalyzed, solution phase chemistry of hydrazine and its derivatives, we have shown that reactions of protonated hydrazine \( (\text{N}_2\text{H}_5^+) \) can be used to specifically detect carbonyl compounds in the gas phase.\(^{162, 163}\) \( \text{N}_2\text{H}_5^+ \) reacts with a carbonyl compound and forms a characteristic product, protonated hydrazone (Scheme 11). This reaction, while exothermic, is slow and observed only when excess neutral \( \text{N}_2\text{H}_4 \) molecules are present.\(^{163}\) Further, formation of hydrazone is dramatically enhanced at high ion kinetic energies in the drift field.\(^{162}\) Additional experiments using deuterated \( \text{N}_2\text{D}_4 \) and density functional calculations\(^{160}\) have identified the origin of the slowness of the reaction: Transition states for the bimolecular reaction (Scheme 11) present kinetic and energetic barriers that can be overcome by translational and/or chemical activation. The energies for the strained transition states are likely to be lowered in the presence of a second \( \text{N}_2\text{H}_4 \) molecule within the intermediate complex.\(^{163}\) The fundamental studies have provided an important guideline for this CIMS scheme to be widely used for future applications.

### 8. Conclusions

Various aspects of fundamental gas phase ion chemistry have been discussed. The SIFT technique is a versatile tool with which to study the kinetics and mechanisms of ion reactions in a very clean, truly thermal environment. This feature has been fully utilized, for example, to reveal nonstatistical behaviors in \( \text{Sn}_2 \) kinetics and to prove otherwise controversial \( \text{Eco}_2 \) reactions of biological interest, as well as address classical questions in organic chemistry and explore new schemes for CIMS applications. By incorporating LIF and triple quadrupole detections and through a powerful coupling with NIPES experiments, the SIFT technique has continued to provide detailed information on ion structure, energetics, and dynamics. These results have stimulated theoretical interest as well. We have shown that a hyperthermal radical source can be used with the SIFT instrument for studies of ion-polynuclear radical reactions. ESI sources are also expected to be useful for loading ions of nonvolatile species in the flow tube. These technical developments will further extend the research horizon with the SIFT technique.

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References

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