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Structure and hydration of the C₄H₄⁺⁺ ion formed by electron impact ionization of acetylene clusters

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Here we report ion mobility experiments and theoretical studies aimed at elucidating the identity of the acetylene dimer cation and its hydrated structures. The mobility measurement indicates the presence of more than one isomer for the C₄H₄⁺⁺ ion in the cluster beam. The measured average collision cross section of the C₄H₄⁺⁺ isomers in helium (38.9 ± 1 Å²) is consistent with the calculated cross sections of the four most stable covalent structures calculated for the C₄H₄⁺⁺ ion [methylene cyclopropane (39.9 Å²), 1,2,3-butatriene (41.1 Å²), cyclobutadiene (38.6 Å²), and vinyl acetylene (41.1 Å²)]. However, none of the single isomers is able to reproduce the experimental arrival time distribution of the C₄H₄⁺⁺ ion. Combinations of cyclobutadiene and vinyl acetylene isomers show excellent agreement with the experimental mobility profile and the measured collision cross section. The fragment ions obtained by the dissociation of the C₄H₄⁺⁺ ion are consistent with the cyclobutadiene structure in agreement with the vibrational predissociation spectrum of the acetylene dimer cation (C₂H₂)₂⁺⁺ [R. A. Relph, J. C. Bopp, J. R. Roscioli, and M. A. Johnson, J. Chem. Phys. 131, 114305 (2009)]. The stepwise hydration experiments show that dissociative proton transfer reactions occur within the C₄H₄⁺⁺(H₂O)ₙ clusters with n ≥ 3 resulting in the formation of protonated water clusters. The measured binding energy of the C₄H₄⁺⁺·H₂O cluster, 38.7 ± 4 kJ/mol, is in excellent agreement with the G3(MP2) calculated binding energy of cyclobutadiene⁺⁺·H₂O cluster (41 kJ/mol). The binding energies of the C₄H₄⁺⁺(H₂O)ₙ clusters change little from n = 1 to 5 (39–48 kJ/mol) suggesting the presence of multiple binding sites with comparable energies for the water–C₄H₄⁺⁺ and water–water interactions. A significant entropy loss is measured for the addition of the fifth water molecule suggesting a structure with restrained water molecules, probably a cyclic water pentamer within the C₄H₄⁺⁺(H₂O)₅ cluster. Consequently, a drop in the binding energy of the sixth water molecule is observed suggesting a structure in which the sixth water molecule interacts weakly with the C₄H₄⁺⁺(H₂O)₅ cluster presumably consisting of a cyclobutadiene⁺⁺ cation hydrogen bonded to a cyclic water pentamer. The combination of ion mobility, dissociation, and hydration experiments in conjunction with the theoretical calculations provides strong evidence that the (C₂H₂)²⁺ ions are predominantly present as the cyclobutadiene cation with some contribution from the vinyl acetylene cation. © 2011 American Institute of Physics. [doi:10.1063/1.3592661]

I. INTRODUCTION

Many complex organics including polycyclic aromatic hydrocarbons (PAHs) are present in flames and combustion processes as well as in outer space.1–5 In fact, well over 100 organic molecules including acetylene, vinylacetylene, cyclobutadiene, benzene, and PAHs, are present in interstellar clouds, molecular clouds, solar nebulae, and in envelopes expelled by evolved stars.5–10 Gas phase polymerization, ion–molecule and intracluster reactions, and catalysis on nanoparticles are important synthetic pathways for the formation of complex molecules in the atmosphere and in space.5,11–14 In ion–molecule reactions, the processes of particular interest are those that lead to larger molecules, which may lead to the PAHs found in soot, meteorites and interstellar clouds.15–17

Intracluster ion–molecule reactions are uniquely suited for the discovery of novel catalytic pathways that can lead to the formation of complex organics. Since acetylene is the smallest organic molecule that can be polymerized, extensive studies have been focused on the ion chemistry of acetylene clusters not only due to the important roles of acetylene in flames and combustion processes including the mechanisms of soot formation, but also for the origin of larger molecular species such as benzene and polycyclic aromatic hydrocarbons (PAHs) in space.16,18–20 For example, the formation of benzene ions within ionized acetylene clusters, (C₂H₂)⁺⁺n, has been suggested by several cluster studies conducted over many years.21–28 Early experiments found the fragment ions observed from the photoionization of acetylene trimers to be identical to those obtained from other stable C₆H₆⁺⁺ isomers such as benzene, 2,4-hexadiyne, 1,3-hexadiyne, and 1,5-hexadiyne.21 However, photoelectron–photoion coincidence experiments and ab initio calculations showed that ionized acetylene dimer and trimer ions rearrange to produce...
stable covalent core ions (C₄H₄•+ and C₆H₆•+, respectively) with a large release of energy that leads to loss of a neutral acetylene molecule from the ionized clusters. Electron impact (EI) ionization of acetylene clusters (C₂H₂)n•+ with n up to ~25 showed magic numbers at n = 3 and 14 which were attributed to the formation of benzene (C₆H₆•+) and C₂₈H₂₈•+ molecular ions, respectively. Mass-selected ion mobility and collisional induced dissociation (CID) experiments coupled with theoretical calculations support the efficient formation of the benzene cations following the EI ionization of large acetylene clusters ((C₂H₂)n•+ with n up to ~50. Subsequent reactions of the benzene cation with acetylene molecules at higher temperatures (650 K) show evidence for sequential covalent addition leading to the formation of naphthalene-type ions. Recently, Relph et al., using infrared predissociation spectroscopy in conjunction with harmonic frequency calculations, showed that electron impact ionization of neutral (C₂H₂)n clusters results in the formation of a covalently bound C₄H₄•+ "core ion" which could lead to the presence of several isomers of the n = 3 species including a weak absorption attributed to the formation of the benzene cation. This work suggests that the structure of the C₄H₄•+ ion could play an important role in the cluster-mediated ion chemistry.

In the present work, we report ion mobility experiments and theoretical studies aimed at elucidating the identity of the acetylene dimer ion, (C₂H₂)₂•+, formed by EI ionization of neutral acetylene clusters. The ion mobility approach requires a comparison of the average collision cross section calculated for likely structures to those formed in the experiment. We employed density functional theory (DFT) (Ref. 29) to determine lowest energy structures of the C₄H₄•+ potential energy surface. We also probe the thermochemical properties and reactivity of the (C₂H₂)₂•+ ion by measuring its hydration energies by one to six water molecules using equilibrium thermochemical measurements at different temperatures. The comparison of the stepwise hydration enthalpy and entropy changes of the acetylene dimer cation with the thermochemical data recently reported for the acetylene trimer cation and the benzene cation provides further evidence for the covalently bound structure of the C₄H₄•+ ion.

The organization of the paper is as follows. In Sec. II, we briefly describe the mass-selected ion mobility system and methods used for measuring the mobility and determining the corresponding collision cross section of the mass-selected ions in helium. We also briefly describe the measurement of ion–molecule equilibrium and the determination of the stepwise enthalpy and entropy changes associated with the hydration of the C₄H₄•+ ion. The computational methods used for the structural calculations are briefly described in Sec. III. In Sec. IV, we present and discuss the results of the mass spectra, dissociation and mobility of the acetylene dimer ion (C₂H₂)₂•+. We also present the calculated structures of the C₄H₄•+ ion and utilize these structures to calculate collision cross sections of the C₄H₄•+ ion for comparison with the experimental cross sections obtained from the mobility measurements. This comparison provides the basis for suggesting the likely structures of the C₄H₄•+ ion involved in our experiments. Finally, we present the successive hydration enthalpy and entropy changes of the C₄H₄•+ ion with up to six water molecules and discuss the calculated hydrated structures with one and two water molecules. In Sec. V, we provide a brief summary of the results and highlight the new physical insights provided by this work on the structure and hydration of the C₄H₄•+ ion.

II. EXPERIMENTAL

The ion mobility, dissociation, and hydration experiments were performed using the VCU mass-selected ion mobility spectrometer. The details of the instrument can be found in several publications and only a brief description of the experimental procedure is given here.

Figure 1 illustrates the essential components of the ion mobility system. In the experiments, acetylene clusters were generated by supersonic expansion of a 2% acetylene/helium mixture (He ~4 atm). The mixture was passed through dry ice and moisture traps to diminish water vapor and acetone impurities (acetone is used as a stabilizing agent for pressurized acetylene). Typical ionizer electron energies ranged from 40 to 80 eV.

The ion mobility measurements are described in Sec. IV. For the ion hydration experiments, mass-selected (C₄H₄•+) ions were injected (in 5–15 µs pulses) into the drift cell containing 0.2–0.6 Torr of pure H₂O vapor. Flow controllers are used to maintain a constant pressure inside the drift cell. The temperature of the drift cell can be controlled to better than ±1 K using four temperature controllers. Liquid nitrogen flowing through solenoid valves is used to cool down the drift cell. The reaction products can be identified by scanning a second quadrupole mass filter located coaxially after

FIG. 1. Schematic diagram of the mass-selected ion mobility system at VCU.
the drift cell. The arrival time distributions (ATDs) are collected by monitoring the intensity of each ion as a function of time. The reaction time can be varied by varying the drift voltage. The injection energies (IEs) used in the experiments (5–20 eV, laboratory frame) are slightly above the minimum energies required to introduce the ions into the cell against the H2O vapor outflow from the entrance orifice. Most of the ion thermalization occurs outside the cell entrance by collisions with the water vapor escaping from the cell entrance orifice. At a cell pressure of 0.2 Torr, the number of collisions that the C4H4++ encounters with the water molecules within the 1.5 ms residence time inside the cell is about 10^4 collisions, which is sufficient to ensure efficient thermalization of the C4H4++ ions.

The ATDs of the injected C4H4++ and the (C4H4++)(H2O)n, formed inside the cell are measured as a function of the drift voltage across the cell. The ion intensity ratio (C4H4++)(H2O)n/(C4H4++)(H2O)n−1 is measured from the integrated peak areas of the ATDs as a function of decreasing cell drift field corresponding to increasing reaction times, and equilibrium is achieved when a constant ratio is obtained. Applying drift fields of 1–4 V/cm inside the drift cell at a temperature of 302 K and water pressure, P(H2O), of 0.4 Torr corresponds to ion residence times between 0.4 and 3.0 ms, respectively. Under these conditions, equilibrium is ascertained by a constant (C4H4++)(H2O)n/(C4H4++)(H2O)n−1 ratio and the equilibrium constants are then obtained using Eq. (1).

\[
K_{eq} = \frac{I[\text{C}_4\text{H}_4^++(\text{H}_2\text{O})_n]}{I[\text{C}_4\text{H}_4^++(\text{H}_2\text{O})_{n-1}]/P(\text{H}_2\text{O})},
\]

where I is the intensity of the ion peak taken from the integrated ATD. The equilibrium constants measured as a function of temperature yield ΔH and ΔS from the slopes and intercepts, respectively of the van’t Hoff plots. All of the results are replicated three or more times.

III. THEORETICAL

Geometries and relative energies for a number of interesting isomers of the empirical formula C4H4+ were calculated using the unrestricted Perdew, Burke, and Enzerhof exchange and correlation functional (UPBEPBE) and the augmented correlation-consistent polarized valence double ζ basis set (aug-cc-pVDZ).33 The aug-cc-pVDZ basis is a 5s2p/3s2p set for H, and a 10s5p2d/4s3p2d set for C. Likely geometry candidates for the dimer ions were derived from the known C4H4+ isomers obtained from the NIST databases.36 All geometry optimizations were followed by vibrational frequency calculations to confirm all minima on the relevant potential energy surface. All relative energies were zero point energy (ZPE) corrected. All calculations were performed using the GAUSSIAN 03 suite of programs.35

IV. RESULTS AND DISCUSSION

A. Mass spectra and dissociation of the acetylene dimer cation

Figure 2 displays a typical mass spectrum obtained by 46 eV EI ionization of neutral acetylene clusters formed by supersonic expansion. The distribution of the cluster ions formed reveals some striking features corresponding to the enhanced intensities (magic numbers) for the (C2H2)n+ ions with n = 2, 3, 14, 17, 22, 29, 35, and 46. The strong magic numbers at n = 2 and 3 are consistent with previous work and suggest the formation of stable C4H4++ and C6H6++ ions in exothermic processes that can lead to extensive evaporation of neutral acetylene molecules from the cluster.21–23 The other magic numbers such as n = 14, 17, etc. probably reflect the solvation of the C4H4++ and C6H6++ ions with acetylene molecules where solvent shells are formed with specific numbers of the acetylene molecules. Others have hypothesized the isomerization of the (C2H2)14+ cluster to a covalent C28H28++ ion.24

Figure 3 displays the mass spectra obtained upon injection of the mass selected (C2H2)2+ ions into the drift cell containing 0.46 Torr of He using different IEs. At lower IE (13 eV), no significant dissociation is observed except for the hydrogen loss to form the even-electron C8H16++ ion. The lack of dissociation is consistent with the formation of a covalent C4H4++ ion as opposed to an ion–molecule acetylene dimer ion (C2H2)2++. At high IE (53 eV), the observed fragments are C4H4++, C4H7+, C4H7++, C4H8+, C4H8++, C5H8+, C5H8++, C6H8++, and C7H8+ corresponding to m/z values of 52, 51, 50, 49, 39, 29, 27, 26, and 25 respectively. The observed fragments are in agreement with previous collision activated spectra of the C4H4+ cation generated from different precursor molecules.36 However, the observation of very minor fragments corresponding to the C13H16+ and C13H18++ ions suggests that losses of CH and CH2 units from the C4H4+ cation are very unlikely. This suggests that the methylenecyclop propane and the 1,2,3-butatriene structures are not likely candidates for the C4H4+ cation formed in our experiment. The apparent stability of the C4H4+ ion towards dissociation (no dissociation observed with injection energies up to 20 eV) and the dominant high energy dissociation channels involving the loss of 1, 2, and 3 hydrogen atoms provide some support for the cyclobutadiene structure.
B. Mobility of the acetylene dimer cation

The mobility $K$ of an ion is defined as:37

$$K = \frac{\nu_d}{E},$$

where $E$ is the drift field ($E = V/z$, $V$ is the drift voltage, and $z$ is the length of the cell (cm)) and $\nu_d$ is the drift velocity ($\nu_d = z/t_d$, $t_d$ is the drift time in s). In order to effectively compare mobility measurements at different cell conditions or different instruments, $K$ is normalized to standard conditions (STP) and referred to as reduced mobility, $K_o$.

$$K_o = K \left( \frac{273.15}{T} \right) \left( \frac{P}{760} \right).$$

Here, $T$ is the buffer gas temperature (K) and $P$ is the buffer gas pressure (Torr). Combination of Eqs. (2) and (3) gives

$$t_d = \left( \frac{z^2 \times 273.15}{T \times 760 \times K_o} \right) \left( \frac{P}{V} \right) + t_o,$$

where $t_o$ is the effective time spent outside the drift cell. A plot of the drift time ($t_d$) versus $PV$ gives a straight line with slope containing $K_o$ and an intercept corresponding to $t_o$. In the experiment, a packet of the mass selected ions of interest is injected into the drift cell and the arrival time distribution (ATD) is collected at varying cell voltages, $V$ (with $T$ and $P$ held constant). All the mobility measurements were carried out in the low-field limit where the ion’s drift velocity is small compared to the thermal velocity and the ion mobility is independent of the field strength ($E/N < 6.0$, where $E$ is the electric field intensity and $N$ is the gas number density and $E/N$ is expressed in units of Townsend (Td), where $1 \text{Td} = 10^{-17} \text{ V cm}^2$).37

The average collision cross section, $\Omega^{(1,1)}$, of the ions in the helium buffer gas is calculated according to the kinetic theory:

$$K = \frac{3q_e}{16N} \left( \frac{2\pi}{k_B T_{\text{eff}}} \right)^{1/2} \left( \frac{M_i + M_b}{M_i M_b} \right)^{1/2} \frac{1}{\Omega^{(1,1)}},$$

where $q_e$ is the ion charge, $N$ is the number density of the buffer gas, $T_{\text{eff}}$ is the effective temperature, $M_i$ and $M_b$ are the masses of the ion and buffer gas, respectively, and $\Omega^{(1,1)}$ is the orientationally averaged collision integral.

Figure 4 displays the ATDs for the mass selected (C$_2$H$_2$)$_2$$^{++}$ ions measured at various cell voltages. The inset in Fig. 4 shows a plot of $t_d$ versus $PV$ with the solid line representing the least square fitting to the data points. The resulting reduced mobility $K_o$ is determined to be $14.2 \pm 0.4$ cm$^2$ V$^{-1}$ s$^{-1}$ which corresponds to a collision cross section in helium of $\Omega = 38.9 \pm 1.4$ Å$^2$.

The ATDs of the injected ions are calculated for a finite packet of ions, $\phi(t)$, exiting a cylindrical drift tube through an aperture area ($a$) of 78.5 μm$^2$ using the transport Eq. (6):38

$$\phi(t) = \frac{sae^{-\alpha t}}{4\sqrt{\pi D_d t}} \left( \frac{V + z}{t} \right) \left( 1 - \exp \left( -\frac{r}{4D_d t} \right) \right) \times \exp \left( -\frac{(z - \nu_d t)^2}{4D_d t} \right),$$

where $r$ and $s$ represent the radius and surface density of a thin disk of injected ions, respectively, with the latter used as a scaling factor, $z$ is the length of the drift cell, and $\alpha$ is the reaction frequency set to zero.

Comparisons of the ATDs of the mass selected acetylene monomer, dimer, and trimer ions with the calculated profiles are shown in Fig. 5. The measured $K_o$, at 303 K for the C$_2$H$_2$$^{++}$, (C$_2$H$_2$)$_2$$^{++}$, and (C$_2$H$_2$)$_3$$^{++}$ ions are found to be $19.02 \pm 0.3$, $14.2 \pm 0.4$, and $11.54 \pm 0.3$ cm$^2$ V$^{-1}$ s$^{-1}$ corresponding to $\Omega$ of $30.2 \pm 1.4$, $38.9 \pm 1.4$, and $47.9 \pm 1.4$ Å$^2$, respectively. As shown in Fig. 5, comparisons of both experimental (open circles) and calculated (solid line) ATDs show excellent fits for the monomer C$_2$H$_2$$^{++}$ and trimer (C$_2$H$_2$)$_3$$^{++}$ ions indicating the presence of one isomer of either ion. However, for the (C$_2$H$_2$)$_2$$^{++}$ ions, the experimental ATD is considerably broader than that calculated using the transport equation (Eq. (6)) suggesting the existence of more than one C$_2$H$_2$$^{++}$ isomer in the cluster beam. The broadened ATD indicates that the collision cross sections of the C$_2$H$_2$$^{++}$ isomers present in the beam are not sufficiently different to be able to resolve their ATDs in our drift cell where isomers with less than 5% difference in collision cross section cannot be resolved.31

C. Calculated structures and collision-cross sections of the C$_4$H$_4$$^{++}$ ions

The four lowest energy structures of the C$_4$H$_4$$^{++}$ ion calculated using the aug-cc-pVDZ basis set are shown in...
Table I. As expected, the most stable C₄H₄⁺⁺ isomer is the methylenecyclopropene ion (MC, C₂ᵥ). The 1,2,3-butatriene ion (OB, D₂) is the second most stable with a relative energy of 12.2 kJ/mol. The third isomer with a relative energy of 25.8 kJ/mol is the planar cyclobutadiene ion (CB, D₂h). Next, is the 1-buten-3-yne ion (vinyl acetylene, VA), predicted to be 37.8 kJ/mol less stable than MC. The fifth most stable C₄H₄⁺⁺ isomer, the methyleneallene ion, is significantly less stable (138.1 kJ/mol relative to MC) than those included in Table I, and therefore is not considered any further.

Interestingly, all the predicted C₄H₄⁺⁺ isomers are of a tight, covalent linear/branched type conformation. No ion–molecule complex of the form C₂H₂⁺⁺C₂H₂ was found using the UPBEPBE/aug-cc-pVDZ method. However, it should be noted that the previously predicted C₂H₂⁺⁺C₂H₂ complex has a T-type structure (similar to the neutral) with C₂ᵥ symmetry with intermolecular distance of ∼4 Å. The relative energies of the C₄H₄⁺⁺ isomers as predicted by the UPBEPBE/aug-cc-pVDZ method are in qualitative agreement with the results reported from the UQCISD/6-31G* calculations but differ from the UB3LYP/6-31G* calculations.

### Table I

<table>
<thead>
<tr>
<th>Name</th>
<th>Optimized structures</th>
<th>Relative energy (kJ/mol)</th>
<th>Ω (Å²)_{l,ck}</th>
</tr>
</thead>
<tbody>
<tr>
<td>MC</td>
<td>Methylene cyclopropene</td>
<td>0</td>
<td>39.9</td>
</tr>
<tr>
<td>OB</td>
<td>1,2,3-Butatriene</td>
<td>12.2</td>
<td>41.1</td>
</tr>
<tr>
<td>CB</td>
<td>Cyclobutadiene (D₂h)</td>
<td>25.8</td>
<td>38.6</td>
</tr>
<tr>
<td>VA</td>
<td>1-Buten-3-yne</td>
<td>37.8</td>
<td>41.1</td>
</tr>
</tbody>
</table>
To investigate the identities of the $\text{C}_4\text{H}_4^{+}$ isomers present in the cluster beam, the calculated structures were used to obtain average collision cross sections and mobilities using the trajectory calculations. \(^4\) Although the calculated collision cross section of the CB isomer (38.6 Å²) is the closest to the measured value (38.9 ± 1.4 Å²), the experimental error precludes a definitive structural conclusion based only on the calculated cross sections. Also, the experimental ATD of the $\text{C}_4\text{H}_4^{+}$ ion indicates the presence of more than one isomer and therefore, a single isomer with a cross section that perfectly matches the experimental value will not reproduce the experimental ATD. To illustrate this point, the calculated mobility for each isomer was used to calculate the corresponding ATD (Eq. (6)) for comparison with the experimental ATD. The effective time the ions spent outside the drift cell ($t_\text{eff}$) (obtained from the intercept of the plot of the drift time ($t_d$) versus $P/V$, Eq. (4)) was added to the calculated ATD of each isomer for comparison with the experimental ATD. The results are shown in Fig. 6. As expected, the calculated ATD of the CB isomer is the closest to the experimental ATD because of the similar mobility. The other three isomers show displacements from the experimental ATD due to the differences between the calculated and experimental mobilities. However, none of the four isomers alone including CB is able to reproduce the experimental ATD of the $\text{C}_4\text{H}_4^{+}$ ion. This is consistent with the broadened ATD shown in Fig. 5 resulting from the presence of more than one isomer of the $\text{C}_4\text{H}_4^{+}$ ion in the experiment.

To investigate the possibility of the presence of more than one $\text{C}_4\text{H}_4^{+}$ isomer in the ionized acetylene cluster beam,
hypothetical relative concentrations of all combinations of the four lowest energy isomer pairs were fitted to the experimental ATD. For example, MC and CB at different relative concentrations (50:50, 40:60, 30:70, etc.) were fitted to the experimental ATD. Of all six $C_4H_4^{+}$ isomer combinations, only mixtures containing the cyclobutadiene ion in combination with the 1-buten-3-yn (vinyl acetylene) ion gave excellent fits to the experimental ATDs as shown in Fig. 7. Slightly reasonable fits could be also obtained by using combinations of the cyclobutadiene and the 1,2,3-butenriene ions as shown in the supplementary material. It is interesting to note that combinations containing methylenecyclopropene ion, the most stable $C_4H_4^{+}$ isomer, with other low energy isomers resulted in poor fits as shown in the supplementary material. Because of the similarity of the calculated collision cross section of CB and the experimental value, isomer combinations containing CB are expected to result in an averaged collision cross section close to the experimental value $(38.9 \pm 1.4 \text{ Å}^2)$. However, as indicated earlier, a successful pair of isomers should be able to reproduce the width of the experimental ATD and not only the peak maximum which determines the mobility value. Therefore, based on the comparison of the experimental and calculated ATDs, it appears that the $C_4H_4^{+}$ isomers produced in the ionized acetylene clusters under our experimental conditions are the cyclobutadiene and vinyl acetylene ions.

The mechanism of formation of the cyclobutadiene and vinyl acetylene ions involves intracluster ion–molecule reactions of the $C_2H_2^{+}$ ion with the neutral acetylene molecule in the cluster. Energy transfer to the low frequency cluster modes leading to evaporation of neutral acetylene molecules can dissipate the excess energy resulting from the covalent addition reactions producing the cyclobutadiene and vinyl acetylene ions. In this way, these isomers can be stabilized within the cluster by evaporative cooling which is analogous to collisional stabilization of the ionic intermediates in the gas phase at high pressures. Formation of the cyclobutadiene ion is the most kinetically favored pathway for all the low
FIG. 7. Comparisons of experimental ATDs of the (C₂H₂)₂⁺ ions (open circles) and the calculated ATDs for different combinations of the cyclobutadiene (CB) and the vinyl acetylene (VA) isomers (solid lines). In order to account for the effective time the ions spend outside the drift cell, the peak maximum of the combined ATDs of each isomer pair was adjusted to coincide with the peak maximum of the experimental ATD.

energy C₄H₄⁺⁺ isomers as it is the only isomer that requires no H shift from its precursor cluster ion, C₂H₂⁺⁺C₂H₂.⁴⁰,⁴³ Therefore, intracluster cyclization to produce the cyclobutadiene ion is expected to be kinetically driven (within the time frame of our experiment) since the heat produced by dimerization of the loose C₂H₂⁺⁺C₂H₂ complex can be efficiently dissipated by evaporation of neutral acetylene molecules from the cluster. In fact, Ono and Ng described a van der Waals type well for the loose C₂H₂⁺⁺C₂H₂ ion;⁴³ which may favor a cyclobutadiene ion route on the reaction coordinate if excess energy dissipation is efficient. Efficient stabilization of the cyclobutadiene ion by evaporation of acetylene molecules from the cluster is analogous to condensed-phase conditions where the cyclobutadiene ion formation is the favorable product.⁴⁴,⁴⁵

The second C₄H₄⁺⁺ isomer suggested by the current mobility experiment, the vinyl acetylene ion, is consistent with the results of Ono and Ng using photoionized acetylene clusters where a linear C₄H₄⁺⁺ isomer was identified as the vinyl acetylene ion.⁴³ Also, photoelectron–photoion coincidence experiments by Booze and Baer⁴¹ concluded that the C₄H₄⁺⁺ species resulting form photoionization of acetylene clusters are likely to be the cyclobutadiene ion in addition to the vinyl acetylene or butatriene ions. Furthermore, detailed theoretical investigation of the C₄H₄⁺⁺ potential energy surface by Hrouda et al.⁴⁰ concluded that isomerization of the (C₂H₂)₂⁺⁺ to cyclobutadiene is kinetically more favorable than any of the other C₄H₄⁺⁺ isomers due to the absence of H shifts in its production. Finally, the most recent results Relph et al., using infrared predissociation spectroscopy in conjunction with harmonic frequency calculations, showed that the C₄H₄⁺⁺ ion produced by sequential addition of acetylene molecule onto the C₂H₂⁺⁺ ion is predominantly present as the cyclobutadiene cation.²⁸
To shed more light on the identity of the \((C_2H_2)_2^{++}\) ion formed from the ionization of acetylene clusters, we studied the sequential addition of water molecules onto the \((C_2H_2)_2^{++}\) ion under thermal conditions. Figure 8 displays the mass spectra obtained upon injection of the mass selected dimer clusters into the drift cell containing water vapor at 0.55 Torr (337 K) and 0.45 Torr (290 K). At these temperatures (337 and 290 K), the main peaks observed are the dimer and its hydrated ions, \((C_2H_2)_2^{++}(H_2O)_n\) with \(n = 1–4\) at 337 K (\(n = 1–6\) at 290 K) and protonated water clusters, \(H^+(H_2O)_n\) with \(n = 3–6\) at 337 K (\(n = 3–7\) at 290 K).

The hydrated \((C_2H_2)_2^{++}(H_2O)_n\) ions are formed according to the stepwise association reaction (7), and the protonated water clusters \(H^+(H_2O)_n\) are attributed to proton transfer reactions within the \(C_4H_4^{++}(H_2O)_n\) clusters with \(n \geq 3\) according to reaction (8). These reactions have been observed for the hydrated benzene cation \(C_6H_6^{++}(H_2O)_n\) and hydrated acetylene trimer ion \((C_2H_2)_3^{++}(H_2O)_n\) with \(n \geq 4\).

\[
C_2H_4^{++}(H_2O)_{n-1} + H_2O \leftrightarrow C_2H_4^{++}(H_2O)_n, \quad n \geq 3.
\]

\[
C_2H_4^{++}(H_2O)_{n-1} + H_2O \rightarrow (H_2O)H^+ + C_4H_7^+, \quad n \geq 3.
\]

The observation of the deprotonation reaction (8) within the \(C_2H_4^{++}(H_2O)_n\) clusters with \(n \geq 3\) indicates that the proton affinity of the radical generated by the deprotonation of the \(C_2H_4^{++}\) ion is lower than that of the water trimer (218 kcal/mol).

Figure 9 displays van’t Hoff plots for the stepwise association of reaction (7), and the resulting \(\Delta H^{\circ}_{n-1,n}\) and \(\Delta S^{\circ}_{n-1,n}\) values are given in Table II. The \(\Delta H^{\circ}_{n-1,n}\) and \(\Delta S^{\circ}_{n-1,n}\) values could not be measured due to diminished ion count of the \(C_4H_4^{++}(H_2O)_2\) cluster as a result of the proton transfer reaction with the next water molecule to generate the \((H_2O)_3H^+\) ion. It is clear that the binding energies of the \(C_4H_4^{++}(H_2O)_n\) clusters change little from \(n = 1\) to 5, similar to the hydration behavior observed for the benzene ion. The significant entropy loss for the addition of the fifth water molecule could be explained by the presence of multiple binding sites with comparable energies for the water molecules to attach to the \(C_4H_4^{++}\) cation and, by the similarity of the bonding strength between the \(C_4H_4^{++}\) ion–water and between water–water interactions. The measured \(\Delta S^{\circ}_{n-1,n}\) also suggest the formation of adducts with restrained water molecules particularly for \(C_4H_4^{++}(H_2O)_5\) cluster.

The significant binding energy of the sixth water molecule could be explained by the formation of a cyclic water pentamer within the \(C_4H_4^{++}(H_2O)_5\) cluster. The measured binding energy for the \(C_4H_4^{++}(H_2O)_5\) cluster is consistent with the model potential/DFT investigation by Hodges and Stone which found the lowest structures of the \(n = 4\) and 5 clusters of \(H_2O^+(H_2O)_n\) to be cyclic. The drop in the binding energy of the sixth water molecule by approximately 14 kJ/mol and the drop in the \(-\Delta S^{\circ}_{4,5}\) value (Table II) suggest a structure in which the sixth water molecule perhaps hangs loose and interacts weakly with the \(C_4H_4^{++}(H_2O)_5\) cluster. This suggests that the \(C_4H_4^{++}(H_2O)_5\) cluster could consist of a cyclobutadiene cation hydrogen bonded to a cyclic water pentamer. As discussed in Sec. IV C, both the cyclobutadiene and the vinyl acetylene isomers are expected to be major constituents of the \(C_4H_4^{++}\) ion in our cluster beam. The measured binding energy for the \(C_4H_4^{++}(H_2O)_5\) cluster, \(38.0 \pm 4\) kJ/mol, is in excellent agreement with the G3(MP2)
TABLE II. Measured thermochemistry ($\Delta H_{n-1,n}^\circ$ and $\Delta S_{n-1,n}^\circ$) of hydration reaction (7).

<table>
<thead>
<tr>
<th>n</th>
<th>$\Delta H_{n-1,n}^\circ$ (kJ/mol)</th>
<th>$\Delta S_{n-1,n}^\circ$ (J/mol K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-58.7</td>
<td>-61.8</td>
</tr>
<tr>
<td>2</td>
<td>- (38.5–40.3) b</td>
<td>-89.0</td>
</tr>
<tr>
<td>3</td>
<td>-48.9</td>
<td>-95.3</td>
</tr>
<tr>
<td>4</td>
<td>-48.1</td>
<td>-108.2</td>
</tr>
<tr>
<td>5</td>
<td>-47.6</td>
<td>-69.6</td>
</tr>
</tbody>
</table>

*aUnits are $\Delta H$ (kJ/mol); $\Delta S$ (J/mol K). Estimated error: $\Delta H$ ±4 kJ/mol, $\Delta S$ ±10 J/mol K.

*bBased on G3(MP2) calculated binding energies for CB-2W and VA-2W since $\Delta H_{1,2}$ and $\Delta S_{1,2}$ values could not be measured due to diminished ion count of the $\text{C}_4\text{H}_4^{+}$(H$_2$O)$_2$ cluster resulting from the deprotonation reaction (8).

The most stable hydrated methylenecyclopropene complex (MC-W) consists of a carbon based $\text{CH}^{+\ldots} \ldots \text{OH}_2$ hydrogen bond (2.0 Å) to one of the methylenecyclopropene hydrogens. Mulliken charge analysis shows that the charge remains on the methylenecyclopropene radical cation. The calculated $\text{CH}^{+\ldots} \ldots \text{OH}_2$ binding energy is 11.0 kcal/mol. The hydrated cyclobutadiene ion (CB-W) exhibits a $\text{CH}^{+\ldots} \ldots \text{OH}_2$ length of 2.1 Å with a calculated binding energy of 41.0 kcal/mol in excellent agreement with the measured binding energy of 38.7 ± 4 kJ/mol. However, no conclusive evidence regarding the $\text{C}_4\text{H}_4^{+\ast}$ isomer identification can be drawn from the calculated binding energies of the hydrated structures since the four hydrated $\text{C}_4\text{H}_4^{+\ast}$ isomers have binding energies within the experimental uncertainty. For example, the hydrated 1,2,3-butatriene ion (OB-W) complex consists of a $\text{CH}^{+\ldots} \ldots \text{OH}_2$ (2.1 Å) type hydrogen bond with a binding energy of 41.0 kJ/mol. The final $\text{C}_4\text{H}_4^{+\ast} \ldots \text{H}_2\text{O}$ complex investigated was that of the 1-buten-3-yne ion (VA-W) where the lowest energy structure has a $\text{CH}^{+\ldots} \ldots \text{OH}_2$ (1.9 Å) type hydrogen bond and a significantly higher binding energy (11.9 kcal/mol) compared to the other three $\text{C}_4\text{H}_4^{+\ast}$ isomers. Although the 1-buten-3-yne radical ion is expected to be the other $\text{C}_4\text{H}_4^{+\ast}$ isomer in our beam, the predicted binding energy of 49.7 kJ/mol is significantly larger than the measured value of 38.0 kJ/mol.

The G3//MP2 predicted geometries, relative energies and binding energies for selected $\text{C}_4\text{H}_4^{+\ast}$(H$_2$O)$_n$ clusters are displayed in Table IV. The most stable bi-hydrated methylenecyclopentene ion, MC-2Wa, has two $\text{CH}^{+\ldots} \ldots \text{OH}_2$ hydrogen bonds with similar bond lengths of 1.9 Å; shorter than that of the mono-hydrated analog MC-W (2.0 Å). Other stable structures, MC-2Wb and MC-2Wc are, respectively, 4.2 and 5.0 kJ/mol less stable than MC-2Wa. It is clear that the second water molecule prefers to form a second hydrogen bond with the MC ion rather than with the first water molecule. The same trend is observed in the bi hydrated cyclobutadiene structures where the most stable isomer CB-2Wa has two bifurcated $\text{CH}^{+\ldots} \ldots \text{OH}_2$ bonds opposite to each other with bond lengths (2.2 and 2.1 Å) similar to the bond length of the first water molecule CB-W (2.1 Å). The second stable structure has two H-bonds with the two water molecules through the opposite hydrogen atoms of the CB ion. The third stable structure CB-2Wc involves water–water hydrogen bonding and is less stable than the structures involving H-bonds to the CB ion. The most stable structure for the hydrated 1,2,3-butatriene cluster (OB-2Wa) consists of two carbon based bifurcated $\text{CH}^{+\ldots} \ldots \text{OH}_2$ hydrogen bonds with bond lengths of 1.5 and 1.6 Å, significantly shorter than that of the mono-hydrated ion OB-W (2.1 Å). For the final bi-hydrated $\text{C}_4\text{H}_4^{+\ast}$ isomer investigated (1-buten-3-yne, VA), the most stable structure consists of the second water molecule forming a $\text{CH}^{+\ldots} \ldots \text{OH}_2$ hydrogen bond across from the mono-hydrated unit. It is interesting to note that in all cases, none of the most stable structures of the bi-hydrated ions are “externally” solvated.

In other words, the most stable structures consist of arrangements in which both water molecules are directly bound to the $\text{C}_4\text{H}_4^{+\ast}$ ion rather than to another water molecule. This trend is somewhat different from the ROHF/6-31+G** predictions for hydrated benzene cation.

**V. SUMMARY AND CONCLUSIONS**

The structure and hydration of the $\text{C}_4\text{H}_4^{+\ast}$ ion formed by electron impact ionization of neutral acetylene clusters have been investigated using the mass-selected ion mobility technique. The mobility measurement indicates the presence of...
TABLE IV. G3(MP2) structures of the \( \text{C}_4\text{H}_4^{+}\cdot\left(\text{H}_2\text{O}\right)_2 \) clusters for selected \( \text{C}_4\text{H}_4^{+} \) isomers. Energies are relative to the most stable species for a particular \( \text{C}_4\text{H}_4^{+} \) isomer. Binding energies are for removal of a water molecule from \( \text{C}_4\text{H}_4^{+}\cdot\left(\text{H}_2\text{O}\right)_2 \).

<table>
<thead>
<tr>
<th>Name</th>
<th>Optimized structures</th>
<th>Relative energies (kJ/mol)</th>
<th>Binding energies (kJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MC-2Wa</td>
<td>0.0</td>
<td>42.2</td>
<td></td>
</tr>
<tr>
<td>MC-2Wb</td>
<td>4.2</td>
<td>37.6</td>
<td></td>
</tr>
<tr>
<td>MC-2Wc</td>
<td>5.0</td>
<td>37.0</td>
<td></td>
</tr>
<tr>
<td>CB-2Wa</td>
<td>0.0</td>
<td>38.5</td>
<td></td>
</tr>
<tr>
<td>CB-2Wb</td>
<td>1.7</td>
<td>36.5</td>
<td></td>
</tr>
<tr>
<td>CB-2Wc</td>
<td>2.5</td>
<td>35.9</td>
<td></td>
</tr>
<tr>
<td>OB-2Wa</td>
<td>0</td>
<td>38.5</td>
<td></td>
</tr>
<tr>
<td>OB-2Wb</td>
<td>0.8</td>
<td>38.0</td>
<td></td>
</tr>
<tr>
<td>OB-2Wc</td>
<td>2.5</td>
<td>35.9</td>
<td></td>
</tr>
<tr>
<td>VA-2Wa</td>
<td>0</td>
<td>40.3</td>
<td></td>
</tr>
</tbody>
</table>
more than one isomer for the C$_4$H$_4$$^{+}$ ion in the cluster beam. The measured average collision cross section of the C$_4$H$_4$H$^{+}$ isomers in helium (38.9 ± 1.4 Å$^2$) is consistent with the calculated cross sections of the four most stable covalent structures calculated for the C$_4$H$_4$H$^{+}$ ion [methylenecyclopropane (39.9 Å$^2$), 1,2,3-butatriene (41.1 Å$^2$), cyclobutadiene (38.6 Å$^2$), and vinyl acetylene (41.1 Å$^2$)]. However, none of the single isomers is able to reproduce the experimental arrival time distribution of the C$_4$H$_4$H$^{+}$ ion. Combinations of cyclobutadiene and vinyl acetylene isomers show excellent agreement with the experimental mobility profile and the measured collision cross section. The fragment ions obtained by the dissociation of the C$_4$H$_4$H$^{+}$ ion are consistent with the cyclobutadiene structure. Both the ion mobility and dissociation experiments suggest the presence of the cyclobutadiene structure in agreement with the vibrational predissociation spectrum of the acetylene dimer cation (C$_2$H$_2$)$_2$$^{+}$.

The stepwise hydration experiments show that both the C$_4$H$_4$H$^{+}$H$_2$O$_n$ clusters with $n = 1$–6 and H$^+$(H$_2$O)$_n$ clusters with $n = 3$–6 are formed. The protonated water clusters are generated by dissociative proton transfer reactions within the C$_4$H$_4$H$^{+}$H$_2$O$_n$ clusters with $n \geq 3$. The measured binding energy of the C$_4$H$_4$H$^{+}$·H$_2$O cluster, 38.7 ± 4 kJ/mol, is in excellent agreement with the G3(MP2) calculated binding energy of cyclobutadiene$^{+}$·H$_2$O cluster (41 kJ/mol). The binding energies of the C$_4$H$_4$H$^{+}$H$_2$O$_n$ clusters change little from $n = 1$ to 5 (39–48 kJ/mol) suggesting the presence of multiple binding sites with comparable energies for the water–C$_4$H$_4$H$^{+}$ and water–water interactions. A significant entropy loss is measured for the addition of the fifth water molecule suggesting a structure with restrained water molecules probably a cyclic water pentamer, within the C$_4$H$_4$H$^{+}$H$_2$O$_5$ cluster. Consequently, a drop in the binding energy of the sixth water molecule is observed suggesting a structure in which the sixth water molecule interacts weakly with the C$_4$H$_4$H$^{+}$H$_2$O$_5$ cluster, presumably consisting of a cyclobutadiene$^{+}$ cation hydrogen bonded to a cyclic water pentamer. The combination of ion mobility, dissociation and hydration experiments in conjunction with the theoretical calculations provides strong evidence that the (C$_2$H$_2$)$_2$$^{+}$ ions are predominantly present as the cyclobutadiene cation with some contribution from the vinyl acetylene cation.

TABLE IV. (Continued)

<table>
<thead>
<tr>
<th>Name</th>
<th>Optimized structures</th>
<th>Relative energies (kJ/mol)</th>
<th>Binding energies (kJ/mol)$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>VA-2Wb</td>
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</tr>
<tr>
<td>VA-2Wc</td>
<td></td>
<td>5.4</td>
<td>34.7</td>
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</tbody>
</table>

$^a$All energies are ZPE corrected.

ACKNOWLEDGMENTS

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