

# To form or not to form a reaction complex: exploring ion–molecule reactions between C<sub>3</sub>H<sub>4</sub> isomers and Xe<sup>+</sup> and O<sub>2</sub><sup>+</sup>

C. Zagorec-Marks, <sup>\*ab</sup> G. S. Kocheril, <sup>ab</sup> O. A. Krohn, <sup>ab</sup>  
T. Kieft, <sup>ab</sup> A. Karpinska, <sup>abc</sup> T. P. Softley<sup>d</sup>  
and H. J. Lewandowski <sup>ab</sup>

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Ion–molecule reactions are an essential contributor to the chemistry of a diverse range of environments. While a great deal of work has been done to understand the fundamental mechanisms driving these reactions, there is still much more to discover. Here, we expand upon prior studies on ion–molecule reactions involving two isomers of C<sub>3</sub>H<sub>4</sub>, allene (H<sub>2</sub>C<sub>3</sub>H<sub>2</sub>) and propyne (H<sub>3</sub>C<sub>3</sub>H). Specifically, we probe the previously observed isomeric dependent reactivity of these molecules by reacting them with two ions with nearly identical ionization potentials, Xe<sup>+</sup> and O<sub>2</sub><sup>+</sup>. Our goal is to determine if the isomer-dependent reaction mechanisms previously observed are universal for C<sub>3</sub>H<sub>4</sub> or if they depend on the ion character as well. Through the combination of experimental measurements and theoretical calculations, we found that both isomeric structure and identity of the ion contribute to the propensity of a reaction complex forming or for only long-range charge transfer to occur.

## 1 Introduction

Cold ion–molecule reactions provide a unique view of the interactions found in many diverse chemical environments, such as the Earth's upper atmosphere and interstellar gas clouds. The long-range interaction between the charged ion and the neutral molecule results in large scattering cross-sections for inelastic or reactive processes.<sup>1</sup> This makes ion–molecule reactions relevant to a variety of environments, creating significant interest in predicting their fundamental behavior.<sup>2</sup> At the single molecule level, the outcomes or products of these ion–molecule reactions are thought to be determined by two primary mechanisms,

<sup>a</sup>Department of Physics, University of Colorado, Boulder, CO 80309, USA. E-mail: chase.zagorec-marks@colorado.edu

<sup>b</sup>JILA, National Institute of Standards and Technology and the University of Colorado, Boulder, CO 80309, USA

<sup>c</sup>Department of Chemistry and Applied Life Sciences, ETH Zürich, 8093 Zürich, Switzerland

<sup>d</sup>School of Chemistry, University of Birmingham, Edgbaston, B15 2TT, UK



direct charge transfer and complex formation.<sup>3</sup> As the name suggests, direct charge transfer largely results in a simple transfer of an electron between an ion and molecule, while complex formation results in the reactants forming an intermediate that has a sufficiently long lifetime for equilibration of vibrational energy in the complex, often resulting in the formation of new chemical bonds.<sup>3</sup> While complex formation is often attributed to the observation of distinct product distributions,<sup>4–7</sup> it is still unclear whether the formation of a complex occurs consistently for reactions of the same neutral with different ions or if the identity of the ion also plays a role. Cold ion techniques, used in conjunction with reactant-ion purification and product time-of-flight analysis, have enabled significant progress towards addressing these questions for reactions with an increasing degree of complexity.

Previously, our lab studied the relatively complex reaction between acetylene ions,  $C_2H_2^+$ , and two structural isomers of  $C_3H_4$ , allene ( $H_2C_3H_2$ ) and propyne ( $H_3C_3H$ ), and found that product formation is determined by the isomeric structure of the neutral.<sup>8,9</sup> Specifically,  $c-C_3H_3^+$  is the sole observed product in the reaction with allene, whereas in the reaction with propyne,  $c-C_3H_3^+$ ,  $C_3H_4^+$ , and  $C_5H_5^+$  are all observed in appreciable amounts originating from complex formation/dissociation. For the allene reaction with  $C_2H_2^+$ , it was found that, following charge exchange, the  $C_2H_2$  was inhibited from forming a reaction complex with the allene cation, likely due to energy localization on the cation that originated from a Jahn–Teller distortion of the allene upon ionization. In contrast, propyne had no structural changes upon ionization, thereby allowing for  $C_2H_2$  to approach and form a complex. This was an interesting finding as it showed that the isomeric structure was in fact the dominant influence on the formation of the reaction complex for a set of reactions on the same global potential energy surface.

Herein, we report an investigation where we extend these studies on reactions with  $C_3H_4$  isomers to simpler non-hydrocarbon ions as the reaction partners to see if the same isomer dependent product distributions are observed for both neutral isomers.  $O_2^+$  and  $Xe^+$  were chosen due to their similar ionization potentials, 12.0697 and 12.12987 eV respectively, and are expected to behave similarly in terms of energetics.<sup>10,11</sup> However, they are expected to follow different reaction mechanisms. While it has been suggested that  $Xe^+$  can perform charge transfer resulting from complex formation,<sup>12,13</sup> it has been shown that  $Xe^+$  rarely forms covalently bound reaction complexes,<sup>14–16</sup> so products of  $Xe^+$  reactions typically arise from direct charge transfer alone.<sup>17–21</sup> Reactions with  $O_2^+$ , on the other hand, have been shown in Selected Ion Flow Tube (SIFT) studies to both react *via* complex formation and *via* direct charge transfer.<sup>22,23</sup> Using these two ions in tandem allows us to further explore the mechanisms of reactivity between the  $C_3H_4$  isomers. Surprisingly, we find here that the ion is now the dominant influence on the reaction mechanisms, and no isomer dependent behavior is observed for either system, which is in direct contrast to the previous reactions involving  $C_2H_2^+$ . All  $Xe^+$  reactions with  $C_3H_4$  isomers undergo exclusively direct charge transfer reactions, whereas  $O_2^+$  reactions show clear complex formation that leads to different product distributions from those observed in the  $Xe^+$  reactions.



## 2 Methods

The experimental setup has been previously described elsewhere, and only a brief description will be given here for clarity.<sup>24</sup>  $\text{Ca}^+$  ions formed *via* non-resonant ionization of an effusive Ca beam using the third harmonic of a Nd:YAG laser were loaded into a linear Paul trap. The  $\text{Ca}^+$  ions were then laser cooled to secular temperatures of  $<1$  K using the frequency doubled and fundamental outputs of two Ti-Sapphire lasers ( $\sim 4$  mW at 397 nm and  $\sim 20$  mW at 866 nm). The resultant Coulomb crystal (comprised typically of  $\sim 1000$  ions) was used for sympathetically cooling co-trapped reactant ions. Fluorescence emitted by the  $\text{Ca}^+$  Coulomb crystal was imaged by an electron multiplying charge-coupled device (EMCCD) camera to qualitatively observe reactions. Reactions were quantitatively monitored *via* time-of-flight mass spectrometry by observing a reduction in reactant ion peaks and correlated growth of product ion peaks. Neutral products were not detected.

### 2.1 Reactant ion loading

After the  $\text{Ca}^+$  Coulomb crystal had formed, Xe was introduced into the chamber *via* a pulsed, skimmed supersonic expansion of Xe (2%) in a buffer gas of  $\sim 1030$  Torr He.  $\text{Xe}^+$  ions were formed in the center of the trap *via* a 2+1 resonance enhanced multi-photon ionization (REMPI) scheme using a focused beam from a pulsed dye laser at 249.6 nm (2 mJ per pulse) overlapped spatio-temporally with the Xe atomic beam.<sup>25</sup> It was expected that approximately 22% of the  $\text{Xe}^+$  ions were formed in the excited  $^2\text{P}_{1/2}$  state using this scheme. However, this excited state has a lifetime of  $\sim 49$  ms, and the  $\text{Xe}^+$  ions were held in the trap for 60 seconds prior to introduction of neutrals to ensure that all the  $\text{Xe}^+$  ions were maximally cooled before reacting; therefore, none of the reacting  $\text{Xe}^+$  ions were in the excited state.<sup>26</sup>

$\text{O}_2$  was also introduced into the chamber *via* a pulsed, skimmed supersonic expansion of  $\text{O}_2$  (12%) in a buffer gas of  $\sim 1030$  Torr He prior to the formation of the  $\text{Ca}^+$  Coulomb crystal.  $\text{O}_2^+$  ions were formed in the center of the trap *via* a 2+1 REMPI scheme using a focused 287.5 nm beam overlapped spatio-temporally with the molecular beam.<sup>27</sup> All  $\text{O}_2^+$  ions were believed to have been formed in the ground electronic state,  $\text{X}^2\Pi_g$ , and are efficiently cooled by  $\text{Ca}^+$ , thereby allowing for the reaction to commence once the Coulomb crystal had fully formed.

### 2.2 Reacting with neutrals

Reactions were initiated by admitting either neutral allene ( $\text{H}_2\text{C}_3\text{H}_2$ ) or propyne ( $\text{H}_3\text{C}_3\text{H}$ ) (14–16% in  $\text{N}_2$ ) into the chamber *via* a pulsed-leak valve to a fixed chamber pressure (with partial pressures at 300 K of  $0.7\text{--}2.5 \times 10^{-10}$  Torr and  $4 \times 10^{-10}$  Torr for  $\text{O}_2^+$  and  $\text{Xe}^+$  reactions, respectively) for fixed periods of time.<sup>28,29</sup> Pressures were measured using a Bayard–Alpert hot cathode ion gauge in the same chamber as the ion trap. Typical pressures in the chamber prior to admittance of gas were  $1.5 \times 10^{-9}$  Torr. Because Bayard–Alpert hot cathode ion gauges have reduced accuracy below  $10^{-8}$  Torr, absolute reaction rates have considerable uncertainty.<sup>30</sup> As such, we report only products and branching ratios for the reactions measured here.

To determine the branching ratios for a reaction, we measured a reaction curve by varying the length of time the neutral gas was admitted into the chamber. The



opening of the pulsed leak valve corresponded to the zero-time point. At each reaction time point, the contents of the trap were ejected into a time-of-flight mass spectrometer (TOF-MS), and a new Coulomb crystal was formed for the next measurement. The contents of the trap at each time point were measured at least 10 times, and the mean and standard error of these measurements were then plotted as a function of time to produce a reaction curve. The reaction curve was fit according to a reaction model representative of the reaction being studied. These reaction models assume that ion numbers within the trap are conserved throughout the course of a reaction.

To verify that no ions were lost from the trap during the reaction, the total ion number was determined for each time step. For the  $\text{O}_2^+$  reactions, the total ion number remained constant during a reaction. However, the case was more complicated for  $\text{Xe}^+$ . We found that not all of the trapped  $\text{Xe}^+$  ions were detected on our micro-channel plate (MCP) detector. (When analyzing the  $\text{Xe}^+$  data, all isotopes of  $\text{Xe}^+$  were summed together and reported as just the total number of  $\text{Xe}^+$  ions.) This underestimate of the number of  $\text{Xe}^+$  ions is because they have a much larger mass than  $\text{Ca}^+$ , which results in inefficient sympathetic cooling and a weaker trapping potential. These two factors combine to decrease the efficiency of detecting  $\text{Xe}^+$  by our TOF-MS, as compared to all other ions present in these reactions. Further, simulations of the  $\text{Xe}^+$  ions' extraction trajectories suggested that a fair number of these ions collided with the trap rods or grounded plate at the entrance of the TOF-MS flight tube and were thus not detected. Heavy ions have previously been observed to be more difficult to detect.<sup>31</sup> While we estimate the detection efficiency of  $\text{Xe}^+$  ions to be 50% less than lighter mass ions, none of the reaction analysis conducted relied on the measured  $\text{Xe}^+$  numbers and was determined solely on product ion numbers, which, being more similar in mass to  $\text{Ca}^+$ , are cooled more efficiently and detected with the same efficiency.

### 2.3 Theoretical

Possible global potential energy surfaces for  $\text{Xe}^+$  and  $\text{O}_2^+$  reactions were determined *via* quantum chemical calculations performed using Gaussian 16 at the unrestricted  $\omega\text{b97xd/aug-cc-pvtz}$  level of theory.<sup>32</sup> Each surface was computed starting with the reactants at infinite separation, with stationary points along the surface identified until the system reached the final observed product. Scans over bond lengths, angles, and dihedrals were used for identifying minima and potential transition states. These potential maxima were explored using optimization and frequency calculations. Transition states were identified by the presence of a vibrational mode with an imaginary frequency and through use of intrinsic reaction coordinate calculations. An effective core potential (ECP) was included for calculations directly involving Xe and  $\text{Xe}^+$  to account for relativistic effects.<sup>33</sup> Only barrierless pathways were considered because of the energetic constraints imposed by the low temperatures of the reactions.

## 3 Results

### 3.1 $\text{Xe}^+$ with $\text{C}_3\text{H}_4$ isomers

A typical TOF-MS trace for the  $\text{Xe}^+$  reactions is shown in Fig. 1. Reaction progress was monitored by the growth of the 39  $m/z$  signal ( $\text{c-C}_3\text{H}_3^+$ ) and decay of the



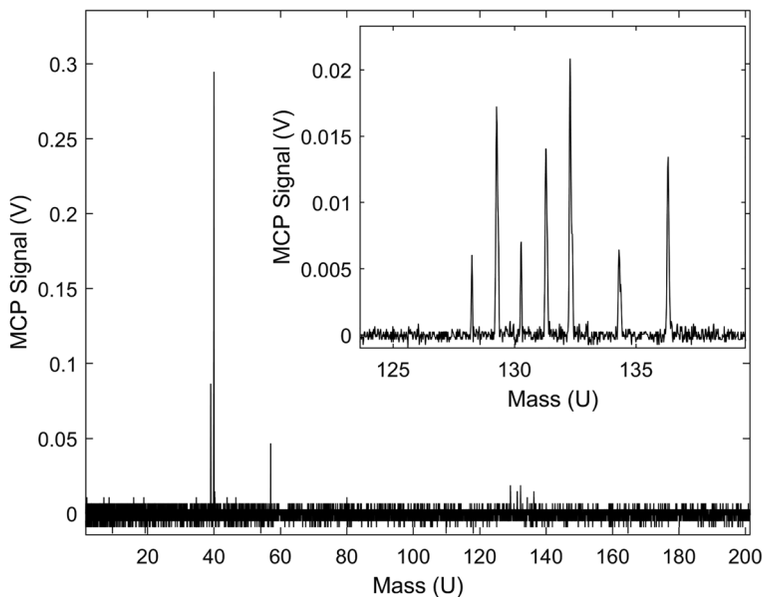


Fig. 1 Typical TOF-MS trace observed from the reaction of  $Xe^+$  with  $C_3H_4$ . Primary plot displays the trace's full signal intensity. Peaks observed in the primary plot, from left to right, correspond to 39  $m/z$  ( $c-C_3H_3^+$ , product), 40  $m/z$  ( $Ca^+$ ), 57  $m/z$  ( $CaOH^+$ ), 128–136  $m/z$  ( $Xe^+$ ). Inset displays the  $Xe^+$  signals when viewed with greater voltage resolution.

various  $Xe^+$  isotope signals, 128–136  $m/z$ . As seen in the inset of the figure, the mass resolution of the experimental setup was sufficient to resolve the seven most abundant isotopes of  $Xe$ . Secondary product growth was observed by the delayed growth of peaks at 77 and 79  $m/z$  ( $C_6H_5^+$  and  $C_6H_7^+$ ). The signals for these secondary products are not shown in the figure for clarity. The peak observed at 57  $m/z$  corresponded to  $CaOH^+$ , which was formed by a side reaction of  $Ca^+$  and trace  $H_2O$  left in the vacuum chamber. This reaction was accounted for by adding the number of ions observed in the 57  $m/z$  channel to the 40  $m/z$  ( $Ca^+$ ,  $C_3H_4^+$ , and  $C_2O^+$ ) channel during the analysis.

**3.1.1  $Xe^+ + H_2C_3H_2$  (allene).** The reaction curve for the reaction between  $Xe^+$  and allene ( $H_2C_3H_2$ ) is shown in Fig. 2. In this reaction, the dominant pathway produced 39  $m/z$  ions. This production was observed by the decay of the reactant  $Xe^+$  ion signal and simultaneous growth of the 39  $m/z$  signal. The 39  $m/z$  signal did not decrease when extending the reaction time an additional 60 seconds, indicating that, once formed, the 39  $m/z$  product was nonreactive. This product was identified as  $c-C_3H_3^+$  based on its charge-to-mass ratio and lack of further reactivity.<sup>34</sup> In addition to the  $c-C_3H_3^+$  product, there was indication of a small number of ions with 40  $m/z$  being formed and further reacting with another allene molecule to form secondary products at 77 and 79  $m/z$ ; however, this constitutes  $\sim 4\%$  of the total products of the reaction. The 40  $m/z$  product was likely  $C_3H_4^+$  based on its charge-to-mass ratio and previous studies showing that  $C_3H_4^+$  reacts with allene to form larger hydrocarbons, such as the  $C_6H_5^+$  and  $C_6H_7^+$ , which we observed here.



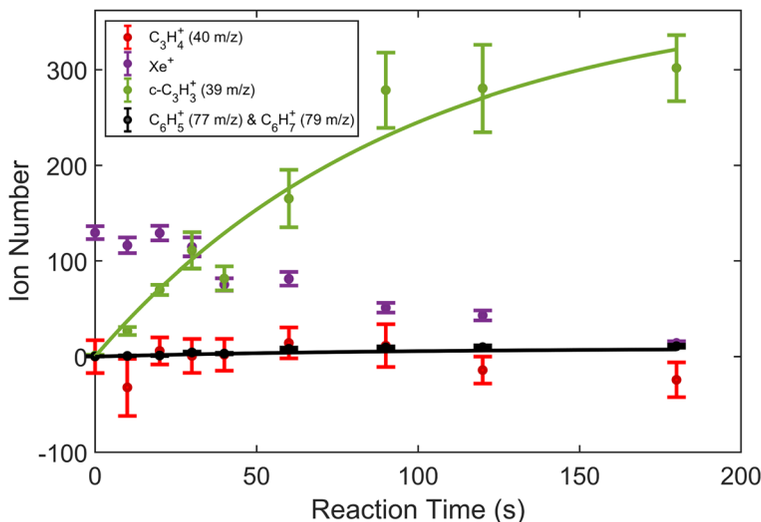


Fig. 2 Product growth in reaction of  $\text{Xe}^+$  with allene ( $\text{H}_2\text{C}_3\text{H}_2$ ) as a function of time. Data points correspond to mean and standard error of 11 measurements. Time was measured from the opening of the pulsed leak valve and subsequent inlet of allene into the chamber. Ion counts corresponding to 40  $m/z$  were adjusted by subtracting the initial number of  $\text{Ca}^+$  ions for clarity. Note that the parasitic reaction of  $\text{Ca}^+ + \text{H}_2\text{O} \rightarrow \text{CaOH}^+ + \text{OH}$  was accounted for by adding ions observed in the 57  $m/z$  channel back to the 40  $m/z$  channel. Additionally, lines plotted correspond to fits produced from eqn (1) and (2).

To extract branching ratios from these data, they were fit assuming pseudo-first order kinetics with allene in excess. Because the detection efficiency of  $\text{Xe}^+$  was lower than for the other ions in the trap, the growth of products were fit independently of the  $\text{Xe}^+$  data using the following reaction model:

$$\frac{dN_{39}}{dt} = k_1 \times (N_{\text{prod}} - N_{39} - N_{77,79}) \quad (1)$$

$$\frac{dN_{77,79}}{dt} = k_2 \times (N_{\text{prod}} - N_{39} - N_{77,79}) \quad (2)$$

where  $N_{39}$  and  $N_{77,79}$  correspond to the number of primary product (39  $m/z$ ) and secondary product (77 and 79  $m/z$ ) ions, respectively.  $N_{\text{prod}}$  corresponds to the number of product ions present at the end of the reaction, and was treated as a fit parameter. The fitting coefficients,  $k_1$  and  $k_2$ , represent rate constants. This model assumes that no lighter ions were lost from the trap and that we detected all products.

**3.1.2  $\text{Xe}^+ + \text{H}_3\text{C}_3\text{H}$  (propyne).** The reaction curve for the reaction between  $\text{Xe}^+$  and propyne ( $\text{H}_3\text{C}_3\text{H}$ ) is shown in Fig. 3. This reaction was nearly identical to that of  $\text{Xe}^+$  with allene ( $\text{H}_2\text{C}_3\text{H}_2$ ). Once again, the dominant pathway produced  $c\text{-C}_3\text{H}_3^+$ . This production was concurrent with the decay of the  $\text{Xe}^+$  signal. There was also indication of a small number of  $\text{C}_3\text{H}_4^+$  ions produced, which react with an additional propyne to form the secondary products of  $\text{C}_6\text{H}_5^+$  and  $\text{C}_6\text{H}_7^+$ ; however, this also constitutes only  $\sim 4\%$  of the total products. The fits to these data were determined from the same model used in the reaction of  $\text{Xe}^+$  with allene and with the same assumptions.



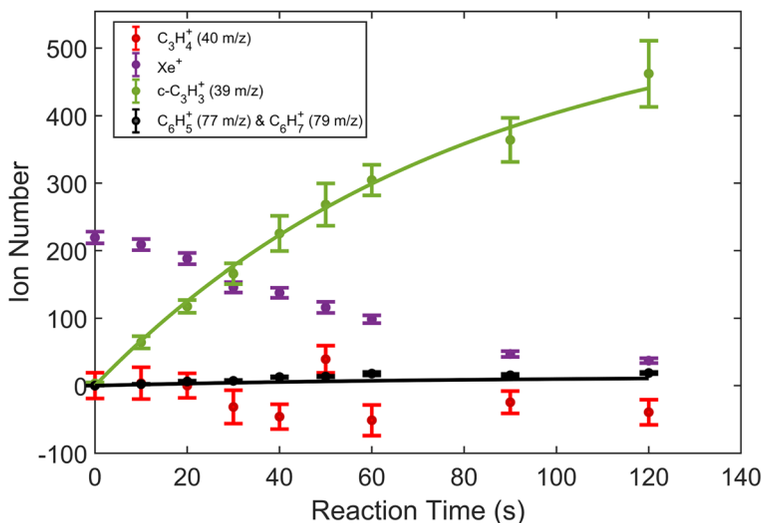


Fig. 3 Product growth in reaction of  $\text{Xe}^+$  with propyne ( $\text{H}_3\text{C}_3\text{H}$ ) as a function of time. Data points correspond to mean and standard error of 12 measurements. Time was measured from the opening of the pulsed leak valve and subsequent inlet of propyne into the chamber. Ion counts corresponding to 40  $m/z$  were adjusted by subtracting the initial number of  $\text{Ca}^+$  ions for clarity. Note that the parasitic reaction of  $\text{Ca}^+ + \text{H}_2\text{O} \rightarrow \text{CaOH}^+ + \text{OH}$  was accounted for by adding ions observed in the 57  $m/z$  channel back to the 40  $m/z$  channel. Additionally, lines plotted correspond to fits produced from eqn (1) and (2).

### 3.2 $\text{O}_2^+$ with $\text{C}_3\text{H}_4$ isomers

In addition to reactions of  $\text{Xe}^+$  with two isomers of  $\text{C}_3\text{H}_4$ , we also studied reactions of these two with  $\text{O}_2^+$  ions. In these reactions, we measured both the growth of the 39  $m/z$  signal and decay of the  $\text{O}_2^+$  signal at 32  $m/z$ . In addition to these channels, secondary product growth was observed by the delayed growth of peaks at 77 and 79  $m/z$ . In contrast to the  $\text{Xe}^+$  reactions, all reactant and product ions were detected with the same efficiency. This detection efficiency was determined by comparing the total number of ions present in the trap across all time points of the reaction and confirming that the number of charges was conserved. With the knowledge that no ions were lost over the course of the reaction, it was possible to determine the number of 40  $m/z$  product ions obscured by the large  $\text{Ca}^+$  signal. Specifically, the number of 40  $m/z$  product ions can be calculated by:

$$N_{40,\text{prod}}(t) = \text{O}_2^+(t=0) - (\text{O}_2^+(t) + N_{39}(t) + N_{77,79}(t)) \quad (3)$$

where  $N_{39}$  and  $N_{77,79}$  are the number of product ions at 39, 77, and 79  $m/z$  at time,  $t$ . This method of determining the number of 40  $m/z$  products relies on two assumptions, conservation of charge and consistent  $\text{O}_2^+$  loading from shot-to-shot. These assumptions were checked and verified by comparing total ion numbers and total reactant/product ion numbers across the reaction profile.

**3.2.1  $\text{O}_2^+ + \text{H}_2\text{C}_3\text{H}_2$ .** The reaction curve for the reaction between  $\text{O}_2^+$  and allene  $\text{H}_2\text{C}_3\text{H}_2$  is shown in Fig. 4. Similarly to the  $\text{Xe}^+$  reactions, there was a dominant 39  $m/z$  product pathway ( $\sim 73\%$  of products); however, in this



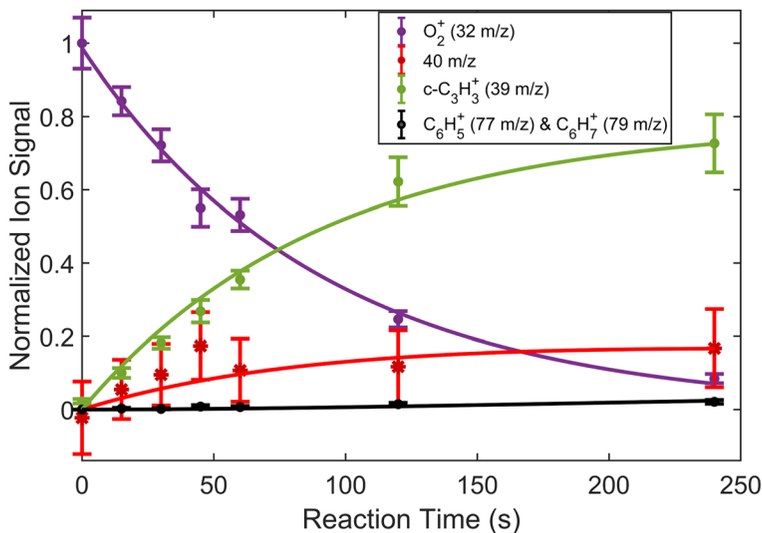


Fig. 4 Fraction of reactant and product ions in the reaction of  $\text{O}_2^+$  with  $\text{H}_2\text{C}_3\text{H}_2$  as a function of time. Circular data points correspond to mean and standard error of 14 measurements. Asterisk data points correspond to calculated mean and standard error of 40  $m/z$  products using eqn (3). All data points have been normalized to the initial number of  $\text{O}_2^+$  ions ( $\sim 180$  ions). Time was measured from the opening of the pulsed leak valve and subsequent inlet of allene into the chamber. Plotted lines correspond to calculated fits for the products and reactants using eqn (4)–(6).

reaction, there was also a clear 40  $m/z$  channel ( $\sim 17\%$  of products) that was absent in the  $\text{Xe}^+$  reactions. The growth of both the 39 and 40  $m/z$  channels was concurrent with the decay of the  $\text{O}_2^+$  signal, suggesting that both were primary products in this reaction. Once again, the 39  $m/z$  product showed no signs of further reacting into another channel. In addition, there was indication that some 40  $m/z$  product ions had reacted with another allene molecule to form secondary products. However, some of the 40  $m/z$  product ions did not react further. Based on the charge-to-mass ratios and presence/absence of further reactivity, we assigned some of the products as  $\text{c-C}_3\text{H}_3^+$  (39  $m/z$ ),  $\text{C}_3\text{H}_4^+$  (40  $m/z$ ),  $\text{C}_6\text{H}_5^+$  (77  $m/z$ ), and  $\text{C}_6\text{H}_7^+$  (79  $m/z$ ). However, the apparent stability of at least some of the 40  $m/z$  product ions suggests that not all the products in this channel were  $\text{C}_3\text{H}_4^+$  and may possibly include other products, such as  $\text{C}_2\text{O}^+$ .

The reaction data were fit assuming pseudo-first order kinetics with allene in excess and the following reaction model:

$$\frac{d\text{O}_2^+}{dt} = -(k_3 + k_4) \times \text{O}_2^+(t) \quad (4)$$

$$\frac{dN_{39}}{dt} = k_3 \times \text{O}_2^+(t) \quad (5)$$

$$\frac{dN_{77,79}}{dt} = k_5 \times (\text{O}_2^+(t=0) - \text{O}_2^+(t) - N_{39} - N_{77,79}) \quad (6)$$

where the fitting coefficients,  $k_3$  and  $k_5$ , represent rate constants, and  $k_4$  represents the rate for an additional pathway that  $\text{O}_2^+$  can react into to form 40  $m/z$ .



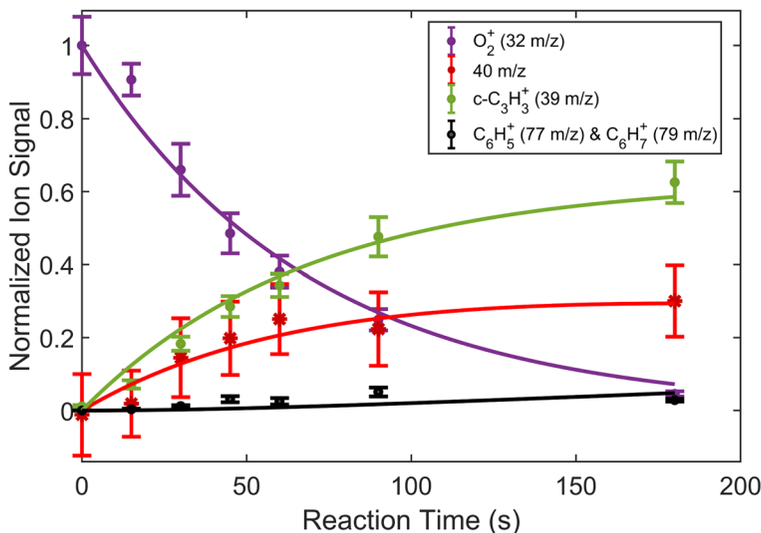


Fig. 5 Fraction of reactant and product ions in the reaction of  $\text{O}_2^+$  with  $\text{H}_3\text{C}_3\text{H}$  as a function of time. Circular data points correspond to mean and standard error of 13 measurements. Asterisk data points correspond to calculated mean and standard error of 40  $m/z$  products using eqn (3). All data points have been normalized to the initial number of  $\text{O}_2^+$  ions ( $\sim 190$  ions). Time was measured from the opening of the pulsed leak valve and subsequent inlet of propyne into the chamber. Plotted lines correspond to calculated fits for the reactants and products using eqn (4)–(6).

**3.2.2  $\text{O}_2^+ + \text{H}_3\text{C}_3\text{H}$ .** The reaction curve for the reaction between  $\text{O}_2^+$  and propyne ( $\text{H}_3\text{C}_3\text{H}$ ) is shown in Fig. 5, where the lines represent a fit to the data using the same model as  $\text{O}_2^+$  with allene ( $\text{H}_2\text{C}_3\text{H}_2$ ). This reaction was similar to that of  $\text{O}_2^+$  with allene. Once again, the dominant pathway produced  $c\text{-C}_3\text{H}_3^+$  ( $\sim 62\%$  of products), which does not go on to react again. Further, there was a pathway that yields a 40  $m/z$  primary product ( $\sim 30\%$  of products) that also appears to be mostly nonreactive. It is clear that some of the 40  $m/z$  ions are  $\text{C}_3\text{H}_4^+$ , as  $\sim 3\%$  of the observed products were found as the secondary products,  $\text{C}_6\text{H}_5^+$  and  $\text{C}_6\text{H}_7^+$ . However, the identity of the nonreactive 40  $m/z$  products is less clear, but may include other products, such as  $\text{C}_2\text{O}^+$ .

## 4 Discussion

Our overall goal for this study was to determine if isomer-specific reactivity previously observed with two isomers of  $\text{C}_3\text{H}_4$ , allene ( $\text{H}_2\text{C}_3\text{H}_2$ ) and propyne ( $\text{H}_3\text{C}_3\text{H}$ ), was a universal trait for ion–molecule reactions involving these molecules, or if the ion character plays an important role for complex formation as well. To explore this question, we studied reactions of allene and propyne with two ions,  $\text{Xe}^+$  and  $\text{O}_2^+$ , chosen because the energy available to the reaction is nearly identical in the two cases due to the very similar ionization potentials of  $\text{Xe}$  and  $\text{O}_2$ .

The results of the reactions with  $\text{Xe}^+$  and  $\text{O}_2^+$  clearly show distinct product distributions, demonstrating that the ionic reactant can be at least partially



responsible for differences in branching ratios. In the following, we investigate how these distinct product distributions provide evidence for these reactions progressing *via* direct charge transfer or *via* complex formation.

#### 4.1 Charge transfer driven reactions with Xe<sup>+</sup>

For the Xe<sup>+</sup> reactions with both C<sub>3</sub>H<sub>4</sub> isomers studied here, c-C<sub>3</sub>H<sub>3</sub><sup>+</sup> is the prevailing product. This result is similar to the case of acetylene (C<sub>2</sub>H<sub>2</sub><sup>+</sup>) with allene (H<sub>2</sub>C<sub>3</sub>H<sub>2</sub>), which was shown to progress *via* direct charge transfer. The dominance of the c-C<sub>3</sub>H<sub>3</sub><sup>+</sup> product for both Xe<sup>+</sup> reactions strongly suggests that these reactions proceed along the same reaction pathway regardless of isomer and theoretical calculations validated this conclusion.

In the Xe<sup>+</sup> reactions, there are two possibilities for how c-C<sub>3</sub>H<sub>3</sub><sup>+</sup> could be formed as a product, hydride abstraction by Xe<sup>+</sup> to form XeH or C–H bond dissociation of C<sub>3</sub>H<sub>4</sub><sup>+</sup>. The former relies on the formation of a complex between the Xe<sup>+</sup> and C<sub>3</sub>H<sub>4</sub>, but the latter does not as C<sub>3</sub>H<sub>4</sub><sup>+</sup> can be formed *via* direct charge transfer. While Xe<sup>+</sup> has been shown to occasionally, but rarely, form reaction complexes, the lack of any XeH<sup>+</sup> ions detected in our experiments suggests that complex formation is very unlikely.<sup>14–16</sup> Further, it was determined *via* quantum chemical calculations that the formation of XeH and C<sub>3</sub>H<sub>3</sub><sup>+</sup> is highly endothermic (>2 eV), making these products impossible to form in the current experimental system. Calculations were also conducted to determine if complex formation is energetically accessible, but all attempts led to endothermic pathways inaccessible for this experiment. Therefore, we find neither experimental nor theoretical support for complex formation in the reaction of Xe<sup>+</sup> + C<sub>3</sub>H<sub>4</sub>.

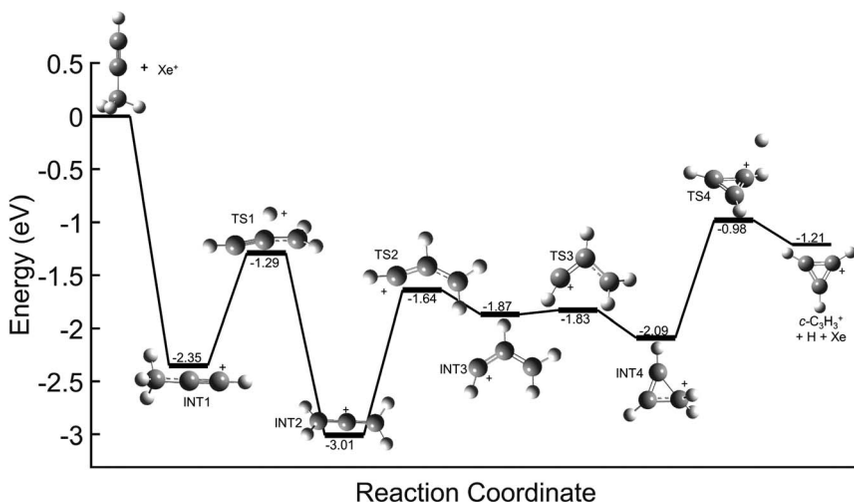


Fig. 6 Global potential energy surface of the reaction between Xe<sup>+</sup> and propyne (H<sub>3</sub>C<sub>3</sub>H) calculated at the ωb97x/aug-cc-pvtz level. All energies are shown relative to reactant energy at infinite separation, with INT1 corresponding to the first stationary point following charge transfer. Note that the allene (H<sub>2</sub>C<sub>3</sub>H<sub>2</sub>) cation is identified as INT2, and the PES for allene with Xe<sup>+</sup> is identical to that shown here with energies being shifted to higher energies by 5 meV.



In contrast, direct charge transfer and resultant dissociation of a hydrogen from  $C_3H_4^+$  is well supported. This can be seen in the potential energy surfaces (PESs) for the reaction of  $Xe^+$  with each  $C_3H_4$  isomer shown in Fig. 6. These surfaces begin with a highly exothermic charge transfer, with exothermicities of 2.35 eV and 2.96 eV for propyne ( $H_3C_3H$ ) and allene ( $H_2C_3H_2$ ) respectively. Because of the ultrahigh vacuum environment of the system, this energy is not quenched by collisions with other ions or neutral molecules. Instead, the ion retains a significant portion of this energy in its rovibrational modes, and the ion rapidly progresses along the PES. In both reactions, the  $C_3H_4^+$  can inter-convert between the allene cation (identified as INT2 in Fig. 6), propyne cation, and INT4 until transitioning to TS4, where the molecule dissociates yielding a hydrogen atom and ultimately goes on to form  $c-C_3H_3^+$ . These PESs demonstrate a barrierless pathway that leads to the production of  $c-C_3H_3^+$  without the formation of a complex between the  $Xe^+$  and  $C_3H_4$  isomers. Further, these PESs are nearly identical to that for the reaction of acetylene ( $C_2H_2^+$ ) with allene, which was determined to progress solely through direct charge transfer.<sup>8,9</sup> Therefore, this reveals that in the  $Xe^+$  reactions direct charge transfer can ultimately lead to formation of  $c-C_3H_3^+$  regardless of isomer.

#### 4.2 Evidence of complex formation with $O_2^+$

Comparisons between the products in the  $O_2^+$  and  $Xe^+$  reactions reveal that complex formation plays a role in the reaction mechanism for  $O_2^+$ . For example, in the reactions of  $O_2^+$  with both  $C_3H_4$  isomers studied here,  $c-C_3H_3^+$  is produced to a lesser extent than in the  $Xe^+$  reactions ( $\sim 60\text{--}75\%$  of total products for  $O_2^+$ , versus  $>95\%$  for  $Xe^+$ ). Further, in contrast to the  $Xe^+$  reactions, there is also noticeable growth in the 40  $m/z$  channel for both  $O_2^+$  reactions. Specifically, in the

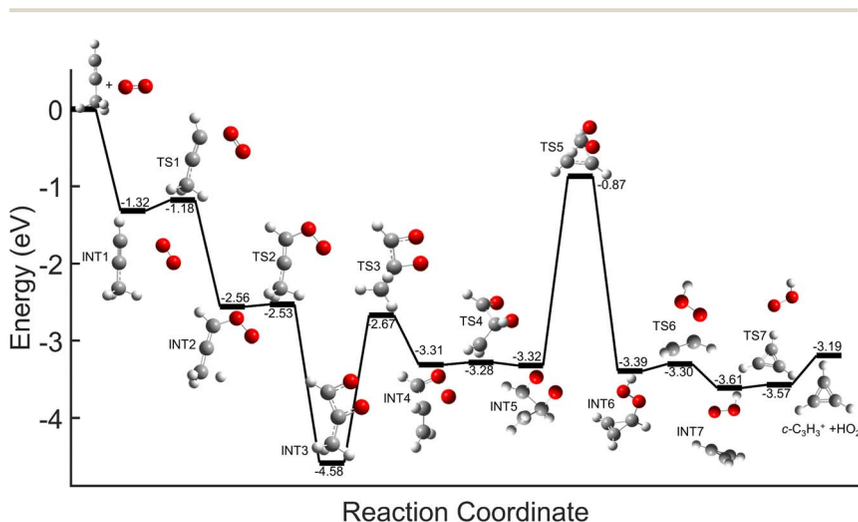


Fig. 7 Global potential energy surface of complex formation leading to  $c-C_3H_3^+$  production for  $O_2^+$  with propyne ( $H_3C_3H$ ) calculated at the  $\omega b97x/aug\text{-}cc\text{-}pvtz$  level. All energies are displayed relative to reactant energy at infinite separation. Note that states INT1–INT4 are identical to those shown in Fig. 8. These surfaces have been separated for clarity.



$O_2^+$  reactions, 40  $m/z$  products are present in branching ratios of  $\sim 17\%$  (allene) and  $\sim 30\%$  (propyne) compared to the branching ratios observed in the  $Xe^+$  reactions of  $<5\%$ . This discrepancy between the  $O_2^+$  and  $Xe^+$  reactions suggests that direct charge transfer is not the sole mechanism present in the  $O_2^+$  reactions. Additionally, the presence of multiple products is similar to the reaction of  $C_2H_2^+$  with propyne ( $H_3C_3H$ ), which progressed *via* complex formation. Therefore, these results suggest that  $O_2^+$  forms a complex with each  $C_3H_4$  isomer, and this complex formation aids in the production of the 40  $m/z$  products,  $C_3H_4^+$  and  $C_2O^+$ .

The conclusion that both isomers of  $C_3H_4$  form a complex with  $O_2^+$  is strengthened through calculation of PESSs. Plausible PESSs by which complex formation can lead to formation of  $C_3H_4^+$  and  $c-C_3H_3^+$  in the reaction with propyne ( $H_3C_3H$ ) are shown separately in Fig. 7 and 8 for clarity. The propyne with  $O_2^+$  surfaces for  $C_3H_4^+$  and  $c-C_3H_3^+$  production both begin with barrierless, exothermic ( $\sim 1.32$  eV) complex formation, and follow similar trajectories for the attachment of both oxygen atoms onto the complex. The two surfaces ultimately separate at INT4 at which point the complex can lead to production of  $C_3H_4^+$  or  $c-C_3H_3^+$ . In the production of  $C_3H_4^+$ , the majority of intermediate states are relatively similar in energy allowing for rapid inter-conversion and sampling of states. Further,  $C_3H_4^+$  can be produced in two separate isomers depending on when and how the  $O_2$  detaches from the complex. Specifically,  $C_3H_4^+$  can be produced either in the  $H_2C_3H_2^+$  isomer if the  $O_2$  detaches at TS6, or it can be produced in the  $H_3C_3H^+$  isomer if the complex regresses at TS1 and detaches  $O_2$ . In contrast to this, the surface for production of  $c-C_3H_3^+$  has a relatively high-energy transition state (TS5, 0.87 eV exothermic compared to reactants) necessary for transferring the hydrogen to the attached  $O_2$ .  $c-C_3H_3^+$  production is finalized upon dissociation of  $HO_2$  from the complex. Surfaces for allene ( $H_2C_3H_2$ ) were calculated, but were not included here because of their similarity with the surfaces for propyne.

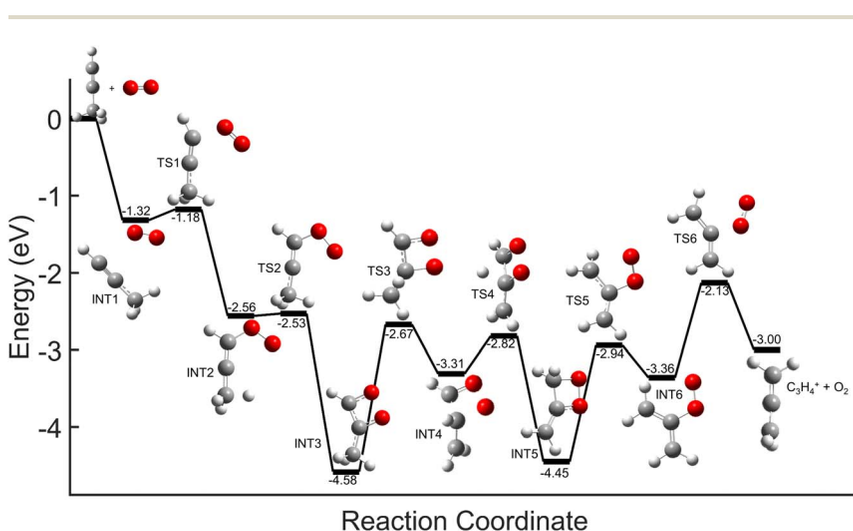


Fig. 8 Global potential energy surface of complex formation leading to  $C_3H_4^+$  production for  $O_2^+$  with propyne ( $H_3C_3H$ ) calculated at the  $\omega b97x/aug-cc-pvtz$  level. All energies are displayed relative to reactant energy. Note that states INT1–INT4 are identical to those shown in Fig. 7. These surfaces have been separated for clarity.



The surfaces provided demonstrate a barrierless pathway for the production of both  $c\text{-C}_3\text{H}_3^+$  and  $\text{C}_3\text{H}_4^+$  given complex formation.

However, not all of the 40  $m/z$  products observed in the  $\text{O}_2^+$  reactions can be accounted for by  $\text{C}_3\text{H}_4^+$ . This is because it is known that  $\text{C}_3\text{H}_4^+$  reacts with  $\text{C}_3\text{H}_4$  to form larger hydrocarbons, and many of the 40  $m/z$  products observed here appear to be stable. While other molecules exist with a charge-to-mass ratio of 40  $m/z$ ,  $\text{C}_2\text{O}^+$  is one of the most likely products. For example,  $\text{C}_2\text{O}^+$  has a simple combination of atoms involved in the reaction that seem plausible given complex formation. Further, calculations find that production of  $\text{C}_2\text{O}^+$  and its co-product,  $\text{CH}_3\text{OH}$ , is exothermic ( $\sim 2.9$  eV). Finding a plausible PES for production of  $\text{C}_2\text{O}^+$  is beyond the scope of our current work. Verifying the presence of  $\text{C}_2\text{O}^+$  *via* isotopic substitution will be the subject of a future study.

## 5 Conclusions

The reactions of  $\text{Xe}^+$  and  $\text{O}_2^+$  with two isomers of  $\text{C}_3\text{H}_4$  demonstrate that the identity of the ion, not just the charge-transfer energetics or the isomer structure, contributes to the reaction pathway.  $\text{Xe}^+$  reactions proceed solely *via* direct charge transfer reactions, principally forming  $c\text{-C}_3\text{H}_3^+$ , which is unsurprising. However,  $\text{O}_2^+$  forms complexes with *both* isomers resulting in a more diverse product distribution, including  $c\text{-C}_3\text{H}_3^+$ ,  $\text{C}_3\text{H}_4^+$ , and possibly  $\text{C}_2\text{O}^+$ . Combining these results with our previous work on reactions between acetylene ions ( $\text{C}_2\text{H}_2^+$ ) and  $\text{C}_3\text{H}_4$  isomers, we find that  $\text{Xe}^+$  and  $\text{O}_2^+$  behave as the two extremes cases where the mechanism is driven by the ion identity, whereas the acetylene cation behaves in a manner whereby the neutral isomer determines the mechanism. While  $\text{Xe}^+$  solely performs direct charge transfer and  $\text{O}_2^+$  undergoes complex formation with both  $\text{C}_3\text{H}_4$  isomers, acetylene performs direct charge transfer with one isomer, allene, and complex formation with the other, propyne. The ability to demonstrate a reaction mechanism that is dependent on the ion identity, independent of the ionization potential of the ion, can be uniquely demonstrated in this pair of reactions. These findings suggest that in the case of the two isomers of  $\text{C}_3\text{H}_4$  studied here, either the identity of the ion, or the structure of the neutral can contribute to the propensity of a reaction complex forming or direct charge transfer occurring. Further studies will be required to better understand how each component independently influences this propensity and their relative importance in doing so.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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