A coupled-cluster ab initio study of triplet C$_3$H$_2$ and the neutral–neutral reaction to interstellar C$_3$H

Christian Ochsenfeld, Ralf I. Kaiser, Yuan T. Lee,a) Arthur G. Suits, and Martin Head-Gordon
Department of Chemistry, University of California at Berkeley, Berkeley, California 94720

(Received 8 October 1996; accepted 6 December 1996)

For the initially formed C$_3$H$_2$ collision complexes of molecular beam experiments ab initio calculations are presented. Resolving energetics and properties of these intermediates is essential for the understanding of the reaction of C$_3$(P) with C$_2$H$_2$ to form interstellar cyclic and linear isomers of C$_3$H. Computed reaction energies agree with results from molecular beam experiments. The combination of crossed molecular beam experiments and ab initio calculations allows us to identify two reaction channels for the carbon–hydrogen exchange and to explain astronomical observations of a higher c-C$_3$H to 1-C$_3$H ratio in dark clouds as compared to hotter envelopes of carbon stars.

© 1997 American Institute of Physics. [S0021-9606(97)03110-3]

I. INTRODUCTION

Reactions involving atomic carbon in its $^3P$ electronic ground state have received great interest both theoretically and experimentally due to the importance in hydrocarbon syntheses, combustion processes, and interstellar chemistry. Isomers of C$_3$H$_2$ and C$_3$H were detected in interstellar clouds and envelopes of evolved carbon stars. Cyclopropenylidene, c-C$_3$H$_2$, was discovered in 1985 in a cold dark cloud, Taurus Molecular Cloud 1 (TMC-1), by Matthews et al. and Thaddeus et al. Another isomer of C$_3$H$_2$, vinylidenecarbene (H$_2$CC=), was first observed in interstellar space by Cernicharo et al. At the same time the hydrocarbon radical C$_3$H was first detected in its linear form (propynlydine) by Thaddeus et al. via microwave spectroscopy towards TMC-1 and the carbon star IRC+10216. A corresponding cyclic isomer, c-C$_3$H (cyclopropynylidene), was found only 2 years later in TMC-1 by Yamamoto et al.

Although interstellar fractional abundances of such hydrocarbons relative to atomic hydrogen of up to 10$^{-8}$ are relatively large compared to other hydrocarbon species, their formation mechanisms in space have not yet been fully understood. Due to very small average kinetic energies of interstellar molecules of typically 0.8 kJ/mol in diffuse clouds and only 0.08 kJ/mol in dark clouds, reactions to these interstellar species must have little or no barriers. Further, only two-body collisions provide reasonable reaction probabilities. Reaction models for interstellar chemistry were focusing on radiative association, dissociative recombination, and exothermic ion–molecule processes (e.g., Refs. 5, 6). However, these mechanisms were not able to reproduce observed number densities and isomer ratios such as for the linear and cyclic form of C$_3$H. Recent kinetic studies by Clary et al. showed no barriers in fast neutral–neutral reactions of atomic C($^3P$) with unsaturated hydrocarbons and initiated the interest in such reaction paths. However, the formation of C$_3$H remained unclear.

To investigate this unresolved problem in interstellar chemistry, the formation process of C$_3$H via neutral–neutral reactions has been studied using crossed molecular beam experiments by three of us, R. I. K., Y. T. L., and A. G. S. In the present theoretical ab initio investigation our main interest is to elucidate energetics and properties of the initially formed triplet C$_3$H$_2$ collision complexes and to compute accurate reaction energies using high level electronic structure theory methods. We restrict ourselves to the triplet surface, as the C$_3$H$_2$ intermediates do not fulfill requirements of intersystem crossing.

Although this molecule has been subject of a multitude of theoretical and experimental studies, a complete investigation of the different possible structures using high level ab initio methods and sufficiently large basis sets for structures, energetics, and vibrational frequencies was still lacking. Similar interest has been attributed to the C$_3$H molecule both theoretically and experimentally. The main difficulty of this system is the occurrence of symmetry breaking in theoretical treatments. Such problems for the cyclic C$_3$H structure have been solved by a state of the art theoretical study of Stanton using the equation of motion coupled cluster singles and doubles approximation for ionized states (EOM-CCSD) method (e.g., Ref. 33), where C$_{3v}$ symmetry is found in agreement with experiments. However, symmetry determination—by accurate calculation of vibrational frequencies—remains an open issue for the linear structure, although attempts have been made to solve these problems. Due to both methodological difficulties and basis set deficiencies, theoretical studies so far do not yet provide conclusive theoretical estimates concerning the symmetry of the linear equilibrium structure of C$_3$H. However, this is an issue which we will not address in this study, as our main interest here in C$_3$H are relative energies and reaction energies which are shown to be only very little

\begin{equation}
\text{C}_2\text{H}_2 + \text{C} \rightarrow \text{C}_3\text{H}_2 \rightarrow \text{C}_3\text{H} + \text{H}
\end{equation}
influenced by stability problems. Relative energies of the cyclic and the linear form of this radical are as well an important issue, as previous results of theoretical calculations reside in a wide range between 1 kJ/mol (Ref. 27) and 75 kJ/mol.

II. DETAILS OF COMPUTATION

All ab initio calculations have been carried out with the program system ACES II.34 Unrestricted Hartree–Fock (UHF) wave functions are used and only pure spherical harmonic components of \(d\), \(f\), and \(g\) functions are included. Throughout this study the single- and double-excitation coupled cluster with a perturbational estimate of triple excitations [CCSD(T)] method has been employed. Some self-consistent field (SCF), Møller–Plesset second order perturbation theory (MP2) (Ref. 36), and CCSD (Ref. 37) relative energies are given for comparison to show the inadequacy of simple approaches like SCF and MP2 for \(C_2H_2\) and \(C_3H\).

Further, Brueckner CC methods38–43 have been used for the treatment of \(C_3H\), where symmetry-breaking problems play an important role. As the single excitation \((T_1)\) amplitudes account for most of the orbital relaxation effects in CC methods, in Brueckner methods the so called Brueckner determinant is used as a reference function, which is defined as the one for which single excitations do not mix with the exact wave function [full configuration interaction (FCI)]. In approximate methods such as the CCSD method, the reference function for which the \(T_1\) amplitudes vanish leads to the B-CCD method.39–43 This approach has proven its usefulness for the treatment of symmetry breaking problems such as encountered for example in case of \(NO_3\) (Ref. 43) or \(O_4^+\).44

Stability of the zeroth-order SCF wave functions (e.g., Ref. 45) has been checked throughout this study. Spin contamination occurring in the SCF wave function (typically 2.40 for \(S^2\)) in triplet states of \(C_2H_2\) is mostly eliminated within the CC scheme [to typically about 2.05 for \(S^2\) for the projected spin multiplicities using \(T_1\) and \(T_2\) (Ref. 46)]. Such a behavior is well known for CC methods and reflects the insensitivity with respect to the choice of orbitals.

All structures were fully optimized at the CCSD(T) level imposing proper symmetries of \(D_{2h}\) and subgroups. To characterize stationary points [local minima or saddle points, IR data, and zero-point vibrational energies (ZPE)], vibrational frequencies have been calculated numerically within the harmonic approximation using analytic CCSD(T) gradients.48 To recognize possible problems in numerical calculations of frequencies due to unstable or differing zeroth-order wave functions, frequencies have been computed as a check both analytically and numerically at the SCF level using the same structure.

The following basis sets have been used:

- DZP:  
  \[
  \begin{align*}
  \text{H}: & \quad (4s1p)/[2s1p] \\
  \text{C}: & \quad (8s4p1d)/[4s2p1d]
  \end{align*}
  \]

- TZP:  
  \[
  \begin{align*}
  \text{H}: & \quad (5s1p)/[3s1p] \\
  \text{C}: & \quad (10s6p1d)/[6s3p1d]
  \end{align*}
  \]

- TZPP:  
  \[
  \begin{align*}
  \text{H}: & \quad (5s2p1d)/[3s2p1d] \\
  \text{C}: & \quad (10s6p2d1f)/[6s3p2d1f]
  \end{align*}
  \]

- QZ2P:  
  \[
  \begin{align*}
  \text{H}: & \quad (7s2p1d)/[4s2p1d] \\
  \text{C}: & \quad (11s7p2d1f)/[6s4p2d1f]
  \end{align*}
  \]

- cc-pVQZ:  
  \[
  \begin{align*}
  \text{H}: & \quad (6s3p2d1f)/[4s3p2d1f] \\
  \text{C}: & \quad (12s6p3d2f1g)/[5s4p3d2f1g]
  \end{align*}
  \]

The double and triple zeta polarization basis sets,49 DZP and TZP, respectively, were used for geometry optimizations and the calculation of vibrational frequencies. The triple zeta double polarization basis (TZPP) (Ref. 49) was used partly for geometry optimizations and the calculation of frequencies to check the accuracy of the smaller basis sets. The quadruple zeta double polarization basis denoted QZ2P (Ref. 49) and the correlation consistent polarization valence quadruple zeta (cc-pVQZ) basis50 (which contains up to \(g\) functions on \(C\) and up to \(f\) on \(H\)) were used only for single-point energy calculations and can be considered to be complete enough to yield reliable relative energies and reaction energies at the CCSD(T) level.

III. DISCUSSION OF RESULTS

In the following, we will discuss first, aspects of methodology and results for the different \(C_2H_2\) structures (Tables I–V). The accuracy of the CCSD(T) calculations using different basis sets will be assessed and the necessity to use such high-level methods is shown. Further, a comparison to results in the literature is made. Finally, selected reaction energies are presented.

A. \(C_3H_2\) minima

Table I shows relative energies of four minima (Fig. 1) and five saddle points (Fig. 2) of \(C_3H_2\) on the triplet surface. The most stable isomer is the so-called propargylene, denoted \(C_{2h}^\mathrm{min}\), which is energetically favored by more than 134.9 kJ/mol [CCSD(T)/QZ2P/ZPE] vs other isomers of \(C_3H_2\). Its \(C_2\) symmetry [Fig. 1(a)] agrees with results of a recent Fourier transform IR (FTIR) experiment, in which the assignment was made by \(^{13}\)C-labeling studies in argon matrices.51 The CCC angle is almost linear, 171.9°, and the torsion angle of the two \(H\) atoms is 88.0° [CCSD(T)/TZP, Table V]. The \(C–C\) bond length of 127.9 pm has to be compared to bond lengths of 120.9 pm for a typical triple bond (\(C_2H_2\)) and 132.8 pm for a double bond (\(C_2H_4\)) within the same approach. The lowest vibrational frequency of 215 cm\(^{-1}\) [CCSD(T)/TZP; Table II] indicates the floppiness of the structure.

Similarly the second-most stable isomer, vinylidenecar-
bene [denoted $C_{2v}$; Fig. 1(b)], with $C_{2v}$ symmetry is also floppy with a vibrational frequency of 222 cm$^{-1}$ [CCSD(T)/TZP]. The C–C bond length to the carbon holding the two hydrogen atoms is 136.9 pm whereas the other C–C bond is much shorter, at 123.8 pm. C–H bond distances of 108.1 pm compare to 108.4 pm as found in C$_2$H$_4$ [CCSD(T)/TZP].

Cyclopropenylidene $c$-C$_3$H$_2$, denoted $C_{1cyclic}^c$, shows no symmetry [Fig. 1(c)] and is 172.4 kJ/mol [CCSD(T)/QZ2P/ZPE] higher in energy compared to the most stable isomer, $C_{2lin}^l$. In contrast to all other structures its lowest vibrational frequency is much higher, 599.1 cm$^{-1}$ [CCSD(T)/TZP]. One hydrogen is placed almost in the plane of the carbon cycle (out-of-plane angle only 0.2$^\circ$), whereas the other H atom is distorted by 46.1$^\circ$ out of the CCC plane [CCSD(T)/TZP]. C–C distances range between 130.4 and 155.1 pm.

Another isomer ($C_s^{trans}$), trans-propenediylidene [Fig. 1(d); trans with respect to the H atoms] shows $C_s$ symmetry and is 250.8 kJ/mol [CCSD(T)/QZ2P/ZPE] less stable than the most stable C$_3$H$_2$ structure. C–C bond lengths of 134.9 and 139.2 pm [CCSD(T)/TZP] have to be compared to the

<table>
<thead>
<tr>
<th>Method/Basis</th>
<th>$C_{2lin}^l$ (Hartree)</th>
<th>$C_{2v}$</th>
<th>$C_{1cyclic}^c$</th>
<th>$C_s^{trans}$</th>
<th>$C_{2lin}^l$</th>
<th>$D_{a+b}$</th>
<th>$C_{2cyclic}^c$</th>
<th>$C_{2cyclic}^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCSD(T)/DZP</td>
<td>-115.042 385</td>
<td>114.3</td>
<td>161.1</td>
<td>238.9</td>
<td>0.5</td>
<td>0.8</td>
<td>3.7</td>
<td>175.5</td>
</tr>
<tr>
<td>CCSD(T)/TZP</td>
<td>-115.106 431</td>
<td>118.8</td>
<td>164.6</td>
<td>240.9</td>
<td>0.5</td>
<td>0.7</td>
<td>2.9</td>
<td>177.4</td>
</tr>
<tr>
<td>CCSD(T)/TZPP</td>
<td>-115.173 001</td>
<td>122.2</td>
<td>160.5</td>
<td>243.1</td>
<td>-0.1</td>
<td>0.1</td>
<td>0.4</td>
<td>173.2</td>
</tr>
<tr>
<td>CCSD(T)/QZ2P</td>
<td>-115.199 261</td>
<td>124.4</td>
<td>160.9</td>
<td>247.1</td>
<td>-0.3</td>
<td>0.0</td>
<td>-0.4</td>
<td>173.6</td>
</tr>
<tr>
<td>CCSD(T)/QZ2P/ZPE</td>
<td>-115.173 119</td>
<td>134.9</td>
<td>172.4</td>
<td>250.8</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>SCF/QZ2P</td>
<td>-114.677 750</td>
<td>104.0</td>
<td>229.6</td>
<td>202.9</td>
<td>0.7</td>
<td>1.0</td>
<td>4.1</td>
<td>238.7</td>
</tr>
<tr>
<td>MP2/QZ2P</td>
<td>-115.129 589</td>
<td>124.5</td>
<td>104.3</td>
<td>256.3</td>
<td>-2.6</td>
<td>-0.8</td>
<td>-5.2</td>
<td>141.5</td>
</tr>
<tr>
<td>CCSD/QZ2P</td>
<td>-115.176 692</td>
<td>122.1</td>
<td>169.3</td>
<td>241.9</td>
<td>-0.5</td>
<td>-0.1</td>
<td>-0.9</td>
<td>182.0</td>
</tr>
</tbody>
</table>

*Second derivatives with respect to nuclear displacements as computed at the CCSD(T)/TZP level. Imaginary frequencies described in text and Tables III and IV.

*b Single-point calculations at the CCSD(T)/TZP geometry.

*c Zero-point vibrational energies included as computed at the CCSD(T)/TZP level (only included for minima).

FIG. 1. Computed minima of C$_3$H$_2$. Relative energies with respect to the most stable isomer C$_{2lin}^l$ are listed in kJ/mol [CCSD(T)/QZ2P/ZPE]. Large grey circles denote carbon atoms and smaller white circles denote hydrogen.
B. Aspects of methodology and connection to literature data

Before discussing the different saddle points found for C$_3$H$_2$ (Fig. 2 and Tables I, III, and IV) it is important to assess the accuracy which might be expected for relative energies, structural properties, and vibrational frequencies of this molecule. For this purpose we use the linear structure with D$_{\infty h}$ symmetry of C$_3$H$_2$, as it allows us to investigate basis set effects not only on structural parameters and vibrational frequencies, but—as it is a saddle point on the hypersurface—effects on imaginary frequencies can be observed as well. The high point group symmetry of the D$_{\infty h}$ structure permits the use of quite large basis sets.

The already mentioned typical spin contamination of C$_3$H$_2$ SCF wave functions suggests the necessity of using high-level ab initio methods like the CC scheme. This is supported by analyzing relative energies of C$_3$H$_2$ structures shown in Table I; the most drastic example represents the cyclic isomer C$_3^{\text{cyclic}}$ for which energy differences relative to the most stable isomer C$_3^{\text{lin}}$ change from 229.6 kJ/mol (SCF), over 104.3 kJ/mol (MP2), to 169.3 kJ/mol (CCSD) and finally to our most accurate result 160.9 kJ/mol [CCSD(T)] using the QZ2P basis. This trend reflects the well known overestimation of correlation effects at the MP2 level and shows the inappropriateness of the SCF and MP2 method for

\begin{table}[h]
\centering
\begin{tabular}{llllll}
\hline
C$_3^{\text{lin}}$ & DZP & TZP & C$_3^{\text{cyclic}}$ & DZP & TZP \\
\hline
b & 221.1 & (95.3) & 215.2 & (96.8) & b$_1$ & 210.6 & (0.4) & 221.8 & (0.4) \\
  & 332.0 & (1.3) & 324.5 & (3.2) & b$_2$ & 391.7 & (1.9) & 374.9 & (2.7) \\
  & 396.5 & (10.2) & 391.8 & (14.2) & b$_3$ & 661.5 & (50.4) & 649.5 & (61.2) \\
  & 401.1 & (21.3) & 398.8 & (27.0) & b$_4$ & 992.1 & (5.5) & 996.9 & (7.0) \\
  & 469.3 & (38.4) & 450.3 & (47.1) & a$_1$ & 1107.3 & (5.7) & 1094.1 & (5.6) \\
  & 1267.1 & (0.0) & 1257.4 & (0.1) & a$_2$ & 1475.5 & (0.0) & 1484.6 & (0.0) \\
  & 1622.6 & (16.2) & 1610.5 & (12.8) & a$_3$ & 1962.9 & (1.0) & 1953.5 & (0.6) \\
  & 3410.8 & (57.6) & 3409.8 & (60.4) & a$_4$ & 3186.9 & (2.1) & 3170.5 & (3.3) \\
  & 3416.7 & (6.0) & 3416.4 & (4.4) & b$_1$ & 3297.8 & (0.2) & 3272.1 & (0.4) \\
\hline
C$_3^{\text{cyclic}}$ & DZP & TZP & C$_3^{\text{trans}}$ & DZP & TZP \\
\hline
a & 591.0 & (46.1) & 599.1 & (55.3) & a' & 178.6 & (10.6) & 187.2 & (11.9) \\
  & 786.1 & (22.3) & 720.1 & (6.5) & a'' & 149.7 & (23.2) & 412.8 & (27.3) \\
  & 934.1 & (12.1) & 919.6 & (72.4) & a'' & 690.3 & (32.5) & 674.3 & (38.9) \\
  & 959.9 & (29.3) & 934.6 & (1.4) & a' & 840.0 & (26.0) & 834.7 & (32.2) \\
  & 1015.2 & (37.9) & 995.4 & (18.6) & a' & 1052.4 & (1.9) & 1063.1 & (1.4) \\
  & 1153.2 & (16.3) & 1089.1 & (28.3) & a' & 1193.9 & (0.1) & 1171.8 & (2.3) \\
  & 1686.5 & (2.8) & 1672.9 & (4.9) & a' & 1383.1 & (50.9) & 1355.4 & (47.5) \\
  & 3136.2 & (25.5) & 3125.0 & (22.3) & a' & 3120.2 & (14.3) & 3115.2 & (13.1) \\
  & 3347.3 & (13.6) & 3343.8 & (15.6) & a' & 3279.6 & (1.2) & 3268.5 & (1.1) \\
\hline
\end{tabular}
\caption{Harmonic vibrational frequencies computed at the CCSD(T) level using DZP and TZP bases, respectively, of C$_3$H$_2$ isomers depicted in Fig. 1. Symmetry of modes, wave numbers (cm$^{-1}$), and IR intensities (km/mol; in parentheses) are listed.}
\end{table}
TABLE III. Harmonic vibrational frequencies computed at the CCSD(T)/TZP level of C₃H₂ saddle points depicted in Fig. 2. Symmetry of modes, wave numbers (cm⁻¹), and IR intensities (km/mol; in parentheses) are listed.

<table>
<thead>
<tr>
<th>Symmetry</th>
<th>C^{lin}_{r}</th>
<th>C^{lin}_{\theta}</th>
<th>C^{cyclic}_{T}</th>
<th>C^{cyclic}_{\theta}</th>
</tr>
</thead>
<tbody>
<tr>
<td>a'</td>
<td>212.0i</td>
<td>258.8i</td>
<td>701.5i</td>
<td>761.7i</td>
</tr>
<tr>
<td>a</td>
<td>232.2</td>
<td>177.2i</td>
<td>732.8</td>
<td>620.1i</td>
</tr>
<tr>
<td>a''</td>
<td>388.5</td>
<td>394.2</td>
<td>785.8</td>
<td>722.7</td>
</tr>
<tr>
<td>a''''</td>
<td>395.0</td>
<td>400.0</td>
<td>809.6</td>
<td>782.7</td>
</tr>
<tr>
<td>a'''</td>
<td>410.0</td>
<td>453.5</td>
<td>873.3</td>
<td>816.7</td>
</tr>
<tr>
<td>a''''''</td>
<td>1241.1</td>
<td>1262.9</td>
<td>1071.7</td>
<td>840.0</td>
</tr>
<tr>
<td>a''''''''</td>
<td>1645.0</td>
<td>1608.9</td>
<td>1556.1</td>
<td>1574.5</td>
</tr>
<tr>
<td>a''''''''''</td>
<td>3388.9</td>
<td>3421.8</td>
<td>3624.8</td>
<td>3365.2</td>
</tr>
<tr>
<td>a''''''''''''</td>
<td>3456.7</td>
<td>3430.5</td>
<td>3272.0</td>
<td>3395.5</td>
</tr>
</tbody>
</table>

C₃H₂. The largest T¹_max and T²_max CC amplitudes range typically around 0.08 and 0.06, respectively.

A recent study of C₃H₂ by Takahashi and Yamashita²² using the SCF and MP2 approximations for the computation of energetics and vibrational frequencies of C₃H₂ also shows inadequacies of these methods for the treatment of this molecule. For example, their MP2/6-31 G(d,p) calculations on a cyclic triplet C₁ isomer of C₃H₂ lead to an unphysical vibrational frequency of 5714 cm⁻¹.²² The occurrence of completely unreasonable force constants is a well known problem at lower levels of theory like SCF and MP2, especially in case of open-shell systems with low-lying excited states.²⁶,²⁸ Such defects within the optimized SCF wave function are only very slowly compensated in a simple perturbation expansion like in many-body perturbation theory [MBPT(n)] [Ref. 53] with higher order n. This is further support for using CC methods.

Using SCF and MP2 Takahashi and Yamashita²² find a different most stable isomer of C₃H₂ on the triplet surface compared to our CCSD(T) study. Instead of the almost linear structure with C₂ symmetry we calculated (C^{lin}_{2}; Fig. 1), whose symmetry agrees with results of recent FTIR experiments,²¹ they find another almost linear structure, C^{cyclic}_{2} (Fig. 2), as the most stable isomer of the C₃H₂ triplet surface. Our CCSD(T)/QZ2P results characterize this structure to be virtually isoenergetic with the most stable isomer of our study (C^{cyclic}_{2}), but it is a saddle point at the CCSD(T) level (Table III). Our calculation supports the C₂ symmetry found for the linear isomer at the QCISD(T)/6-31G* level by Herges et al.²¹ The second most stable isomer of C₃H₂ (tripllet), vinylidenecarbene (C^{trans}_{2}), was not considered in the study of Takahashi and Yamashita.²² Further, trans-propenediylidene (C^{trans}_{2}), found as a saddle point by Takahashi and Yamashita²² at the MP2 level (structure 2a in Ref. 22), is a minimum on the CCSD(T) hypersurface (Table II).

Another serious problem of ab initio studies are instabilities and symmetry-breaking in the zeroth-order SCF wave function, which we will discuss extensively for the C₂H molecule. Related to this often underestimated difficulty, the published cis-propenediylidene (cis with respect to the H atoms; structure 2b in Ref. 22) structure of C₃H₂ found as a local minimum by Takahashi and Yamashita²² is based on an unstable HF wave function (for which the energy was listed as −114.556 17 Hartree, Table III in Ref. 22), although in this case a stable HF solution within the same symmetry can be found by running along the mode of the negative eigenvalue describing the corresponding orbital rotations. In such a case MP2 enhances the error introduced in the zeroth-order wave function as reflected by even bigger energy differences between using unstable and stable wave functions (whereas CC methods tend to largely overcome such errors). This will be discussed more extensively for symmetry-broken HF wave functions in context of the C₂H molecule. The cis-propenediylidene structure of C₃H₂ found using the unstable

TABLE IV. Comparison of equilibrium geometries (bond length, r, in pm), harmonic vibrational frequencies (cm⁻¹), IR intensities (km/mol, in parentheses), and zero-point vibrational energies, ZPE, (kJ/mol) as computed at the CCSD(T) level using various basis sets for the linear structure (D_2h) of C₃H₂ shown in Fig. 2(c).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>DZP</th>
<th>TZP</th>
<th>TZPP</th>
<th>QZ2P</th>
</tr>
</thead>
<tbody>
<tr>
<td>r(CC)</td>
<td>128.2</td>
<td>127.4</td>
<td>127.2</td>
<td>127.0</td>
</tr>
<tr>
<td>r(CH)</td>
<td>107.3</td>
<td>106.3</td>
<td>106.1</td>
<td>106.1</td>
</tr>
<tr>
<td>ω(σ')</td>
<td>398.1i</td>
<td>374.8i</td>
<td>247.8i</td>
<td>196.2i</td>
</tr>
<tr>
<td>ω(π')</td>
<td>302.1i</td>
<td>308.8i</td>
<td>198.2i</td>
<td>149.3i</td>
</tr>
<tr>
<td>ω(σ')</td>
<td>397.1</td>
<td>390.6</td>
<td>408.7</td>
<td>414.1</td>
</tr>
<tr>
<td>ω(π')</td>
<td>1282.6</td>
<td>1274.5</td>
<td>1275.8</td>
<td>1283.6</td>
</tr>
<tr>
<td>ω(σ')</td>
<td>1626.7</td>
<td>1617.5</td>
<td>1633.1</td>
<td>1647.1</td>
</tr>
<tr>
<td>ω(π')</td>
<td>3457.4</td>
<td>3449.6</td>
<td>3437.6</td>
<td>3469.3</td>
</tr>
<tr>
<td>ω(σ')</td>
<td>3470.0</td>
<td>3463.1</td>
<td>3448.8</td>
<td>3481.8</td>
</tr>
<tr>
<td>ZPE</td>
<td>63.59</td>
<td>63.32</td>
<td>63.48</td>
<td>64.06</td>
</tr>
</tbody>
</table>
SCF wave function by Takahashi and Yamashita,\textsuperscript{22} was not located using the CCSD(T) method (using of course a stable zeroth-order wave function), which lead on several optimizations from different starting points to a linear structure. However, this does not safely exclude the existence of this structure as a stationary point, it can only be concluded that if its surrounding hypersurface is probably very flat.

After we have shown the problematic nature of low level methods for the computational treatment of C$_3$H$_2$, it is important to continue the evaluation of accuracies which might be expected at the CCSD(T) level for different properties and aspects of this system. The CCSD(T)/QZ2P relative energies are expected to be highly accurate and basis set effects to be small, as indicated by a change of less than 4 kJ/mol in increasing the basis from TZPP to QZ2P. Due to its high point group symmetry the linear structure D$_{lin}$ allows us to investigate basis set effects on both structural properties and vibrational frequencies. The data listed in Table IV show the excellent agreement of structural constants computed at the CCSD(T)/TZP level as compared to data obtained using the bigger basis sets TZPP and QZ2P, with deviations of less than 0.4 pm. Vibrational frequencies [CCSD(T)/TZP] differ by less than 30 cm$^{-1}$ using basis sets up to QZ2P, whereas imaginary frequencies are more sensitive to basis set effects (Table IV). However, no change in the number of imaginary frequencies was observed. The good accuracy achieved for vibrational frequencies leads to only small changes in zero-point vibrational energies (ZPE) with deviations of less than 1 kJ/mol (Table IV).

The examination of relative energies including zero-point vibrational energies for the C$_3$H$_2$ triplet structures, $C^0_{lin}$ (which is almost isoenergetic with the saddle point $C^0_s$ found in Ref. 22 as a minimum) and structure $C^0_{cycl}$, which we might compare to the \textit{ab initio} study of Takahashi and Yamashita,\textsuperscript{22} reveals a surprisingly good coincidence of their MP4/6-311G(d,p)//MP2/6-31G(d,p) value (MP4 single-point at MP2 structure) with our CCSD(T)/QZ2P/ZPE data. However, the unexpected small deviation of only slightly more than 10 kJ/mol is a result of a fortuitous cancellation of errors in their study;\textsuperscript{22} they used the zero-point energy which includes the unreasonable vibrational frequency of 5714 cm$^{-1}$, mentioned earlier, for the calculation of relative energies [the ZPE effect on the relative energy of these two structures is 38 kJ/mol Ref. 22].

### C. Saddle points of C$_3$H$_2$

Three almost linear structures, $C^0_{s}$, $C^2_{s}$, and the exactly linear one $D_{sh}$ [Figs. 2(a)–2(c)], are saddle points [CCSD(T)/TZP] with one and two imaginary frequencies, respectively (Tables III and IV; for $D_{sh}$ the two imaginary modes are degenerate). The first one, $C^0_{s}$, has been previously found to be a minimum at lower levels of theory (e.g., Ref. 22). However, all these structures are virtually isoenergetic [within 1 kJ/mol, CCSD(T)/QZ2P] with the most stable minimum of C$_3$H$_2$, $C^0_{lin}$. For example, the energy difference between the most stable isomer $C^0_{lin}$ and the saddle point $C^0_{s}$ is extremely small (see Table I). The occurrence of negative values (−0.3 to +0.5 kJ/mol) within the CCSD(T)/TZP and CCSD(T)/QZ2P approach is attributed to the non-optimized structures used in the single-point approach [at the CCSD(T)/TZP structures], a fact which has been checked using the TZPP basis (nonoptimized vs optimized). These small energy differences reflect again the flatness of the hypersurface in the vicinity of the most stable C$_3$H$_2$ isomer, $C^0_{lin}$, which is indicated as well by its lowest vibrational frequency of 215.2 cm$^{-1}$ [CCSD(T)/TZP].

A cyclic C$_3$H$_2$ structure with both hydrogen atoms attached to the same carbon has not been investigated in this study, as it is considered to be unimportant for the reaction of C$^1(P)$ with C$_3$H$_2$,\textsuperscript{14} where the two H atoms are bound to different carbons and hydrogen migration can not take place within the lifetime of this collision complex, see Ref. 9. Its energy difference with respect to the the most stable linear triplet isomer of C$_3$H$_2$ was found by Jonas et al.\textsuperscript{20} to be 256 kJ/mol, as obtained by using a spin-projected MP4 method at MP2 geometries [MP4/6-311G(2df)//MP2/6-31G(d)].

The cyclic isomer of C$_3$H$_2$, $C^0_{cycl}$, which is a minimum on the CCSD(T)/TZP hypersurface, has been derived by distortion from higher symmetry analogs, $C^0_{cycl}$ and $C^0_{cycl}$, showing one and two imaginary frequencies, respectively [Figs. 2(d) and 2(e), and Table III]. The energy gain by the distortion from the $C^0_{cycl}$ structure with the highest symmetry is 26.4 kJ/mol [CCSD(T)/QZ2P, Table I].

---

**Table V. Structural data of C$_3$H$_2$ isomers (Figs. 1 and 2) as computed at the CCSD(T)/TZP level.**

<table>
<thead>
<tr>
<th>Isomer$^a$</th>
<th>Structural data</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C^0_{lin}$</td>
<td>(C$^<em>$–C)=127.9 (C–H)=106.7 (C,C$^</em>$,C’)=171.9°</td>
</tr>
<tr>
<td>(C$^<em>$,C,H)=156.5° (H–C$^</em>$–C–H)=88.0°</td>
<td></td>
</tr>
<tr>
<td>$C^2_{s}$</td>
<td>(C$^*$,H,C)=136.9 (C–H,C)=123.8 (C–H)=108.1</td>
</tr>
<tr>
<td>(C–C,H,C)=120.5°</td>
<td></td>
</tr>
<tr>
<td>$C^0_{cycl}$</td>
<td>(C$^*$–C)=144.8 (C–C)=130.4 (C–H,C)=151.5</td>
</tr>
<tr>
<td>(C–C,H,C)=141.1° (H,C,C,C,H)=161°</td>
<td></td>
</tr>
<tr>
<td>$C^0_{s}$</td>
<td>(C$^*$–C)=134.9 (C–C)=139.2 (C–H,C)=109.2</td>
</tr>
<tr>
<td>(C–C,H,C)=134.1°</td>
<td></td>
</tr>
<tr>
<td>$C^0_{lin}$</td>
<td>(C$^*$–C)=129.6 (C–C)=126.2 (C–H)=106.9</td>
</tr>
<tr>
<td>(C–C,H,C)=172.6°</td>
<td></td>
</tr>
<tr>
<td>$C^0_{cycl}$</td>
<td>(C$^*$–C)=135.5 (C–C)=159.3 (C–H)=107.7</td>
</tr>
<tr>
<td>(H,C,C$^*$)=146.4°</td>
<td></td>
</tr>
<tr>
<td>$C^0_{cycl}$</td>
<td>(C$^*$–C)=134.9 (C–C)=159.2 (H,C)=106.9</td>
</tr>
<tr>
<td>(H,C,C$^*$)=144.8°</td>
<td></td>
</tr>
</tbody>
</table>

$^a$For structures with a C$_2$-symmetry axis (and not all C atoms situated on this axis, as in the $C^0_{s}$ structure, Fig. 1) the symmetry-unique C atom is labeled C$. C^*$ is the symmetry-corresponding atom of C. If no such symmetry is found, C and H atoms, respectively, are numbered from left to the right in Figs. 1 and 2 (for the cyclic isomer $C^0_{cycl}$, atom C2 is characterized by holding no H atoms). Notation: (C$^*$–C)=127.9 denotes bond length in pm, (C$^*$,C$^*$,C)=171.9° denotes bond angle with apical atom C$^*$, (H,C,C,C,H)=46.1° stands for the out of plane angle between the bond H1–C1 and the plane C2–C1–C3, and (H–C$^*$–C–H$’$)=88.0° stands for the torsion angle H–C–C$’$–H$’$.}
D. \( \text{C}_3\text{H} \)

Before we discuss computed reaction energies of the neutral–neutral reaction (1) presented in the introduction of this paper, we need to investigate the product of the reaction, \( \text{C}_3\text{H} \). This study focuses on its energetics, and, as mentioned before, it is not our intention to treat the still open issue of what symmetry has to be assigned to the equilibrium geometry of the linear isomer using \textit{ab initio} methods. This question has only negligible influence on reaction energies, which are our only concern about \( \text{C}_3\text{H} \) in the present study.

Table VI lists relative energies of three \( \text{C}_3\text{H} \) structures; a cyclic \( \text{c-C}_3\text{H} \) (\( C_{2v} \) symmetry), a linear \( \text{l-C}_3\text{H} \) (\( C_{\infty v} \)), and a bent \( \text{b-C}_3\text{H} \) (trans) form with \( C_2 \) symmetry (see Fig. 3). The \( \text{c-C}_3\text{H} \) isomer is 13.0 kJ/mol more stable than the linear form \( \text{l-C}_3\text{H} \) at the CCSD(T)/cc-pVQZ level. The change in increasing the basis from QZ2P to cc-pVQZ is less than 1 kJ/mol, indicating the sufficient size of the employed basis sets. The two linear structures \( \text{l-C}_3\text{H} \) and \( \text{b-C}_3\text{H} \) are virtually isoenergetic with an energetic difference of less than 1 kJ/mol [CCSD(T)/cc-pVQZ].

Comparison of our most reliable CCSD(T) data with results at lower levels of theory like SCF and MP2 reveals, as expected, an even much more pronounced deficiency of these methods for the treatment of \( \text{C}_3\text{H} \) than for \( \text{C}_3\text{H}_2 \). SCF favors the linear structures by more than 30 kJ/mol, whereas for MP2 a strong overestimation of correlation effects can be observed, preferring the linear structure by more than 80 kJ/mol (Table VI). The problem is reflected by high maximal amplitudes \( T^{\text{max}}_1 \) and \( T^{\text{max}}_2 \); although reasonable values of 0.04–0.05 and 0.06–0.07, respectively, occur for \( \text{c-C}_3\text{H} \), much higher values of \( T^{\text{max}}_1=0.17–0.22 \) and \( T^{\text{max}}_2=0.08–0.10 \) are observed for linear structures of \( \text{C}_3\text{H} \). This is closely related to the problem that for \( \text{c-C}_3\text{H} \) stable HF solutions can be found using basis sets like TZP and larger, whereas for the linear structures no stable SCF wave function within the true symmetry (always \( D_{2b} \) and subgroups) can be found. However, energy differences between using the true symmetry wave function and the stable \( C_1 \) zeroth-order wave function are very small (negative sign if \( C_1 \) solution is lower in energy; basis TZP),

\[
\text{l-C}_3\text{H} \quad -0.4/+/5.0/+0.3+0.2 \quad [\text{SCF}/\text{MP2}/\text{CCSD}/\text{CCSD}(\text{T})],
\]

\[
\text{b-C}_3\text{H} \quad -0.3/+4.4/+0.2+0.1 \quad [\text{SCF}/\text{MP2}/\text{CCSD}/\text{CCSD}(\text{T})].
\]

![Fig. 3. Considered \( \text{C}_3\text{H} \) structures and energy differences with respect to the most stable isomer \( \text{c-C}_3\text{H} \) (\( C_{2v} \) symmetry) in kJ/mol [CCSD(T)/cc-pVQZ/ZPE].](image-url)
TABLE VII. Equilibrium geometry, harmonic vibrational frequencies (cm\(^{-1}\)), IR intensities (km/mol, in parentheses), and zero-point vibrational energies, ZPE, (kJ/mol) of \( b\)-C\(_3\)H (\( C_1 \) point group; Fig. 3). Stability difficulties in the zeroth-order wave function are discussed in text. Labeling used for internal coordinates as described in Table V.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>CCSD(T)/TZP</th>
</tr>
</thead>
<tbody>
<tr>
<td>( r(\text{H–C}1) )</td>
<td>107.2 pm</td>
</tr>
<tr>
<td>( r(\text{C}1–\text{C}2) )</td>
<td>125.3 pm</td>
</tr>
<tr>
<td>( r(\text{C}2–\text{C}3) )</td>
<td>133.6 pm</td>
</tr>
<tr>
<td>( \angle(\text{H}1\text{C}1\text{C}2) )</td>
<td>156.5°</td>
</tr>
<tr>
<td>( \angle(\text{C}1\text{C}2\text{C}3) )</td>
<td>174.0°</td>
</tr>
<tr>
<td>( \omega(\text{a}^\prime) )</td>
<td>207.8</td>
</tr>
<tr>
<td>( \omega(\text{a}^\prime') )</td>
<td>351.2</td>
</tr>
<tr>
<td>( \omega(\text{a}^\prime') )</td>
<td>368.6</td>
</tr>
<tr>
<td>( \omega(\text{b}^\prime') )</td>
<td>1170.1</td>
</tr>
<tr>
<td>( \omega(\text{b}^\prime') )</td>
<td>1876.1</td>
</tr>
<tr>
<td>( \omega(\text{b}^\prime') )</td>
<td>3379.9</td>
</tr>
<tr>
<td>ZPE</td>
<td>43.98</td>
</tr>
</tbody>
</table>

As mentioned before such energy differences between a true symmetry and a symmetry-broken solution are increased by the MP2 method, whereas the CCSD(T) method compensates the already small energy difference within the SCF approach to 0.2 and 0.1 kJ/mol, respectively. This shows that energetics are only slightly influenced by such effects.

In addition we performed Brueckner CC calculations B-CCD, known for their ability to be often efficient for the treatment of symmetry-breaking problems and the occurrence of larger \( T_{\text{max}} \) amplitudes.\(^{33,44}\) The change in relative energies as compared to the CCSD data (basis Q2ZP) is less than 2 kJ/mol, confirming the validity of the conventional CCSD approach for this molecule.

Our relative energies are in excellent agreement with multireference CI (MRCI) data including the Davidson correction published by Takahashi and Yamashita\(^ {22} \) using a valence triple zeta basis. They found the \( c\)-C\(_3\)H isomer to be favored by 8.6 kJ/mol vs \( l\)-C\(_3\)H, and the slightly bent one, \( b\)-C\(_3\)H, to be separated by less than 2 kJ/mol from the exactly linear \( l\)-C\(_3\)H structure.

Structural data for \( c\)-C\(_3\)H [CCSD(T)/TZP] shows excellent agreement with both theoretical and experimental results. Bond lengths differ by less than 0.3 pm from data obtained by microwave experiments.\(^ {30} \) For the cyclic isomer of C\(_3\)H we use the most accurate zero-point vibrational energy (49.94 kJ/mol) of Stanton,\(^ {26} \) obtained by applying the EOMIP-CCSD method.

Problems are more difficult for linear structures of C\(_3\)H considered in this study, as no stable wave function within the correct symmetry restrictions can be found. However, as mentioned above, energetic differences in using symmetry-restricted and stable \( C_1 \) zeroth-order wave functions are negligible. It shows that the influence of optimizing the structures of \( l\)-C\(_3\)H (\( C_{\infty v} \) point group) and \( b\)-C\(_3\)H (\( C_1 \) symmetry) using these different zeroth-order wave functions is extremely small.

In Table VII vibrational frequencies for the \( b\)-C\(_3\)H structure at the CCSD(T)/TZP level are listed. These results were obtained using the symmetry-constrained \( C_s \) wave function, as one would expect in general, that the use of an optimized wave function which does not belong to any irreducible representation at a point of high symmetry leads to an artificial cusp in the energy and to a nondifferentiable function (see Refs. 43, 54). The use of the symmetry-constraint \( C_s \) wave function is further supported by the agreement of the corresponding numerical and analytical frequencies at the SCF level. Our zero-point vibrational energy for \( b\)-C\(_3\)H, which is our main interest in this study, of 44.0 kJ/mol is in good agreement with complete active space SCF (CASSCF) studies\(^ {22,27,31} \) with a deviation of less than 2.7 kJ/mol.

An experimental study of \(^{13}\)C isotopomers for C\(_3\)H in a solid Ar matrix using FTIR spectroscopy\(^ {32} \) assigned absorptions at 1159.8 and 1824.7 cm\(^{-1}\) to the linear form of this radical. The agreement of these observed frequencies with our corresponding CCSD(T)/TZP data for the almost linear isomer \( b\)-C\(_3\)H of 1170.1 and 1876.1 cm\(^{-1}\) (Table VII) is excellent, considering theoretical difficulties encountered for \( l\)-C\(_3\)H and the slightly distorted linear conformation.

We use the zero-point vibrational energy of \( b\)-C\(_3\)H as computed at the CCSD(T)/TZP level for the correction of reaction energies of both \( l\)-C\(_3\)H and \( b\)-C\(_3\)H, as these structures are very similar. The calculation of vibrational frequencies for \( l\)-C\(_3\)H remains an open issue, which we do not address in this publication. The difficulties encountered for the \( l\)-C\(_3\)H structure are reflected by three CASSCF studies; one by Takahashi et al.\(^ {22} \) leading to a minimum with a lowest vibrational frequency of 325 cm\(^{-1}\), the other by Aoki et al.\(^ {27} \) finding \( l\)-C\(_3\)H as a transition state (355 cm\(^{-1}\)), and a third CASSCF study of Kanada et al.\(^ {31} \) which yields as well an imaginary frequency of 247 cm\(^{-1}\). However, the influence of basis set effects has not yet been investigated, so that no final conclusion concerning vibrational frequencies of the exactly linear structure can be drawn.

A better understanding of the linear C\(_3\)H structure can be gained by comparing experimental to theoretical structure data. A recent microwave study of Kanada et al.\(^ {31} \) finds in combination with theoretical data a quasilinear structure. This is strongly supported by an exceptionally short C–H bond of 101.7 pm found experimentally,\(^ {31} \) whereas the corresponding bond length of the exactly linear \( l\)-C\(_3\)H has been computed to be significantly larger [106.5 pm at CCSD(T)/TZP]. The short distance may be interpreted as due to the very low lying vibronic state corresponding to the CCH bending mode as a result from a large Renner–Teller effect (e.g., Ref. 55). Further, it is possible that the bent isomer, \( b\)-C\(_3\)H, represents the equilibrium structure. Projection of the C–H bond length on to the \( z \)–axis (for simplicity we just take the C1–C3 axis) leads to a bond length of 100.8 pm [CCSD(T)/TZP] compared to 101.7 pm found in the experiment,\(^ {31} \) which is within error bars for the angle and bond length used for projection. As for the other computed bond lengths (here without projection, as the CCC bond angle is with 174.0° quite close to a linear form), they compare well to experimental values. The computed C–C distance to the terminal CH group is 125.3 pm, which is much shorter and closer to a typical triple bond than the second
TABLE VIII. Selected reaction energies in kJ/mol computed at CCSD(T)/TZP geometries.a

<table>
<thead>
<tr>
<th>Method/Basis</th>
<th>$\Delta E_{R_1}$</th>
<th>$\Delta E_{R_2}$</th>
<th>$\Delta E_{R_3}$</th>
<th>$\Delta E_{R_4}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCSD(T)/TZP</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>CCSD(T)/TZP</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>CCSD(T)/QZ2P</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>CCSD(T)/cc-pVQZ</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>CCSD(T)/QZ2P/ZPE</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>CCSD(T)/cc-pVQZ/ZPE</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>SCF/QZ2P</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>SCF/cc-pVQZ</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>MP2/QZ2P</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>MP2/cc-pVQZ</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>CCSD/QZ2P</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
<tr>
<td>CCSD/cc-pVQZ</td>
<td>33.1</td>
<td>39.4</td>
<td>45.0</td>
<td>52.2</td>
</tr>
</tbody>
</table>

a Full point-group symmetries (D3h and subgroups) are used for the wave functions (see discussion in text).

The carbon–carbon bond length of 133.6 pm at the CCSD(T)/TZP level (Table VII). These data are in good agreement with experimental values of 125.4 and 132.6 pm, respectively.31 For comparison the corresponding C–C bond lengths of the exactly linear structure $l$-C3H are 124.3 and 134.7 pm, respectively, differing by less than 1 pm from those in $b$-C3H.

E. Reaction energies

In Table VIII reaction energies for the neutral–neutral reaction of $C(3P)$ with C2H2 (1) are displayed. In addition to our most reliable CCSD(T) data, we also list in this table data computed at lower levels of theory, which confirms the already discussed deficiencies of the SCF and MP2 methods for the treatment of C2H2 and C3H.

The initial formation of C2H2 collision complexes is exothermic by 385.4 kJ/mol [CCSD(T)/QZ2P/ZPE]. The whole reaction to the cyclic isomer, $c$-C3H, is computed to be exothermic by 8.6 kJ/mol [CCSD(T)/cc-pVQZ/ZPE]. The change of 18 kJ/mol for this energy in increasing the basis from TZP to cc-pVQZ shows the importance of using an adequate basis set. The cc-pVQZ basis is expected to yield reaction energies for this reaction that are accurate to a few kJ/mol at the CCSD(T) level. As mentioned before, the linear form of C3H is only slightly higher in energy compared to $c$-C3H leading to a still slightly exothermic reaction energy (Table VIII).

IV. EXPERIMENTAL IMPLICATIONS

In this section, we briefly discuss the experimental implications of our ab initio calculations for the reaction of ground state carbon, C(3P), with acetylene, C2H2, as studied via the crossed molecular beams technique described elsewhere.8,9 This approach allows insight into chemical dynamics of the reaction and reveals information on reaction intermediates as well as C3H isomers, when the experimental data are compared to what is expected based on our ab initio structures and energetics. For more complete discussions, see Refs. 9, 10.

Reaction dynamics inferred from the experimental data together with our ab initio calculations suggest two micro-channels on the triplet ground state surface initiated by addition of C($3P$) either to one acetylenic carbon to form trans-propargenediylidene, $C\;_{trans}$, or to two carbon atoms to yield $c$-C3H2, cyclopropenylidene ($C\;_c^{cyclic}$). Propargenediylidene itself undergoes $[2,3]$-H-migration to propargylene, followed by the final C–H bond rupture to $l$-C3H and H. Direct stripping dynamics contribute to the second microchannel to form $c$-C3H via a short lived triplet $c$-C3H2 intermediate.

The high energy cutoffs of the translational energy distributions, $P(E_T)$s, suggest a reaction exothermicity of about 10–15 kJ/mol to C3H, which stands in excellent agreement with our ab initio value of 8.6 kJ/mol. The difference falls within error limits of the experimental peak collision energies and the accuracy of the calculations. Further, the peaks of all $P(E_T)$s range between 5 and 10 kJ/mol, indicating an almost simple bond-rupture via a loose exit transition state from decomposing C3H2 reaction intermediates to the products. This framework correlates with minor geometry changes from triplet C3H2 complexes to $l/c$-C3H isomers, cf. C3H description and Tables V and VII. Compared to triplet propargylene, the C–C and C–H distances alter by less than 1 pm in $l$-C3H, i.e., the C–C bond distance changes from 127.9 pm to 124.3 pm and 134.7 pm, respectively. The bending angle of the three propargylene carbon atoms deviates by only 8.1° from the linear geometry, and the HCC bond angle widens from 156.5° to 180.0° or even remains unchanged (156.5°), if the bent structure is adopted (Table VII). Larger geometry changes up to about 17 pm are found in the cyclic C3H isomer as compared to triplet $c$-C3H2. Finally, the experimental center-of-mass angular distributions support the $C_2$ rotation axis of the propargyl isomer. This geometry and a rotation around the symmetry axis lead to a symmetric transition state to $l$-C3H, since both hydrogen atoms can be interconverted and depart with equal probability in the center-of-mass angles $\Theta$ and II–$\Theta$ to yield a symmetric, collision energy invariant isotropic shape.

With rising collision energies from 8.8 kJ/mol, over 28.0 kJ/mol, to 45.0 kJ/mol, the shape of the observed center-of-mass angular distributions changes significantly, showing a decrease of the forward–backward intensity ratio at 0° and 180° (see Refs. 9, 10). This indicates the occurrence of two microchannels. We show elsewhere,8,9 that the combination of ab initio information characterizing the C3H2 surface and molecular beam experiments allows us to attribute the strong forward peaking microchannel to the formation of $c$-C3H, a channel which becomes quenched with rising collision energy. The second microchannel leads to $l$-C3H through the propargylene C3H2 complex. This explains a higher $c$-C3H to $l$-C3H number density ratio observed in dark clouds vs hotter envelopes of carbon stars.
V. CONCLUSION

High level *ab initio* calculations [CCSD(T)] have been performed to investigate energetics of the initially formed triplet C$_2$H$_2$ collision complexes. Resolving energetics and properties of these isomers is crucial for the neutral–neutral reaction of C$_3$P with C$_2$H$_2$ to the interstellar isomers of C$_2$H, studied by crossed molecular beam experiments. The most stable almost linear isomer of C$_2$H$_2$ shows C$_2$ symmetry in agreement with results of a recent FTIR experiment. Similar almost linear structures of C$_2$H$_2$ are found to be virtually isoenergetic, although they are characterized to be saddle points on the CCSD(T) hypersurface. Other isomers are energetically unfavored relative to the most stable $c_{\perp}$ isomer by more than 1.34 kJ/mol.

The exothermicity for the reaction of C$_3$P with C$_2$H$_2$ to $l$-C$_3$H is computed as $-8.6$ kJ/mol at the present highest level of theory. This value is consistent with experimental data inferred from the molecular beam experiments. The reaction energy to linear (or quasilinear) C$_2$H is calculated to be slightly less exothermic.

The combination of *ab initio* calculations and crossed molecular beam experiments, described extensively elsewhere, allows for the explicit identification of both isomers of C$_2$H, the cyclic and the linear form. This reaction under single collision conditions avoids the need for successive binary encounters as in ion–molecule processes. Reaction dynamics in the molecular beam experiment show that the microchannel leading to $c$-C$_2$H is quenched with rising collision energy. This explains hitherto unresolved astronomical observations of number density ratios $c$-C$_2$H vs $l$-C$_3$H of 1 in cold molecular clouds (e.g., TMC-1) compared to $0.2 \pm 0.1$ found in the hotter envelope surrounding IRC +10216.

ACKNOWLEDGMENTS

C.O. thanks Professor Jürgen Gauss (Universität Mainz, Germany) and Dr. Dage Sundholm (University of Helsinki, Finland) for support in using the ACES II program system and providing a DEC version of this package. C.O. further thanks Professor John F. Stanton (University of Texas at Austin) and Professor Jürgen Gauss for valuable discussions. C.O. and R.I.K. acknowledge financial support by postdoctoral fellowships from the DFG (‘Deutsche Forschungsgemeinschaft’). M.-H.G. acknowledges support of the National Science Foundation (CHE-9357129) and by fellowships from the David and Lucile Packard Foundation and the Alfred P. Sloan Foundation. This work was further supported by the Director, Office of Energy Research, Office of Basic Energy Sciences, Chemical Sciences Division of the U.S. Department of Energy under Contract No. DE-AC03-76SF00098.

---

36. C. M. Pless and M. S. Pless, Phys. Rev. 46, 618 (1934).