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Reactions of Atomic Carbon with Butene Isomers: Implications for Molecular Growth in Carbon-Rich Environments

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ABSTRACT

Product detection studies of C(3P) atom reactions with butene (C4H8) isomers (but-1-ene, cis-but-2-ene, trans-but-2-ene) are carried out in a flow tube reactor at 353 K and 4 Torr under multiple collision conditions. Ground state carbon atoms are generated by 248 nm laser photolysis of tetrabromomethane, CBr4, in a buffer of helium. Thermalized reaction products are detected using synchrotron tunable VUV photoionization and time of flight mass spectrometry. The temporal profiles of the detected ions are used to discriminate products from side or secondary reactions. For the C(3P) + trans-but-2-ene and C(3P) + cis-but-2-ene reactions, various isomers of C4H5 and C5H7 are identified as reaction products formed via CH3- and H-elimination. Assuming equal ionization cross sections for all C4H5 and C5H7 isomers, C4H5:C5H7 branching ratios of 0.63:1 and 0.60:1 are derived for the C(3P) + trans-but-2-ene and the C(3P) + cis-but-2-ene reactions, respectively. For the C(3P) + but-1-ene reaction, two reaction channels are observed: the H-elimination channel, leading to the formation of the ethylpropargyl isomer, and the C3H5 + C2H5 channel. Assuming equal ionization cross sections for ethylpropargyl and C3H5 radicals, a branching ratio of 1:0.95 for the C3H5 + C2H5 and H + ethylpropargyl channels is derived. The experimental results are compared to previous H-atom branching ratios and used to propose the most likely
mechanisms for the reaction of ground state carbon atoms with butene isomers.

I. INTRODUCTION

Carbon is the fourth most abundant element in the universe and unbounded carbon atoms have been detected in a wide range of environments, from interstellar media (dense and diffuse clouds, protoplanetary nebulae, circumstellar envelopes...)^1-5 to planetary atmospheres (e.g. Mars, Venus, Titan),^6-8 and plasmas.^9-11 In these environments, their barrierless reactions^12-35 with abundant saturated and unsaturated hydrocarbons, nitrogen bearing molecules, and small free radicals are expected to initiate the growth of complex molecules^9,36,37 even at very low temperatures.^12-14,33,34 Consequently, carbon atom reactions are expected to play a role in the formation of polycyclic aromatic hydrocarbons (PAHs), aerosols, and soot.^9,36,37 Improving the accuracy of models trying to reproduce the chemistry of such environments requires a better understanding of the mechanisms of elementary reactions between carbon atoms and abundant hydrocarbons. This goal can be achieved by probing the products of these reactions experimentally and theoretically over a wide range of physical conditions.

Most of the previous research on C(\(^3\)P) reactivity focused on reactions with hydrocarbons containing fewer than 4 carbon atoms, investigated using various experimental techniques such as crossed molecular beams,^16,19-21,27,35,38,39 flow tubes,^23,26,31,40 or CRESU experiments,^13,14,41 often coupled with ab initio calculations.^18,24,25 The accepted dominant reaction mechanism derived from these experiments is addition of the C(\(^3\)P) to the hydrocarbon followed by the loss of a hydrogen atom or methyl group to form an open-shell molecule.^23,40 For large unsaturated hydrocarbons such as butene isomers (C\(^4\)H\(_8\)), exit channels other than H-atom or CH\(_3\) loss may become accessible.^23

Butene is one of the smallest hydrocarbons with structural isomers and is a good archetype molecule for the study of gaseous fuel reactivity. It is a negligible component in
gasoline but is produced during the combustion of larger hydrocarbons.\textsuperscript{42-44} The high temperature (>1200 K) pyrolysis of butene isomers leads to the formation of reactive free radicals such as methyl (CH\textsubscript{3}), propargyl (C\textsubscript{3}H\textsubscript{3}) and allyl (C\textsubscript{3}H\textsubscript{5}) and ultimately to the formation of larger molecules such as benzene and C\textsubscript{3}H\textsubscript{6} isomers.\textsuperscript{45} Under plasma conditions, butene is known to decompose easily into CH\textsubscript{x} and C\textsubscript{2}H\textsubscript{x} (x=1,2,3) species and is used as a source of carbon atoms.\textsuperscript{46} Butene isomers are also believed to play a role in the photochemistry of organic molecules in planetary atmospheres (e.g. Titan).\textsuperscript{47,48} They are expected to react rapidly with atoms such as C, O, or N and with small hydrocarbons radicals such as CH or C\textsubscript{2}H. Although rate constants and product distributions of butene reactions have been shown to depend on the specific butene isomer,\textsuperscript{49} computational model of these environments do not consider isomeric C\textsubscript{4}H\textsubscript{8} structures, likely leading to uncertainties in the derived abundances of large species. Atkinson \& Aschmann\textsuperscript{49} determined rate constants for the reactions of OH radical with but-1-ene, cis-but-2-ene, trans-but-2-ene and isobutene at 295±1 K to be (3.13±0.16)×10\textsuperscript{-10} cm\textsuperscript{3} molecules\textsuperscript{-1} s\textsuperscript{-1}, (5.60±0.13)×10\textsuperscript{-10} cm\textsuperscript{3} molecules\textsuperscript{-1} s\textsuperscript{-1}, (6.39±0.13)×10\textsuperscript{-10} cm\textsuperscript{3} molecules\textsuperscript{-1} s\textsuperscript{-1} and (5.13±0.24)×10\textsuperscript{-10} cm\textsuperscript{3} molecules\textsuperscript{-1} s\textsuperscript{-1}, respectively. Loison \textit{et al.}\textsuperscript{50} investigated the lighter products formed by the reaction of OH radicals with trans-but-2-ene, and but-1-ene in a fast flow reactor at room temperature. The branching fractions for the H-atom abstraction channel were measured to be 8±3\% for OH + but-1-ene, and 3±1\% for OH + trans-but-2-ene. More recently Bouwman \textit{et al.}\textsuperscript{51} investigated C\textsubscript{2}H radical reactions with butene isomers (but-1-ene, cis-but-2-ene, trans-but-2-ene and isobutene) in a Laval expansion at 79 K. They found similar rate constants for all reactions (ca. 2×10\textsuperscript{-10} cm\textsuperscript{3} molecules\textsuperscript{-1} s\textsuperscript{-1}) in agreement with previous measurements performed at room temperature,\textsuperscript{52,53} but observed an isomer dependence of the reaction products. It is worth noting that this work also showed that the C\textsubscript{2}H + C\textsubscript{4}H\textsubscript{8} reactions contribute to molecular weight growth with formation of larger species (4-penten-1-yne, 3-penten-1-yne,
2-methyl-1-buten-3-yne, 4-methyl-3-penten-1-yne) through methyl radical elimination in addition to H-atom elimination.

The reaction kinetics of C(3P) + butene isomers was first studied by Haider & Husain\(^{12}\) at room temperature in a slow flow reactor. They measured overall rate constants for reactions with but-1-ene, cis-but-2-ene and trans-but-2-ene of \((4.3\pm0.2)\times10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}\), \((3.2\pm0.2)\times10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}\) and \((2.9\pm0.2)\times10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}\), respectively. More recently, Loison & Bergeat\(^{23}\) measured an overall rate coefficient for the C(3P) + trans-but-2-ene reaction of \((1.9\pm0.6)\times10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}\), which is a factor of 1.5 smaller than the value of Haider & Husain. Loison & Bergeat\(^{23}\) suggest the initial formation of a triplet H\(_3\)C–HC•–C=CH–CH\(_3\) adduct. From this intermediate, the accessible exothermic channels are H + C\(_5\)H\(_7\) and CH\(_3\) + C\(_4\)H\(_5\) through isomerization and C–H or C–C bond breaking, respectively. In addition Loison & Bergeat measured an absolute atomic hydrogen branching fraction of 0.33 ± 0.08 using H-atom resonance fluorescence. The authors stress the fact that the isomerization of the initial reaction adduct (H\(_3\)CHCCCHCH\(_3\)) may greatly affect the final isomer distribution. Subsequently, Li et al.\(^{54}\) performed a theoretical study of the ground state carbon atom reaction with trans-but-2-ene by probing the complex triplet potential energy surface. They predict two major products: dimethylpropargyl (H\(_3\)CHCCCH\(_3\)) and a methylpropargyl isomer (CH\(_3\)CHCCH), in agreement with Loison and Bergeat.\(^{23}\) The authors confirm a barrierless association reaction followed by isomerization and dissociation of the initial adduct. The large reaction rate coefficient measured at room temperature is therefore expected to remain high at low temperatures, making this reaction of interest for both interstellar medium and plasma assisted combustion chemistry.

The role of the C(3P) + butene reactions in gas phase environments depends on the importance of the H elimination exit channel relative to elimination of larger hydrocarbon
radicals, as well as on the isomeric structure of the final products. If C₅H₇ isomers are produced, the reactions will contribute to the formation of long linear carbon chains in carbon rich environments. If C₄H₅ isomers are formed, the reactions could play a role in the chemical scheme leading to the formation of aromatic rings. C₅H₇-isomers (i-C₄H₅ and methylpropargyl isomers) have been observed both in fuel-rich flames and as C(3P) + propene reaction products. The reaction of acetylene (C₂H₂) with the i-C₄H₅ isomer is suggested as a possible route to benzene (c-C₆H₆) formation under certain flame conditions. The reaction would initially form fulvene (c-C₅H₄=CH₂) which could then convert to benzene by H-atom-assisted isomerization. The role of the methylpropargyl isomers (CH₃CHCCH and CH₃CCCH₂) in ring formation processes is still unknown and requires further investigation. For instance, CH₃CHCCH, is completely absent from flame models whereas CH₃CCCH₂ is suggested to form toluene C₇H₈ (C₆H₅–CH₃) or benzyl radical C₇H₇ (C₆H₅–CH₂) by reaction with the propargyl radical (CH₃CCH). In this work we present an investigation of the products formed by reactions of triplet ground state carbon atoms C(3P) with three butene isomers: but-1-ene, cis-but-2-ene and trans-but-2-ene. In order to probe the reaction products, experiments are performed in a flow tube reactor under multiple collision conditions (353 K & 4 Torr) at the Advanced Light Source (ALS) synchrotron of Lawrence Berkeley National Laboratory. Detection of the products in the flow is carried out using tunable Vacuum Ultraviolet (VUV) photoionization and time of flight mass spectrometry. Kinetic traces and photoionization spectra supported by thermodynamic and Franck-Condon factor calculations of the C₄H₅ and C₅H₇ species have been used to infer the primary products of the reactions.

II. EXPERIMENTAL PROCEDURE

A description of the apparatus has been given elsewhere and only a brief overview is presented here. Reactions are performed in a quartz flow tube with a length of 62 cm and
1.05 cm inner diameter. The gas pressure inside is typically 4 Torr with a total gas flow of 250 sccm (standard cubic centimeters per minute). All experiments are carried out at 353 K corresponding to a total number density of $1.09 \times 10^{17} \text{ cm}^{-3}$ and a slow flow velocity ($\sim 10 \text{ m s}^{-1}$). The buffer gas (helium), carbon atom precursor ($\text{CBr}_4$) and reagent gas (butene isomers) are metered by mass-flow controllers. Reagent gases and the precursor were used without further purification: but-1-ene ($> 99\%$, Aldrich), cis-but-2-ene ($> 99\%$, Aldrich), trans-but-2-ene ($> 99\%$, Aldrich) & CBr$_4$ (99%, Aldrich). CBr$_4$ was introduced in the reactor with a typical density of $\sim 4 \times 10^{12} \text{ cm}^{-3}$, by flowing a small amount of He over solid CBr$_4$ upstream of the reaction tube. At $\sim 30 \text{ cm}$ along the tube, a 650 µm pinhole allows a small portion of the gas mixture to expand into a vacuum chamber. There, a skimmer generates a beam of molecules entering a differentially pumped ionization chamber in which molecular and atomic species are ionized by quasi-continuous VUV synchrotron radiation. Ions formed are detected using an orthogonal-acceleration time-of-flight mass spectrometer pulsed at 50 kHz. All detected ions are time-tagged relative to the excimer laser pulse (at $t=0$) operating at 10 Hz and accumulated in a transient digitizer to collect complete mass spectra as a function of time. It should be noted that the operation frequency of the laser allows sufficient time between laser pulses to completely refresh the gas mixture. The total ion detection time-window is 50 ms, beginning 10 ms before the laser pulse. Time-and energy-resolved mass spectra are obtained by averaging at least 500 laser shots at each photon energy. All data are normalized to account for variations in VUV photon flux using a NIST-calibrated photodiode (SXUV-100, International Radiation Detectors, Inc.). At each mass-to-charge ($m/z$) ratio, the average signal present before the laser pulse is subtracted from the time dependent data. The photoionization spectra at a given $m/z$ ratio are obtained by integrating the data set over the desired mass and time windows. The ionizing photon energy is scanned from 7.4 to 9.0 eV in steps of 0.025 eV. Kr or Ar is used in a gas filter to absorb higher energy photons at
harmonics of the undulator energy. Up to three independent datasets are recorded for each
reaction and averaged. The photon energy resolution is determined to be of the order of 40
meV by measuring an atomic resonance of Xe.

Carbon atoms were generated by 248 nm photolysis of CBr$_4$, using a pulsed excimer
(KrF) laser operating at 10 Hz. The laser power output is typically 266 mJ per pulse (20 ns),
with a fluence inside the reaction flow tube of ~20-50 mJ cm$^{-2}$. The multiphoton
photodissociation of CBr$_4$ at 248 nm eliminates multiple halogen atoms, leading to the
formation of carbon atoms$^{62}$ and CBr radicals.$^{63,64}$ Under our experimental conditions 20% of
the introduced CBr$_4$ is photodissociated. Shannon et al.$^{63}$ determined a C(1D)/C(3P) ratio
following CBr$_4$ photodissocation at 266 nm of 0.15 ± 0.04 at 296 K. Molecular nitrogen is
known to efficiently quench C(1D) atoms through C(1D) + N$_2$ →7 C(3P) + N$_2$ with a reaction
rate coefficient of $k = 5.3 \times 10^{-12}$ cm$^3$ molecules$^{-1}$ s$^{-1}$. In order to avoid potential chemical
interferences from the reaction between C(1D) and butene, the present experiments were
performed using a large excess of N$_2$ (1.8 × 10$^{16}$ cm$^{-3}$). In the present experimental conditions
(353 K & 4 Torr), for known densities of reactant (C$_4$H$_8$ ~ 10$^{14}$ cm$^{-3}$) and considering a C(3P)
+ C$_4$H$_8$ reaction rate constant of $k = 1.9 \times 10^{-10}$ cm$^3$ molecules$^{-1}$ s$^{-1}$, the characteristic decay
times are calculated to be five times shorter for C(1D) reaction with molecular nitrogen than
for the C(3P) reaction with butene isomers. Consequently, in the following sections the
influence of C(1D) atoms reactions with butene isomers is neglected. In addition, due to the
high efficiency of C(3P) fine structure relaxation by collision with He, the spin-orbit level
population distribution of ground state carbon atoms, C(3 P$^0,1,2$) in the flow tube are expected
to be thermalized.$^{13,66}$ The impact of CBr reactions with butene on product identification is
discussed in the Results section. Finally, it is worth noting that the absorption cross-section of
1-butene was measured to be 6.8 × 10$^{-21}$ cm$^2$ at 220 nm,$^{67}$ likely leading to negligible
dissociation of the butene isomers at the longer wavelength used here.
III. COMPUTATIONAL METHODS

Electronic structure calculations of the neutral C₅H₇ species and their cations are carried out using the Gaussian09 package employing the B3LYP/CBSB7 method to obtain optimized geometries. Theoretical calculations with the same methods were applied for C₄H₅ isomers in a previous study.²⁶ Heats of reaction and adiabatic and vertical ionization energies are obtained using the CBS-QB3 composite method. All the vibrational frequencies are real, indicating that the optimized geometries represent minima on the potential energy surface.

Photoionization spectra of C₅H₇ isomers are calculated at 298 K using the G09 package within the Franck–Condon approximation.⁶⁸ The simulated factors are convolved with a 40 meV (FWHM) Gaussian response function and integrated to simulate direct photoionization spectra.

IV. RESULTS

Table 1 reports the optimized structures for seven C₅H₇ isomers together with their computed ionization energies. Because the cationic form of the cyclic isomer cannot be optimized, the value reported is the vertical ionization energy. As mentioned above, C₄H₅ isomers have been investigated in a previous study.²⁶

<table>
<thead>
<tr>
<th>Isomer</th>
<th>Optimized structure</th>
<th>Ionization energy (eV)</th>
<th>Isomer</th>
<th>Optimized structure¹</th>
<th>Ionization energy (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trans C₅H₇</td>
<td>![Image]</td>
<td>7.10</td>
<td>Ethylpropargyl</td>
<td>![Image]</td>
<td>7.88</td>
</tr>
<tr>
<td>Cis C₅H₇</td>
<td>![Image]</td>
<td>7.19</td>
<td>Trans 3-C₅H₇</td>
<td>![Image]</td>
<td>7.40</td>
</tr>
<tr>
<td>Dimethylpropargyl</td>
<td>![Image]</td>
<td>7.41</td>
<td>Cis 3-C₅H₇</td>
<td>![Image]</td>
<td>7.41</td>
</tr>
<tr>
<td>c-C₅H₇</td>
<td>![Image]</td>
<td>8.19</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 2 presents computed enthalpies of reaction at 298 K for all the predicted reaction channels and investigated \( \text{C}_4\text{H}_8 \) isomers. All of the reaction channels are found to be exothermic. The difference between isomers is outside the error of the method which is \( \sim 5 \text{ kJ mol}^{-1} \) on averaged prediction of \( \Delta f H^\circ(298.15 \text{ K}) \) for CBS–X methods.\(^6\) Within these uncertainties, the enthalpies of reaction displayed in Table 2 for the \( \text{C}(^3\text{P}) + \text{trans–C}_4\text{H}_8 \) reaction are similar to those calculated by Li et al. (G3B3//B3LYP/6-311G(d,p)).\(^5\) These values are all lower than the enthalpies calculated by Loison and Bergeat based on experimental enthalpies of formation.\(^2\)

<table>
<thead>
<tr>
<th>Channels</th>
<th>Trans–C(_3\text{H}_7) + H</th>
<th>Cis–C(_4\text{H}_8)</th>
<th>I–C(_4\text{H}_8)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trans–C(_3\text{H}_7) + H</td>
<td>-204.52</td>
<td>-209.75</td>
<td>-215.00</td>
</tr>
<tr>
<td>Cis–C(_3\text{H}_7) + H</td>
<td>-207.13</td>
<td>-212.37</td>
<td>-217.62</td>
</tr>
<tr>
<td>Dimethylpropargyl + H</td>
<td>-209.75</td>
<td>-215.00</td>
<td>-220.24</td>
</tr>
<tr>
<td>Ethylpropargyl + H</td>
<td>-199.26</td>
<td>-204.51</td>
<td>-209.75</td>
</tr>
<tr>
<td>3-C(_3\text{H}_7) + H</td>
<td>-201.89</td>
<td>-207.13</td>
<td>-212.37</td>
</tr>
<tr>
<td>c-C(_3\text{H}_7) + H</td>
<td>-49.82</td>
<td>-55.06</td>
<td>-60.30</td>
</tr>
<tr>
<td>i-C(_3\text{H}_7) + CH(_3)</td>
<td>-243.84</td>
<td>-249.08</td>
<td>-254.32</td>
</tr>
<tr>
<td>CH(_3)CHCCH + CH(_3)</td>
<td>-238.59</td>
<td>-243.84</td>
<td>-249.08</td>
</tr>
<tr>
<td>CH(_3)CCCH(_2) + CH(_3)</td>
<td>-249.08</td>
<td>-254.32</td>
<td>-259.57</td>
</tr>
<tr>
<td>C(_3\text{H}_3) + C(_3\text{H}_5)</td>
<td>-236.64</td>
<td>-247.66</td>
<td>-247.14</td>
</tr>
</tbody>
</table>

\(^{a}\text{CBS–QB3}\)

\(^{b}\text{Cis and Trans}\)

Photoion spectra were recorded with and without \( \text{CBr}_4 \) and with or without the photolysis laser pulse. In the absence of butene in the reactor flow, no ion signals were detected at \( m/\ell \) 15, 53 and 67, corresponding to the expected \( \text{C}(^3\text{P}) + \text{butene} \) product masses (\( \text{CH}_3, \text{C}_4\text{H}_5 \) and \( \text{C}_5\text{H}_7 \)). The reactivity of \( \text{CBr} \) radicals with butene isomers has to be considered to ensure that products formed at \( m/\ell \) 53 and 67 do not originate from reactions involving \( \text{CBr} \). The rate coefficient of \( \text{trans–C}_4\text{H}_8 \) reaction with \( \text{CBr} \) has been determined by Ruzsicska et al.\(^7\) to be \( 1.23 \times 10^{-11} \text{ cm}^3 \text{ molecules}^{-1} \text{ s}^{-1} \), which is fifteen times lower than that of the \( \text{C}(^3\text{P}) \) reaction with butene.\(^2\) Consequently, under the present experimental conditions, \( \text{CBr} + \text{butene} \) products are formed more slowly than the \( \text{C}(^3\text{P}) + \text{butene} \) products. Previous
investigations on the C(\(^3\)P) + ethylene and C(\(^3\)P) + propylene reactions have shown that such slower secondary processes and side reactions with characteristic times of the order of few hundreds of microseconds can be discriminated against primary processes by inspecting the product time traces.\(^{26}\) Furthermore, for all the results presented below, the contributions from slow reactions or subsequent reactions are minimized by integrating mass spectra only over the 0–1 ms time range.

In the sections below, we analyze mass spectra and photoionization spectra obtained upon irradiation of a mixture of CBr\(_4\) with three different butene isomers: but-1-ene, \(\text{cis}\)-but-2-ene, \(\text{trans}\)-but-2-ene. Photoionization spectra acquired at 4 and 8 Torr do not show significant differences. In the following section only spectra recorded at 4 Torr are presented. Comparing photoionization spectra to previous results or to calculated photoionization spectra identifies the isomeric structure of the reaction products.

The time behavior of the non-normalized ion signal after the laser pulse is used to discriminate between C(\(^3\)P) reaction products and products from side reactions as well as to infer product branching fractions. The signals \(S_i\) for the main products are fitted using Eq. 1 corresponding to a pseudo-first order model of sequential reactions.\(^{71}\)

\[
S_i(t) = \frac{S_i \cdot k_{1,i}}{k_{2,i} - k_{1,i}} \cdot e^{-k_{1,i}t}
\]

Eq.1

The coefficients \(k_{1,i}\) and \(k_{2,i}\) are the pseudo first order rate constants for the formation and loss of the product \(i\). \(S_i\) is proportional to the amount of reactants consumed to form product \(i\).

Branching ratios are obtained from the ratio of the \(S_i\) values for two different channels assuming equal ionization cross sections.

**A. C(\(^3\)P) + trans-but-2-ene**
FIG. 1 displays the mass spectrum obtained following photolysis of a mixture of CBr₄ and trans-but-2-ene in helium and nitrogen. The data are recorded at 9.0 eV photon energy.

The negative signals at m/z 28, 41, 56 and 127 are due to imperfect subtraction of ion signal from butene or from impurities present in the reaction flow. Over the short reaction window (1 ms), the main ion signals are observed at m/z 53, 55, 67, 68, 91, 93, 146, 147, 148 & 149. Products at m/z 91 & 93 are identified to be C⁷⁹Br and C⁸¹Br, formed by photodissociation of CBr₄ at 248 nm.

FIG. 1. Mass spectrum obtained by photolysis of a CBr₄ and trans-but-2-ene mixture in helium at 9.0 eV photon energy integrated over the 0–1 ms time range.

FIG. 2 shows the normalized averaged kinetic traces of m/z 146/148 (red circles) and m/z 147/149 (blue squares) together with modeled C₄H₇Br (red line) and C₅H₈Br (blue line) temporal traces. The m/z 147/149 and m/z 146/148 temporal profiles are modeled using the following scheme:

\[
\begin{align*}
\text{CBr} + \text{C}_4\text{H}_8 & \rightarrow \text{C}_5\text{H}_7\text{Br} + \text{H} \quad \text{(R1a)} \\
\text{CBr} + \text{C}_4\text{H}_8 & \rightarrow \text{C}_5\text{H}_8\text{Br}^* \quad \text{(R1b)} \\
\text{C}_5\text{H}_8\text{Br}^* & \rightarrow \text{C}_5\text{H}_7\text{Br} + \text{H} \quad \text{(R2)}
\end{align*}
\]
The metastable $\text{C}_5\text{H}_8\text{Br}^*$ at m/z 147/149 is formed by the $\text{CBr} + \text{C}_4\text{H}_8$ reaction (R1b) $(k_{1a+1b}=1.23 \times 10^{-11} \text{ cm}^3 \text{ molecules}^{-1} \text{ s}^{-1})^{70}$ and decays $(k_2=500 \text{ s}^{-1})$ to give $\text{C}_5\text{H}_7\text{Br} + \text{H}$ (R2).

The temporal profile for $\text{C}_5\text{H}_7\text{Br}$ at m/z 146/148 signal is the sum of the contributions from $\text{CBr} + \text{C}_4\text{H}_8$ (R1a) and $\text{C}_5\text{H}_8\text{Br}$ decomposition to $\text{C}_5\text{H}_7\text{Br} + \text{H}$ (R2). The data are best reproduced with branching ratios of 0.5–0.6 for reaction R1a and 0.5–0.4 for reaction R1b.

FIG. 2. Normalized averaged kinetic traces of m/z 146/148 (red circles) and m/z 147/149 (blue squares) obtained by photolysis of a CBr$_4$ and trans-but-2-ene mixture in helium, at 9.0 eV photon energy. The thick lines are modeled traces for $\text{C}_5\text{H}_7\text{Br}$ (red line) and $\text{C}_5\text{H}_8\text{Br}$ (blue line).

FIG. 3 shows the normalized temporal traces of (a) m/z 53 (C$_4$H$_5$, blue down triangles) and 67 (C$_5$H$_7$, purple triangles), (b) m/z 55 (C$_4$H$_7$, cyan diamonds) and 67 (purple triangles), and (c) m/z 68 (C$_5$H$_8$, dark yellow left triangles) and 67 (purple triangles). Thick blue, cyan and purple lines are double exponential fits to m/z 53, 55 and 67, respectively. The thick dark yellow line is a single exponential fit to m/z 68. According to these kinetic traces, m/z 53 & 67 are likely to be formed by the fast C($^3\text{P}$) + C$_4$H$_8$ reaction, as suggested by the
350 µs time constant of the observed rise, corresponding to a large value of $k_{1st} (\geq 3.500 \text{ s}^{-1})$. The diffusion corrected$^{72,73}$ pseudo first order rate ($\sim 9,000 \text{ s}^{-1}$) is still lower than the predicted one ($19,000 \text{ s}^{-1}$), likely due to the instrument response function. Here we assume that the rising edges of these two signals represent creation of $\text{C}_4\text{H}_3$ and $\text{C}_5\text{H}_7$. By contrast, the kinetic traces of $m/z$ 55 & 68 show a slower increase ($k_{1st} \sim 1,000 \text{ s}^{-1}$), suggesting that they are formed by reactions slower than $\text{C}(^3\text{P}) + \text{C}_4\text{H}_8$.

FIG. 3. Normalized kinetic traces obtained by photolysis of a CBr$_4$ and trans-but-2-ene mixture in helium, at 9.0 eV photon energy of (a) $m/z$ 53 (blue down triangles) & 67 (purple
triangles) (b) m/z 55 (cyan diamonds) & 67 (purple triangles) and (c) m/z 68 (dark yellow left triangles) & 67 (purple triangles). Thick blue, cyan & purple lines are double exponential fits to m/z 53, 55 & 67 respectively. The thick dark yellow line is a single exponential fit to m/z 68.

From the above analysis, C₅H₇ and C₄H₅ isomers are likely to be the primary products of the C(3P) + butene reaction through H-elimination and CH₃-elimination. The branching ratio between these two channels can be estimated directly from the values of Sᵢ obtained from the fit of the temporal traces corrected by the relative ionization cross sections and the mass discrimination factor. As these cross sections are unknown, they are assumed to be equal. Consequently, the ratio of the Sᵢ values gives only an estimate of the H– vs. CH₃–loss branching. The fit of the m/z 53 and 67 temporal signals suggests a 0.63:1 C₄H₅/C₅H₇ branching ratio. No data were recorded at photon energies above the IE of methyl radical (9.83 eV), although it is expected it is formed.

FIG. 4 displays the m/z 53 photoionization spectrum obtained by irradiating the CBr₄ and trans-but-2-ene mixture at 248 nm. The data are integrated over the 0-40 ms time range and displayed from 7.4 to 9.0 eV photon energy. Within the experimental uncertainty, the absence of an ion signal onset at 7.55 eV suggests that the i-C₄H₅ isomer is not formed by the C(3P) + trans-but-2-ene reaction. The ion onset at 7.9 eV suggests the presence of one or both of the methylpropargyl isomers (IE = 7.94 eV for H₂CCCCH₃ and IE = 7.95 eV for HCCCHCH₃). Within the experimental photon energy resolution (40 meV) it is not possible to differentiate between the two methylpropargyl isomers. FIG. 4 also displays the integrated Franck-Condon factor simulations for the i-C₄H₅ (dashed cyan line) and methylpropargyl isomers (dashed and dotted blue & dotted green lines). The spectra for each isomer are normalized to unity at 9.0 eV, because the simulated spectra are constant above this photon
energy. The thick red line is the simulated spectrum of the methylpropargyl isomer H₂CCCCH₃ normalized to the experimental data. The data are well reproduced with no contribution from i-C₄H₅. The absence of the i-C₄H₅ isomer and the formation of one or both methylpropargyl isomers differs from previous studies on C₄H₅ formation.²⁶,⁵⁷

FIG. 4. Photoionization spectrum of m/z 53 obtained by photolysis of a CBr₄ and trans-but-2-ene mixture in helium integrated over the 0-40 ms time range and displayed from 7.4 to 9.0 eV photon energy. The lines are the integrated Franck-Condon factor simulations for i-C₄H₅ (dash cyan line) and methylpropargyl isomers (H₂CCCCH₃ dashed-doted blue line & HCCCHCH₃ doted green lines). The thick red line is the integrated Franck-Condon factor simulation of H₂CCCCH₃ normalized to the experimental data.

Due to small signal-to-noise-ratio in the photoionization spectrum at m/z 67 it is not possible to infer the contribution from the various C₅H₇ isomers. The faster decay of this signal compared with m/z = 53 makes the C₅H₇ spectra much noisier. However, the detection of ion signal above 7.5 eV is consistent with the possible formation of the dimethylpropargyl isomers as suggested by Bergeat & Loison²³ and Li et al.⁵⁴
In FIG. 3, the decay rate of the m/z 67 signal (428 s⁻¹) is greater than that of the m/z 53 signal (42 s⁻¹). From the product identification above, these signals are likely to be due to ionization of dimethylpropargyl and methylpropargyl. The presence of a methyl group on either terminal carbon of the propargyl radical may result in hyperconjugation between the partially filled p orbital and the σ orbital from a C–H bond in beta position. Although hyperconjugation stabilizes the radical site, it weakens the methyl C–H bond, facilitating H-abstraction by Br or H atoms. The large difference in decay rate between the two radicals could be due to the dependence of the hyperconjugation strength on the isomeric structure of the substituted propargyl radical and the number of methyl groups. Abstraction of a H-atom from dimethylpropargyl is consistent with the detection of a signal at m/z 66 with a slow formation rate (not shown). Within the experimental signal-to-noise, no signal is detected at m/z 52.

Photoionization spectra were also recorded for secondary products, m/z 55 and 68 (not shown here). The presence of an ion onset at ca. 7.5 eV for m/z 55 suggests the contribution of one or two 1-methylallyl isomers (C₄H₇). They are likely formed by abstraction of an H atom from butene by reactive species such as Br, CBr, or H. The presence of an ion onset at 8.8 eV in the photoionization spectrum for m/z 68 suggests its identification as isoprene.

B. C(3P) + cis-but-2-ene

The mass spectrum registered (not shown here) following photolysis of a CBr₄ and cis-but-2-ene mixture in helium and nitrogen leads to the observation of the same four major peaks m/z 53, 55, 67 & 68 as for trans-but-2-ene. FIG. 5 shows the normalized temporal traces of (a) m/z 53 (blue down triangles) and 67 (purple triangles) (b) m/z 55 (cyan diamonds) and 67 (purple triangles) and (c) m/z 68 (dark yellow left triangles) and 67 (purple triangles). Thick blue, cyan and purple lines are double exponential fits to m/z 53, 55 and 67 respectively. The
thick dark yellow line is a single exponential fit to m/z 68. The first-order rise of m/z 53 and 67, around 3000–4000 s⁻¹, suggests that they are primary reaction products as for the reaction of C(3P) with trans-but-2-ene. The first-order rise coefficients of m/z 55 and 68, are 1230 and 1386 s⁻¹, suggesting products formed by secondary or slower reactions. The fit of the m/z 53 and 67 signals from FIG. 5 lead to a 0.60:1 C₄H₅/C₅H₇ branching ratio.

FIG. 5. Normalized kinetic traces obtained by photolysis of a CBr₄ and cis-but-2-ene mixture in helium, at 9.0 eV photon energy of (a) m/z 53 (blue down triangles) & 67 (purple triangles)
(b) m/z 55 (cyan diamonds) & 67 (purple triangles) and (c) m/z 68 (dark yellow left triangles) & 67 (purple triangles). Thick blue, cyan & purple lines are double exponential fits to m/z 53, 55 & 67 respectively. The thick dark yellow line is a single exponential fit to m/z 68.

FIG. 6 displays the m/z 53 photoionization spectrum obtained by irradiating a CBr₄ and cis-but-2-ene mixture at 248 nm. The data are integrated over the 0-40 ms time range and displayed from 7.4 to 8.8 eV photon energy. Similar to the results for trans-but-2-ene, there is no evidence for i-C₄H₅ isomer, and clear evidence for one or both methyl propargyl isomers.

FIG. 6. Photoionization spectrum of m/z 53 obtained by photolysis of a CBr₄ and cis-but-2-ene mixture in helium integrated over the 0-40 ms time range and displayed from 7.4 to 8.8 eV photon energy. The lines are the integrated Franck-Condon factor simulations for i-C₅H₅ (dash cyan line) and methylpropargyl isomers (H₂CCCCH₃ dashed-dotted blue line & HCCCHCH₃ dotted green lines). The thick red line is the integrated Franck-Condon factor simulation of H₂CCCCH₃ normalized to the experimental data.

FIG. 7 displays the m/z 67 photoionization spectrum obtained by irradiating a CBr₄ and cis-but-2-ene mixture at 248 nm. The red solid line represents the normalized integrated
Franck Condon factors of the dimethylpropargyl isomer while the dashed line is that of the ethylpropargyl isomer. The relatively good match between the dimethylpropargyl simulated spectrum and experimental spectrum suggests that dimethylpropargyl is formed by the reaction. Due to the low signal however, it is not possible to determine the contribution from the ethylpropargyl isomer to the signal.

FIG. 7. Photoionization spectrum of \textit{m/z} 67 obtained by photolysis of a CBr$_4$ and \textit{cis}-but-2-ene mixture in helium integrated over the 0-40 ms time range and displayed from 7.4 to 8.8 eV photon energy. The thick red line and the dash green line are the integrated Franck-Condor factor simulation for dimethylpropargyl (red line) and ethylpropargyl (green line) respectively.

\textbf{C. \textit{C}(^3\textit{P}) + but-1-ene}

FIG. 8 displays a time resolved mass spectrum obtained by photolysis of a CBr$_4$ and but-1-ene mixture in helium and nitrogen at 9.0 eV photon energy from -2.5 to 35 ms relative to the laser pulse. The main time-resolved signals are observed at \textit{m/z} 39, 55, 66, 67 and 68. By contrast with experiments performed with \textit{cis}- and \textit{trans}-but-2-ene, \textit{m/z} 53 is not
observed, and signals at m/z 29 and 39 appear after the photolysis laser pulse. Experiments performed while doubling the concentrations of all the reactants (not shown here) led to similar kinetic traces. As for the reaction with trans-C4H8, signals at m/z 55 and 68 are identified as methylallyl and isoprene. Signal at m/z 29, 39 and 67 are likely to originate from ionization of C2H5, C3H3, and C5H7 respectively, corresponding to the C3H3 + C2H5 and H + C3H7 exit channels of the C(3P) + but-1-ene reaction. Signal at m/z 66 could come from abstraction of a hydrogen atom from m/z 67 as discussed in Section IV A. The signal at m/z 39 is clearly identified as propargyl radical due to an ion onset at the propargyl ionization energy (IE=8.70 eV) and clear autoionization resonances. It should be noted, that when the concentration of butene is increased, the signal at m/z 29 is not observed anymore, while the signal at m/z 39 is still present. This is likely due to fast reaction of the C2H5 radical, produced by the C(3P) + butene reaction, with butene. This is consistent with the fact that C2H5 reactions with hydrocarbons are expected to be faster than reactions with the resonance stabilized propargyl (C3H3) radical (see KIDA datasheet at http://kida.obs.u-bordeaux1.fr/).

![3D mass spectrum obtained by photolysis of a CBr4 and but-1-ene mixture in helium](image)
at 9.0 eV photon energy from -2.5 to 35 ms relatively to the laser pulse.

FIG. 9 displays the \textit{m/z} 67 photoionization spectrum obtained by irradiating a CBr$_4$ and but-1-ene mixture in helium and nitrogen at 248 nm. The data are integrated over the 0-40 ms time range and displayed from 7.4 to 8.8 eV photon energy. The green dashed line is the integrated Franck-Condon factor simulation for ethylpropargyl and the thick red line that for dimethylpropargyl. The reasonable match between the integrated Franck-Condon factor simulation for ethylpropargyl and the experimental data suggests that ethylpropargyl is the main C$_3$H$_7$ isomer formed by the reaction C($^3$P) + but-1-ene. The product branching ratio for the C$_3$H$_3$/C$_5$H$_7$ isomers can be estimated directly from the ratio of their ion signals assuming equal ionization cross sections. The \textit{m/z} 39 and 67 signals from the mass spectrum lead to a 1:0.95 branching ratio for the C$_3$H$_3$ + C$_2$H$_5$ and H + C$_5$H$_7$ exit channels.

![Photoionization spectrum of m/z 67](image)
respectively.

V. DISCUSSION

For the C(3P) reactions with the trans- and cis-but-2-ene isomers, assuming equal ionization cross sections for the C4H5 and C5H7 isomers, C4H5:C5H7 branching ratios are found to be 0.63:1 and 0.60:1 ratio, respectively. This suggests that the stereoisomery has no effect on the product ratios. The results for the trans-but-2-ene isomer differs from the 0.33±0.06 absolute H-atom production branching fraction measured in fast flow reactor monitoring H-atom fluorescence by Loison and Bergeat\textsuperscript{23} corresponding to a 1:0.5 C4H5:C5H7 branching ratio. Li et al. propose a possible 1:1 branching fraction based solely on the energetics of the products and transition states.\textsuperscript{54} Previous work on branching fraction for the C(3P) + C3H6 reaction led to reasonable agreement between our results using tunable VUV photoionization with time of flight mass spectrometry and those obtained Loison and Bergeat using H atom fluorescence detection.\textsuperscript{26} The discrepancy reported here could be due to the low signal in the present experiments for the C5H7 isomers combined with the assumed equivalence of the photoionization cross section of these isomers. For these reasons the values reported by Loison and Bergeat\textsuperscript{23} are likely to be more accurate. For the C(3P) reaction with but-1-ene, the 1:0.95 branching fraction for the C3H3:C3H7 isomers derived from our experiments suggests similar probability of losing a H or C2H5 group from the initial C3H8 reaction adduct. No experimental or theoretical values are available for comparison.

C3H7 production, corresponding to the H elimination channel, is observed for all three C(3P) + butene isomer reactions. In the case of the C(3P) + trans-but-2-ene reaction, the theoretical investigation by Li et al. suggests that dimethylpropargyl (CH3CHCCCH3) is the most likely isomer produced. Although this cannot not be confirmed by the present study for the C(3P) + trans-but-2-ene reaction, the fit of the photoionization spectrum presented in FIG. 7 is consistent with the formation of the dimethylpropargyl isomer by the C(3P) + cis-but-2-
nene reaction. Regarding the $\text{C}^3\text{P} + \text{but-}1\text{-ene}$ reaction, the photoionization spectrum in FIG. 9 suggests ethylpropargyl as the main product isomer.

FIG. 10 presents a proposed mechanism for the direct formation of the dimethylpropargyl and ethylpropargyl isomers from $\text{C}^3\text{P}$ reactions with the butene isomers. As previously proposed$^{26}$ the carbon atom is likely to interact with the $\pi$-electron system of the unsaturated hydrocarbon to form a 3-carbon ring. In the case of but-2-ene, different stereoisomers of the cyclic intermediate will be formed by reaction with the cis- or trans-but-2-ene. The strained isomer is likely to have a very short life time ($<1\text{ps}$) as suggested by Loison et al.$^{23}$ for reaction with butene and Kaiser et al. for reaction with ethylene.$^{16}$ Upon ring opening it will form a substituted triplet allene-like intermediate. Elimination of a hydrogen atom from one of the C–H groups in alpha position to the triplet carbon leads to the formation of dimethylpropargyl for reaction with trans/cis-but-2-ene and ethylpropargyl with but-1-ene. The photoionization spectra at $m/z$ 67 reported here are consistent with the previously discussed mechanism, although a more complex isomerization schemes could lead to other $\text{C}_5\text{H}_7$ isomers. The formation of the energetically accessible $\text{HCCC(CH}_3\text{)}\text{CH}_3$ would require a $\text{CH}_3$ transfer after formation of the cyclic intermediate.

![Proposed mechanisms](image.png)

FIG. 10. Proposed mechanisms for the formation of $\text{C}_5\text{H}_7$ isomers from $\text{C}(^3\text{P})$ reactions with butene isomers.
C₄H₅ isomers are observed for both C(³P) + trans- and cis-but-2-ene reactions, while they are not detected for the C(³P) + but-1-ene reaction. The absence of C₄H₅ products in the case of but-1-ene is consistent with the absence of a methyl group on the carbon double bond of the initial reactants. This suggests a rapid dissociation of the acyclic adduct through breaking of the C–H or C–C bonds initially on the C=C group of the reactant without further isomerization. FIG. 11 shows the decomposition of the initial C₅H₈ adduct into ethyl + propargyl.

![Diagram](https://example.com/diagram.png)

FIG. 11. Formation of propargyl radical from C₂H₅-loss from the C(³P) + but-1-ene reaction adducts.

Interestingly, in the case of C(³P) + trans- and cis-but-2-ene reactions where C₄H₅ is formed, the photoinization spectra, displayed in FIG. 4 and 6, differ significantly from those obtained for the C + C₃H₆ reactions by Capron et al.²⁶ or for fuel-rich flames experiments by Hansen et al.⁵⁷ The main difference seems to be due to the absence of the i-C₄H₅ isomer (IE=7.55 eV).²⁶ This contrast can be explained by the different loss channels of the intermediates formed by ring opening of the adducts from the C + propene or C + butene reactions. FIG. 12 displays the adduct formed by reaction of carbon atoms with cis-but-2-ene (top) and propene (bottom). In the case of the reaction with propene, the i-C₄H₅ isomers are formed directly by elimination of an H-atom from the methyl group.²⁶ In the case of CH₃ loss from the C₅H₈ intermediate, the isomer formed by the most direct pathway is the CH₂CHCCH methylpropargyl isomer. Formation of the i-C₄H₅ isomer would require a hydrogen transfer from the methyl group to a carbon in γ position. Although within our experimental energy resolution it is not possible to distinguish between the two methylpropargyl isomers, the detection of one or both of the methylpropargyl isomers is
consistent with mechanisms proposed by Loison & Bergeat and Li et al. for the C(3\(^3\)P) reaction with \textit{trans}-but-2-ene. Li et al. suggest CH\(_2\)CHCCH as a likely candidate for C\(_4\)H\(_5\) isomers from C + \textit{trans}-C\(_4\)H\(_8\) reaction due to either kinetic or thermodynamic factors. Indeed, from FIG. 12 formation of CH\(_3\)CCCH\(_2\) would require a H-atom transfer to a carbon in \(\beta\) position. Distinguishing between the two isomers would be of interest as CH\(_3\)CHCCH is completely absent from flames models whereas CH\(_3\)CCCH\(_2\) is proposed to form toluene (C\(_7\)H\(_8\)) or benzyl radical (C\(_7\)H\(_7\)) by reaction with another CH\(_3\)CCH\(_2\) or with propargyl radical.\(^{58}\)

![Diagram of CH\(_3\)CCCH\(_2\) and CH\(_3\)CHCCH isomers formation from H-loss and CH\(_3\) loss.](image)

FIG. 12. Formation of C\(_4\)H\(_5\) isomer from CH\(_3\) or H-loss from the C(3\(^3\)P) + \textit{cis/trans}-but-2-ene (top) and C + propene (bottom) reaction adducts.\(^{29}\)

Although carbon atoms may not be formed in standard flames and engines, they are known to be present in plasma and plasma-assisted environments.\(^{9-11}\) Similar to the CH radical, carbon atoms will react with the most abundant fuel molecules through addition/elimination pathways to form larger molecules. Such mechanisms still need to be validated, especially for large hydrocarbon molecules used as fuels. Coupling plasmas to combustion systems has the potential to greatly improve the combustion process through both an increase in combustion efficiency and a reduction in the emission of pollutants (e.g. NOx, CO\(_2\), soot).\(^{78}\) Feedback control algorithms using emission concentration as error signals and applied plasma as control signals could allow the combustion device to run at its stability limit. The design of such plasma-assisted technologies requires a basic understanding of the
combined combustion and plasma chemistry leading to increased heat release and decreased emissions.

Isomers of the \( \text{C}_4\text{H}_5 \) and \( \text{C}_5\text{H}_7 \) hydrocarbon radicals have been detected in standard combustion flames where they contribute to the formation of larger molecules, including aromatic hydrocarbons.\(^{58,79} \) Their detection from the \( \text{C}(^3\text{P}) + \text{butene} \) isomers reactions suggests that the formation of these radicals will be enhanced under plasma assisted combustion conditions potentially changing the molecular growth chemical scheme. The estimated branching ratios obtained in the present experiments are important data for the design of new combustion models under high-energy conditions such as those found in plasma. Ultimately these models will be used to tailor new combustion strategies with lower emission and higher efficiency.

Carbon-containing molecules can take various forms such as carbon chain molecules, polycyclic aromatic hydrocarbons (PAHs), carbon clusters and carbonaceous solids.\(^{80} \) \( \text{C}(^3\text{P}) \) atoms have been detected in dense interstellar clouds\(^{81-83} \) where a large fraction of the observed interstellar molecules are formed. None of the \( \text{C}_4\text{H}_8 \) isomers however, have been observed so far, likely due to their low electric dipole moments (\( \leq 0.5 \) D). Their predicted abundances in astrochemical models, formed through \( \text{C} \) and \( \text{C}^+ \) reactions, seems to be sensitive to reactions involving atomic oxygen and nitrogen.\(^{84} \) In methane rich atmospheres such as Titan’s, hydrocarbon formation is initiated by the photochemistry of methane that produces more complex and less saturated hydrocarbons.\(^{47,48,85-88} \) Although \( \text{C}_4\text{H}_8 \) has not been detected in Titan’s atmosphere,\(^{47} \) Lavvas and coworkers\(^{48} \) for instance, included the reaction of carbon with 1-butene isomer in their photochemical model of Titan’s atmosphere with no information on the products. Zhang \( \textit{et al.} \)\(^{6} \) used a 1D-photochemical model of Titan to produce an atomic carbon profile reproducing the atomic carbon emission at 1657Å observed by the Cassini Ultra-violet Imaging Spectrograph and stress the need of a full set of
photochemical reactions involving chemical species containing at least three carbon atoms to
describe carbon atom chemistry above 600 km. Recently, Dobrijevic and coworkers\textsuperscript{88} derived
abundance profiles from a 1D-photochemical model coupling neutral and ion chemistries,
showing that C\textsubscript{4}H\textsubscript{8} (with no isomer distinction) is among the most abundant C\textsubscript{4}H\textsubscript{x} species
above 600 km. Such large hydrocarbons could also play a role in the formation of organic
hazes observed in Titan’s atmosphere.\textsuperscript{89}

The rate coefficient of the reaction C(\textsuperscript{3}P) trans-but-2-ene was measured by Loison et
al. at room temperature ($k = 1.9\times10^{-10}$ cm$^3$ molecules$^{-1}$ s$^{-1}$)\textsuperscript{23} and is likely to be of the same
order of magnitude for cis-but-2-ene and but-1-ene. A large number of reactions between
ground state carbon atoms and small saturated or unsaturated hydrocarbons have been shown
to be fast at room temperature\textsuperscript{12,23,33,40,90} with rate coefficients increasing with decreasing
temperature.\textsuperscript{13,14,91} Similarly, it is expected that reactions of C(\textsuperscript{3}P) with butene isomers
remain fast down to very low temperatures. Furthermore, the present investigation shows that
even for such a simple system, the title reaction can form quite a large number of
hydrocarbon radicals such as C\textsubscript{4}H\textsubscript{5} isomers (H\textsubscript{2}CCCCH\textsubscript{3} and/or HCCCHCH\textsubscript{3}), propargyl
(C\textsubscript{3}H\textsubscript{3}), dimethylpropargyl and ethylpropargyl. Although the methyl radical is not detected it
is expected to be formed by reaction of carbon atoms with trans- and cis-butene. The nature
and branching ratios of these radicals are found to depend on the C\textsubscript{4}H\textsubscript{8} isomers considered,
and are consistent with simple addition/elimination schemes shown in Figs.10 - 12. All these
radicals are susceptible to participate in the formation and destruction of other molecules
present in these environments. This stresses the importance of including molecular isomer
resolution for both reactants and products in interstellar astrochemical and atmospheric
photochemical models in order to estimate their abundances as well as their potential
importance in chemical schemes leading to the formation and destruction of complex
molecules.
VI. CONCLUSION

The ground state C(\(^3\)P) reactions with butene isomers (but-\(1\)-ene, \textit{cis}-but-2-ene, \textit{trans}-but-2-ene) were studied under multiple collision conditions\(^{26}\) at 353 K and 4 Torr using tunable photoionization coupled to time-of-flight mass spectrometry. Products from secondary or side reactions are distinguished from primary products using the temporal profile of the detected ions. The reactions are performed under multiple collision conditions from 4 to 8 Torr with no significant pressure dependence of the product distributions. Although the initial energized adduct is expected to have a very short lifetime (<1ps),\(^{16,23}\) statistical calculations have shown that some of its isomers may have longer lifetime, up to tens of nanoseconds for intermediates of the C(\(^3\)P) + C\(_2\)H\(_2\) reaction for instance.\(^{18}\)

Experimental or theoretical studies over a wider range of pressure are required to fully investigate the effect of collisional quenching on the final product distribution and the role of intersystem crossing.

In the case of C(\(^3\)P) + \textit{trans}-but-2-ene and C(\(^3\)P) + \textit{cis}-but-2-ene reactions, the observed reaction exit channels are C\(_4\)H\(_5\) + CH\(_3\) and C\(_5\)H\(_7\) + H. Assuming similar ionization cross sections for the CH\(_3\)- and H-coproducts, the C\(_4\)H\(_5\):C\(_5\)H\(_7\) branching ratios are 0.63:1 and 0.60:1 for reactions with \textit{trans}- and \textit{cis}-2-butene, respectively. These values differ from the 1:0.5 CH\(_3\):H-loss ratio measured in fast flow reactor\(^{23}\) or the 1:1 ratio suggested by Li et al.\(^{54}\) from the analysis of the C\(_5\)H\(_8\) potential energy surface. The difference between the experimental branching ratios is likely to be in part due to the absence of data on the photoionization cross sections for the C\(_5\)H\(_7\) and C\(_4\)H\(_5\) isomers. Further experimental or theoretical studies are required to obtain C\(_4\)H\(_5\) and C\(_5\)H\(_7\) photoionization cross-section and therefore more accurate branching ratios.

For the C(\(^3\)P) + \textit{but-1-ene} reaction, the observed reaction channels are C\(_3\)H\(_3\) + C\(_2\)H\(_5\) and C\(_3\)H\(_7\) + H. The measured C\(_3\)H\(_3\):C\(_5\)H\(_7\) signal ratio from the present flow data is found to
be 1:0.95 assuming equal photoionization cross sections for the C$_3$H$_3$ and C$_3$H$_7$ radicals. No C$_4$H$_5$ formation is observed for this reaction. The non-detection of C$_4$H$_5$ for reaction with but-1-ene is consistent with direct addition of the carbon atom onto the carbon double bond followed by rapid ring opening to form a triplet allene-like structure. The most likely fate of the energized intermediate is a C–C bond breaking in β position from the carbene center. In the case of reaction with but-1-ene, this direct addition-elimination mechanism leads to the formation of the observed C$_2$H$_5$ and C$_3$H$_3$ radicals.

Photoionization spectra at the mass of the C$_4$H$_5$ isomers for the *trans* and *cis* isomers show formation of one or both methylpropargyl isomers and negligible formation of *i*-C$_4$H$_5$. Within the experimental energy resolution, it is not possible to distinguish between the two methylpropargyl isomers. Photoionization spectra at the mass of the C$_5$H$_7$ isomers display very low signal-to-noise ratio and do not allow the unequivocal identification of the formed isomers. The detection of signal below 7.8 eV is consistent with the formation of dimethylpropargyl for reaction with the *trans* and *cis* isomers. In the case of the reaction with but-1-ene, the photoionization spectrum is well reproduced by the simulated spectrum of only ethylpropargyl.

Overall, the isomer-resolved product detection for the C($^3$P) + butene isomer reactions is consistent with the addition of the carbon atom to the double bond to form a cyclic intermediate. This intermediate is likely to have a very short lifetime and to isomerize through ring opening to a triplet allene-like intermediate. The experimental data can all be explained using direct H-, CH$_3$-, or C$_2$H$_5$-elimination without further isomerization of the energized intermediates. Experimental and theoretical determinations of open-shell molecule photoionization cross-sections are required in order to improve the accuracy of branching ratios inferred from photoionization mass spectrometry studies. The reaction of ground state carbon atoms with butene isomers in carbon rich environments such as the ISM or plasma
assisted combustion is therefore likely to contribute to the formation of large carbon radicals and therefore to molecular growth chemical schemes.

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