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The cyclopropene radical cation: Rovibrational level structure at low energies from high-resolution photoelectron spectra

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The cyclopropene radical cation (c-C\textsubscript{3}H\textsubscript{4}+) is an important but poorly characterized three-membered-ring hydrocarbon. We report on a measurement of the high-resolution photoelectron and photoionization spectra of cyclopropene and several deuterated isotopomers, from which we have determined the rovibrational energy level structure of the X\textsuperscript{+} B\textsubscript{2} \textgreek{e} ground electronic state of c-C\textsubscript{3}H\textsubscript{4}+ at low energies for the first time. The synthesis of the partially deuterated isotopomers always resulted in mixtures of several isotopomers, differing in their number of D atoms and in the location of these atoms, so that the photoelectron spectra of deuterated samples are superpositions of the spectra of several isotopomers. The rotationally resolved spectra indicate a C\textsubscript{2h}-symmetric R\textsubscript{0} structure for the ground electronic state of c-C\textsubscript{3}H\textsubscript{4}+. Two vibrational modes of c-C\textsubscript{3}H\textsubscript{4}+ are found to have vibrational wave numbers below 300 cm\textsuperscript{-1}, which is surprising for such a small cyclic hydrocarbon. The analysis of the isotopic shifts of the vibrational levels enabled the assignment of the lowest-frequency mode (fundamental wave number of \(\approx\)110 cm\textsuperscript{-1} in c-C\textsubscript{3}H\textsubscript{4}+) to the CH\textsubscript{2} torsional mode (\(\nu_{6}^\text{t}\), A\textsubscript{3} symmetry) and of the second-lowest-frequency mode (\(\approx\)210 cm\textsuperscript{-1} in c-C\textsubscript{3}H\textsubscript{4}+) to a mode combining a CH out-of-plane with a CH\textsubscript{2} rocking motion (\(\nu_{15}^\text{t}\), B\textsubscript{2} symmetry). The potential energy along the CH\textsubscript{2} torsional coordinate is flat near the equilibrium structure and leads to a pronounced anharmonicity.

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I. INTRODUCTION

The physical and chemical properties of three-membered-ring molecules have been the object of extensive research.\textsuperscript{1-4} Especially cyclopropene (see Fig. 1), with many atypical structural parameters\textsuperscript{5-7} and a strain energy exceeding 50 kcal/mol,\textsuperscript{8,9} represents an extreme case of chemical bonding.

The existence and structure of c-C\textsubscript{3}H\textsubscript{4} was first established by Dunitz \textit{et al.} using the electron diffraction method.\textsuperscript{5} A larger set of structural parameters was derived later by Kasai \textit{et al.},\textsuperscript{6} who measured the microwave spectra of cyclopropene and three of its deuterated isotopomers, c-CH\textsubscript{3}CD\textsubscript{2}H, c-CH\textsubscript{2}(CD)\textsubscript{2}H, and c-CDH(CH)\textsubscript{2}, and determined their rotational constants. The microwave spectrum of cyclopropene was further investigated by Stiglioni \textit{et al.}\textsuperscript{7} who also studied the \textsuperscript{13}C\textsubscript{1} and \textsuperscript{13}C\textsubscript{2} isotopomers and were thus able to complete the structural determination of cyclopropene. Because three-membered-ring hydrocarbon molecules are of astrophysical interest,\textsuperscript{10} the investigations were extended to the millimeter-wave and submillimeter-wave regions of the spectrum.\textsuperscript{11,12} The analysis of these data combined with the previous microwave spectroscopic data yielded a more accurate set of rotational and centrifugal distortion constants, which was used to predict the positions of spectral lines over the entire radio-frequency range that could help in the detection of cyclopropene in the interstellar space.

The identification of a cyclopropene ring in two naturally occurring fatty acids, malvalic and sterculic acids (see Ref. 13 and references therein), and the interpretation of the IR spectra of these disubstituted cyclopropenes stimulated investigations of the IR spectra of c-C\textsubscript{3}H\textsubscript{4}. Part of the infrared spectrum of cyclopropene was reported in Ref. 14 in the context of a broader discussion of the structure and the spectra (IR, UV, and NMR) of a series of saturated and unsaturated cyclic compounds. More detailed studies of the vibrational spectrum of cyclopropene were then reported by Eggers \textit{et al.}\textsuperscript{15} and by Mitchell \textit{et al.},\textsuperscript{16} who also prepared and investigated partially deuterated samples. However, the rather low isotopic purity rendered the interpretation of the spectra difficult. The vibrational spectrum of cyclopropene was re-investigated by Yum \textit{et al.}\textsuperscript{17} who synthesized six deuterated isotopomers of cyclopropene and studied their infrared spectra at a resolution sufficiently high to enable a reliable analysis of the vibrational structure of all six isotopomers.

In contrast to the neutral species, the information currently available on the cyclopropene radical cation is limited to the partially resolved vibrational structure of He I and He II photoelectron spectra.\textsuperscript{18-21} The first photoelectron band of cyclopropene arises from the ejection of an electron out of the highest occupied molecular orbital (2b\textsubscript{2}) of cyclopropene, which has predominantly \(\pi_{\text{C=C}}\) character. Its partially resolved vibrational structure revealed a vibrational interval of \(\sim\)1300 cm\textsuperscript{-1}, indicative of a C–C stretching mode. However, the observed anharmonicity raised questions concerning this assignment and the role of the methylene-centered 1b\textsubscript{1} \(\pi\) orbital in the three-membered rings.\textsuperscript{19} A significant mixing of the \(\pi_{\text{C=C}}\) and the 1b\textsubscript{1}, \(\pi\) molecular orbital would indeed imply a \(\pi\)-electron delocalization over the entire ring and result in a significant excitation of the scissoring mode of the methylenic hydrogens (\(\nu_{5}\)) upon ionization.

Whereas the equilibrium geometry of neutral cyclopropene in its ground electronic state is of C\textsubscript{2v} symmetry,\textsuperscript{11} high-level \textit{ab initio} calculations performed by Horný \textit{et al.} for the cyclopropene radical cation in the X\textsuperscript{+} state with...
extrapolation to a complete basis-set limit predict that the Born-Oppenheimer potential-energy surface along the \( \text{CH}_2 \) torsional coordinate has two equivalent minima corresponding to \( C_2 \)-symmetric structures.\(^{22}\) These two minima are separated by a potential-energy barrier with a maximum (saddle point) of about 470 cm\(^{-1}\) at the \( C_{2v} \)-symmetric position. The height of the potential barrier is predicted to increase to \( \sim 690 \text{ cm}^{-1} \) upon inclusion of zero-point-energy corrections.\(^{22}\) The equilibrium rotational constants of the \( \tilde{X}^+ \) ground state of the cyclopropene radical cation and several deuterated isotopomers, and the fundamental vibrational wave numbers of the \( C_2 \)-symmetric structure and the \( C_{2v} \) transition state were also reported in Ref. 22.

The lack of high-resolution spectroscopic information on the energy-level structure of \( c\text{-C}_3\text{H}_4^+ \) in the literature up to date reflects the experimental difficulties associated with spectroscopic measurements on molecular cations. Photoelectron spectroscopy enables studies of the energy-level structure of cations starting from the parent neutral molecules and circumvents the necessity to generate large densities of the cationic species. Moreover, photoionization selection rules provide access to a wide range of rovibronic levels of the cation. Cyclopropene polymerizes at room temperature\(^{24}\) and is not commercially available. Moreover, it has not been possible so far to deuterate selectively specific sites of the molecule and previous studies of isotopically substituted cyclopropene had to be carried out using mixtures of several isotopomers.\(^{6,17}\) Given that photoelectron spectroscopy does not provide direct information on the mass of the investigated species, additional information must be acquired to identify the spectral structures of individual isotopomers. In the present case, the photoelectron spectroscopic data were complemented by photoionization mass-spectrometric data, which enabled us to assign most lines of the photoelectron spectra to specific isotopomers.

To determine the structure of the cyclopropene radical cation, we first determined the molecular-symmetry group from the rotational structure of the origin band of the \( \tilde{X}^+ \leftarrow \tilde{X} \) photoionizing transition of cyclopropene utilizing general rovibronic photoionization selection rules.\(^{25}\) With the same selection rules, we then also determined the vibronic symmetry of the low-lying excited vibrational levels of \( c\text{-C}_3\text{H}_4^+ \) from the rotational structure of the corresponding bands of the photoelectron spectrum. Finally, we characterized the vibrational modes by analyzing the isotopic shifts observed in the photoelectron spectra of \( c\text{-C}_3\text{H}_4, c\text{-CH}_2\text{CHCD}, c\text{-CH}_2\text{)(CD)}_2, c\text{-CD}_2\text{(CH)}_2, c\text{-CHD(\text{CD})}_2, c\text{-CD}_2\text{CHCD}, \) and \( c\text{-C}_3\text{D}_4 \).

**II. EXPERIMENTAL**

**A. Synthesis of \( c\text{-C}_3\text{H}_4 \) and its deuterated isotopomers**

The synthesis of cyclopropene and its deuterated isotopomers (see Figs. 2–4) was performed by dropwise addition of the respective 3-chloroprop-1-ene (allyl chloride) isotopomer (1) into a boiling solution of sodium bis(trimethylsilyl)amide (NaHMDS) in toluene (Fig. 2). The synthesis was carried out as described by Binger et al.\(^{26–28}\) with only minor changes; the 3-chloroprop-1-ene isotopomers were added via an automatic syringe dispenser and the deuterated cyclopropene products were condensed in a trap cooled with liquid nitrogen and further transferred into a lecture bottle.

The reaction presented in Fig. 2 was performed with five different allyl chloride isotopomers as precursors, as indicated in Figs. 3 and 4: 1,1,2-d₃-3-chloroprop-1-ene (1a), 1,1,2,3-d₄-3-chloroprop-1-ene (1b), 1,1,2,3,3-d₅-3-chloroprop-1-ene (1c), 2,3,3-d₅-3-chloroprop-1-ene (1d) (with an admixture of 25% 3,3-d₅-3-chloroprop-1-ene (1d′)), and 3-chloroprop-1-ene (1). (1) was purchased (Acros) and used without further purification, whereas the deuterated species (1a-1d) were synthesized in two to three steps.

3-d-prop-2-yn-1-ol-OD (4) was prepared from prop-2-yn-1-ol (3) by three exchanging steps in NaOD/D₂O according to Korth et al.\(^{29}\) In a second step, 3,3-d₃-prop-2-1-ol-OD (7a) and 2,3,3-d₅-prop-2-prop-1-ol-OD (7b) were prepared by reduction of 4 with lithium aluminum hydride and lithium aluminum deuteride, respectively, with subsequent hydrolysis with deuterium oxide according to Stewart et al.\(^{30}\)

For the synthesis of 1,1,2,3,3-d₅-prop-2-1-ol-OD (7c), however, a different synthetic route was followed. Prop-2-ynoic acid methyl ester (5) was deuterated in a mixture of dichloromethane and a solution of potassium carbonate in deuterium oxide by phase-transfer catalysis with tetrabutylammonium iodide to give 3-d-prop-2-ynoic acid methyl ester (6) according to Svenda and Myers.\(^{31}\) In a second step, 7c was prepared by reduction of 6 with lithium aluminum deuteride and subsequent hydrolysis with deuterium oxide according to Orfanopoulos et al.\(^{32}\)

Prop-2-ynoic acid methyl ester (5) was also reduced with lithium aluminum deuteride in tetrahydrofuran and subsequent hydrolysis with water\(^{33}\) to get a mixture of

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**FIG. 1.** Cyclopropene and its principal inertial axes in the \( \tilde{I} \) convention for near-prolate asymmetric-top molecules.

**FIG. 2.** Synthesis of cyclopropene.
FIG. 3. Synthesis of 1,1-d2-3-chloroprop-1-ene (1a), 1,1,2-d3-3-chloroprop-1-ene (1b), and 1,1,2,3,3-d5-3-chloroprop-1-ene (1c).

In the last step, the chlorination of the deuterated prop-2-en-1-ols 7a–7d to the deuterated allyl chlorides 1a–1d, respectively, was performed by the reaction with 2,2,2-trichloroacetic acid ethyl ester in dioxothiolane (sulfolane) according to Matveeva et al.34 As 7d was synthesized with an admixture of 25% 7d′, the chlorination (second step in the reaction scheme depicted in Fig. 4) also resulted in the formation of 25% 1d′.

To our knowledge, the synthesis of the deuterated species with this method is presented here for the first time. Surprisingly, every attempt at deuterating a specific site of the cyclopropene molecule yielded a mixture of several different isotopomers. The composition of the different deuterated samples of cyclopropene depended on the choice of the deuterated allyl chloride used as precursor. Four substituted samples were used in our studies, the compositions of which are summarized in Table I. The percentage of each isotopomer in the mixtures was determined by VUV photoionization mass spectrometry.

As illustration, the time-of-flight spectrum of Sample 2 (see Table I) recorded following photoionization at a wave number of 78 220 cm−1 is presented in Fig. 5. Five equally spaced peaks are observed in the spectrum indicating that the sample contains isotopomers of cyclopropene with all possible degrees of deuteration. The first peak in the spectrum at ∼5.12 μs corresponds to the time of flight of undeuterated cyclopropene cations and the last one at ∼5.37 μs to the time of flight of the perdeuterated species. The good mass resolution enabled the simultaneous recording of the photoionization spectra of different isotopomers in separate mass channels (see Fig. 10 and discussion in Subsection IV C).

FIG. 4. Synthesis of 2,3,3-d3-3-chloroprop-1-ene (1d) and 3,3-d2-3-chloroprop-1-ene (1d′).

B. Photoionization and pulsed-field-ionization zero-kinetic-energy photoelectron spectra

To record the photoelectron spectra of the $\tilde{X}^+ \xleftarrow{} \tilde{X}$ photoionizing transition of cyclopropene and its deuterated isotopomers we used a pulsed, tunable vacuum-ultraviolet (VUV) laser system in combination with a pulsed-field-ionization zero-kinetic-energy (PFI-ZEKE) photoelectron spectrometer. The spectra were obtained by monitoring the field ionization of high Rydberg states (principal quantum number n ≥ 200) located immediately below the successive ionization thresholds35 as a function of the VUV wave number. The VUV radiation was generated from the outputs of two tunable Nd:YAG pulsed dye lasers by resonance-enhanced two-color difference-frequency mixing ($\tilde{\nu}_{\text{VUV}} = 2\tilde{\nu}_1 - \tilde{\nu}_2$) in a krypton cell using the $(4p)^5 (2P_{3/2})$ state.
The laser-induced fluorescence spectrum of I$_2$ simultaneously discriminated pulse and a pulse electric-field sequence consisting of a ionization at a wave number of 78 220 cm$^{-1}$.

FIG. 5. Time-of-flight mass spectrum of Sample 2 (see Table I) recorded following ionization at a wave number of 78 220 cm$^{-1}$.

To record high-resolution photoelectron spectra, we used a tunable VUV laser with a Fourier-transform-limited bandwidth of better than 0.01 cm$^{-1}$ (Ref. 37) in combination with a multipulse electric-field ionization sequence,$^{38}$ delayed by 2.3 $\mu$s relative to the time of photoexcitation. An electric-field pulse sequence consisting of a positive discrimination pulse of +167 mV/cm followed by negative extraction pulses of −83 mV/cm, −100 mV/cm, −117 mV/cm, −133 mV/cm, −150 mV/cm, −167 mV/cm, −183 mV/cm, −200 mV/cm, −217 mV/cm, −250 mV/cm, and −333 mV/cm enabled us to record PFI-ZEKE photoelectron spectra with a resolution of about 0.15 cm$^{-1}$.

Mixtures of about 15% cyclopropene-$d_0$ or deuterated cyclopropenes in Ar at a total stagnation pressure of 2–4 bar were introduced into the chamber through a pulsed nozzle, forming a supersonic expansion. Polymerisation of cyclopropene at room temperature in the reservoir caused a gradual decrease of the signal with time, typically 30% during 6–8 h of measurement time. From the analysis of the rotationally resolved spectra, the rotational temperature of the sample in the supersonic beam was estimated to be in the range of 5–7 K.

III. GROUP-THEORETICAL CONSIDERATIONS, SYMMETRY LABELS, AND ROVIBRONIC PHOTOIONIZATION SELECTION RULES

The recent $ab$ initio calculations performed by Horný $et$ $al.$ for the cyclopropene radical cation in the $X^+$ state predict a $C_2$ equilibrium structure, the plane containing the CH$_2$ group being tilted away from that of the $C_{2v}$ structure of cyclopropene.$^{22}$ Two equivalent $C_2$ structures arise along this methylenic torsional coordinate that are separated by a potential barrier having its maximum at the $C_{2v}$ point. When including the zero-point energy of the 14 remaining modes, the height of the barrier along the CH$_2$ torsional coordinate is predicted to increase from 470 cm$^{-1}$ to 690 cm$^{-1}$.$^{22}$ One therefore expects the ground state of c-C$_4$H$_3^+$ to be split into two components by the tunneling effect.

When treating the rovibrational energy-level structure of c-C$_4$H$_3^+$ group theoretically, two situations must be considered. If the tunneling splitting associated with the motion along the CH$_2$ torsional coordinate cannot be resolved, the tunneling motion can be considered to be infeasible,$^{39,40}$ and the appropriate molecular-symmetry group is $C_2(M)$.$^{41}$ In this group, the CH$_2$ torsional mode ($\nu_{2v}^t$) is totally symmetric and transitions from the neutral ground state to torsional levels of...
the cation having even and odd numbers of vibrational quanta are allowed by the Franck-Condon principle. If the tunneling splitting is resolved, or if the lowest level of $c$-C$_3$H$_4^+$ lies above the barrier, the $C_{2v}$(M) group should be used. In this case, the torsional mode has $A_2$ symmetry and only the torsional levels of the cation with an even number of quanta, i.e., levels with $v_{3a}^\pm = 0, 2, 4, \ldots$, are expected to be observable from the neutral ground state, which has $A_1$ vibrational symmetry.

The rovibronic photoionization symmetry selection rules were derived in the relevant molecular-symmetry groups using the equation\textsuperscript{25}

\[ \Gamma_{\text{ve}}^\pm \otimes \Gamma_{\text{ve}}'' \cong (\Gamma^*)^\pm. \tag{1} \]

In this equation, $\Gamma_{\text{ve}}^\pm$ ($\Gamma_{\text{ve}}''$) represents the irreducible representation of the rovibronic state of the neutral (cation), $\Gamma^*$ is the dipole-moment representation, and $\ell$ the orbital angular momentum quantum number of the outgoing photoelectron partial wave. A double prime and a plus sign are used to designate quantum numbers of the neutral and cationic states, respectively. The analysis of the rotational structure of cyclopropene was performed using a rigid-rotor Hamiltonian for near-prolate asymmetric tops. The choice of axes of the molecule-fixed coordinate system is presented in Fig. 1 and corresponds to the convention $\Gamma$.\textsuperscript{42} The $z$ axis coincides with the twofold axis of symmetry, the $x$ axis is parallel to the carbon-carbon double bond, and the $y$ axis is perpendicular to the molecular plane.

The rovibronic photoionization symmetry selection rules can be expressed in terms of the changes $\Delta K_a = K_a^+ - K_a^-$ and $\Delta K_c = K_c^+ - K_c^-$ in the asymmetric-top quantum numbers $K_a^+(\pm)$ and $K_c^+(\pm)$ of the neutral molecule (ion).\textsuperscript{25} Assuming that the equilibrium geometry of the cyclopropene radical cation is $C_{2v}$, Eq. (1) translates into

\[ \Delta K_a = \text{even/odd}, \quad \Delta K_c = \text{odd for } \ell = \text{odd}, \tag{2a} \]

\[ \Delta K_a = \text{odd}, \quad \Delta K_c = \text{odd for } \ell = \text{even}, \tag{2b} \]

for the $\tilde{X}^+ (0^0, \Gamma_{\text{ve}} = B_2) \leftrightarrow \tilde{X}(0^0, \Gamma_{\text{ve}} = A_1)$ ionizing transition of cyclopropene. Equivalently, the selection rules can be expressed in the notation $\Gamma_{\text{ve}}'' = \Gamma_{\text{ve}}$ as

\[ A_1 \leftrightarrow A_1; A_2 \leftrightarrow A_2; B_1 \leftrightarrow B_1; B_2 \leftrightarrow B_2 \text{ for } \ell = \text{odd}, \tag{3a} \]

\[ A_1 \leftrightarrow A_2; B_1 \leftrightarrow B_2 \text{ for } \ell = \text{even}. \tag{3b} \]

If the molecular cation is treated in the $C_{2v}$(M) molecular-symmetry group, the selection rules can be easily deduced from Eqs. (3a) and (3b) by dropping the indices “1” and “2”:\n
\[ A \leftrightarrow B; B \leftrightarrow B, \tag{4} \]

or equivalently

\[ \Delta K_a = \text{odd}, \quad \Delta K_c = \text{even or odd} \tag{5} \]

for both even- and odd-$\ell$ partial waves.

The analysis of the rotational structure observed in the photoelectron spectra of $c$-CHD(CH)$_2$ and $c$-CD$_2$CHCD must be performed in the $C_{2v}$(M) molecular-symmetry group. In the first case, the plane of symmetry contains the $a$ and $c$ inertial axes and the symmetry of the ground electronic state of the cation is $\Gamma_a^+ = \Gamma_c^+ \approx \Gamma'$. Using the convention $\Gamma$,$^{42}$ the selection rules

\[ \Delta K_a = \text{even/odd}, \quad \Delta K_c = \text{even/odd for both } \ell = \text{even and odd} \]

result for the $\tilde{X}^+(0^0, \Gamma_{\text{ve}} = A') \leftrightarrow \tilde{X}(0^0, \Gamma_{\text{ve}} = A')$ ionizing transition of $c$-CHD(CH)$_2$.

In the case of $c$-CD$_2$CHCD, the symmetry plane is that defined by the $a$ and $b$ inertial axes and the symmetry of the ground electronic state of the cation is $\Gamma_a^+ = \Gamma_b^+$, and the photoionization selection rules for the $\tilde{X}^+(0^0, \Gamma_{\text{ve}} = A') \leftrightarrow \tilde{X}(0^0, \Gamma_{\text{ve}} = A')$ ionizing transition are

\[ \Delta K_a = \text{even/odd}, \quad \Delta K_c = \text{odd for } \ell = \text{odd, and} \]

\[ \Delta K_a = \text{even/odd}, \quad \Delta K_c = \text{even for } \ell = \text{even}. \tag{7b} \]

To designate the vibrational modes, we use the convention described by Herzberg,\textsuperscript{43,44} with mode indices given in the order of the irreducible representations and, within each irreducible representation, in order of decreasing wave number. Using this convention, the same mode (e.g., the CH$_2$ torsional mode) can have different mode indices in different isotopomers. Table II summarizes the mode indices for the low-frequency vibrations of the different isotopomers observed in this study. The mode associated with the CH$_2$ torsion is $v_{3a}^+$ in the $C_{2v}$(M) and $C_{2v}$(M) group, but $v_{15}^+$ in the $C_{2v}$(M) group.

### IV. RESULTS

#### A. Rovibronic structure of the photoelectron spectrum of $c$-C$_3$H$_4$ and the molecular-symmetry group of $c$-C$_3$H$_4^+$

The overview PFI-ZEKE photoelectron spectrum of the $\tilde{X}^+ \leftrightarrow \tilde{X}$ transition of $c$-C$_3$H$_4$ in the vicinity of the adiabatic ionization threshold is presented in Fig. 6. Five bands are observed in the region up to $\sim$700 cm$^{-1}$ of internal energy: the origin band at $\sim$77932 cm$^{-1}$, a very weak band at 78043 cm$^{-1}$, two relatively strong bands located 230 cm$^{-1}$ and 414 cm$^{-1}$ above the origin band, and a weak band at 78503 cm$^{-1}$ (see Table III). Additional very weak bands are observed beyond 78400 cm$^{-1}$.

The high-resolution PFI-ZEKE photoelectron spectrum of the origin band of the $\tilde{X}^+ \leftrightarrow \tilde{X}$ transition of cyclopropene is displayed in Fig. 7(a). The sharpest lines in the experimental spectrum correspond to isolated transitions and have a full width at half maximum of 0.15 cm$^{-1}$. These lines are labelled using the notation $N_{k_2}^N_{k_1}N_{k_2}^N_{k_1}$. The analysis of the rotational structure observed in the photoelectron spectrum of $c$-C$_3$H$_4$ was first performed in the $C_{2v}$(M) and $C_{2v}$(M) groups using the photoionization selection rules (5), and (2a) and (2b), respectively. The rotational intensity distribution of the spectrum was calculated using the
orbital ionization model described in detail in Ref. 45. In this model, the angular momentum of the photon is assumed to be fully absorbed by the outgoing photoelectron, creating a hole of the same angular-momentum composition as the molecular orbital out of which the electron is ejected. With the choice of axes presented in Fig. 1, the highest occupied molecular orbital of neutral cyclopropene is of $b$ and $b_2$ symmetry restricting the expansion to orbitals having a quantum number $\lambda = 1$ for the projection of the orbital angular momentum onto the principal axis $z = a$ (see Fig. 1).

The dominant rotational branches are therefore expected to be those obeying the conditions $|\Delta N| = |N^+ - N^-| \leq 2$ and $\Delta K_a = K_a^+ - K_a^- = \lambda = \pm 1$. The parameters that were optimized in the analysis of the spectrum included the adiabatic ionization energy, the rotational constants of $c$-$C_3H_2^+$, the rotational temperature of the sample, and the relative intensities of the different rotational branches. The rotational constants of the neutral ground state were kept fixed at the values reported in Ref. 6.

The rovibronic photoionization selections rule appropriate for the $C_2v$ group (Eq. (5)) are less restrictive than those appropriate for the $C_2v(M)$ group (Eqs. (2a) and (2b)) and lead to a higher density of lines than observed experimentally. Consequently, all attempts to reproduce the spectrum using the $C_2v(M)$ group remained unsuccessful, and the analysis was performed using the $C_2v(M)$ group. The optimized molecular constants of the radical cation determined from the analysis of the rotational structure are presented in Table IV. The best agreement with the experimental spectrum was achieved for a rotational temperature of $\sim 5$ K and by taking into account the angular-momentum components $|\ell^+| = (1, \pm 1)$ and $|\ell^-| = 2, \pm 1$ with weights of 0.65 and 1.9, respectively. Cyclopropene has two pairs of identical protons, giving rise to nuclear-spin statistical factors of 6 or 10 depending on the even or odd nature of $K_a^\rho$, respectively. The calculated spectrum is presented in Fig. 7(b) and reproduces the observed line positions and intensity distribution satisfactorily. The density of lines in the PFI-ZEKE photoelectron spectrum of the $0_0^+$ band is high and only a selected number of transitions are labelled in Fig. 7(a) for clarity. To nevertheless provide a complete assignment, the calculated spectrum shown in Fig. 7(b) was decomposed into groups of transitions corresponding to $(K_a^\rho, K_b^\rho)$ values of $(2, 1), (1, 0), (0, 1)$, and $(1, 2)$ (traces (c)–(f)) which exhibit branches for all possible values of $\Delta N = 0$ allowed by the orbital ionization model.

![FIG. 6. Overview PFI-ZEKE photoelectron spectrum of the $0_0^+$ ionizing transition of $c$-$C_3H_4$ at low energies. Two vibrational progressions have been identified and assigned to the $v_4^\rho$ (CH$_2$ torsion) and $v_2^\rho$ (mainly CH$_2$ rocking) vibrational modes. The positions of the CH$_2$ torsional levels are marked along the assignment bar. The dashed lines indicate the positions we expect for the levels that were not observed experimentally. The wave number region where the 15$^4_B$ band is anticipated is denoted by a horizontal bar with diagonal stripes.](image-url)
Because the next two most intense bands of the photoelectron spectrum of c-C₃H₄ (the third and fourth bands in Fig. 6) are about three to four times weaker than the origin band, a larger extraction electric field of ~330 mV/cm had to be applied to obtain PFI-ZEKE photoelectron spectra with a sufficient signal-to-noise ratio, which limited the resolution to about 0.5 cm⁻¹. The partially rotationally resolved spectra of these bands are displayed in panels (a) and (b) of Fig. 8 (middle traces).

In these panels, the inverted traces correspond to the spectra calculated based on the assumption that the ionic state has A₁ vibrational symmetry. The rotational constants and the intensity weighting factors of the different rotational branches were varied at the values determined from the analysis of the origin band (see Fig. 7(a)). The only parameters that were varied were the ionization energies, the optimized valence wave numbers.

All modes of A₁ vibrational symmetry of the cyclopropene radical cation have fundamental wave numbers larger than 600 cm⁻¹ according to the ab initio calculations presented in Ref. 22. Consequently, the observed vibrationally excited states must be overtones of nontotally symmetric vibrational modes, which, in turn, implies extremely low fundamental wave numbers.

To characterize these modes, we recorded spectra of isotopically substituted cyclopropene, and analyzed the isotopic shifts arising from the deuterium of specific sites. As explained in Subsection II A, it was not possible to synthesize isotopically pure samples and the photoelectron spectra of isotopically substituted cyclopropane consisted of contributions from several isotopomers. In order to assign the bands to specific isotopomers in cases where the rotational structure could not be resolved, complementary information had to be obtained from photoionization spectroscopic measurements (see Subsection IV C). The PFI-ZEKE photoelectron spectra obtained from Samples 1, 2, and 4 (see Table I) are discussed in Subsection IV D and confirm the existence of several low-lying vibrational levels of the cation.

### B. Rotational structure of the origin bands of the photoelectron spectra and adiabatic ionization energies of c-CHD(CH₂)₂, c-CD₂(CH₂)₂, c-CD₂CDCH, and c-C₃D₄

The high-resolution photoelectron spectrum of the overlapping origin bands of the X⁺ ← X' ionizing transition of the deuterated Samples 2 (c-CD₂(CH₂)₂ (53%), c-CD₂CDCH (35%), and c-C₃D₄ (8%)), see Table I) and 3 (c-CHD(CH₂)₂ (15%), c-CD₂(CH₂)₂ (77%), and c-CD₂CDCH (6%)) are...
The adiabatic ionization energy of the different isotopomers of cyclopropene have been corrected for the field-induced shift of the ionization energy. Because the adiabatic ionization energy of c-CHD(CH)₂ lies significantly lower in energy than those of c-CD₂(CH)₂ and c-CD₂CDCH, and below the range depicted in Fig. 9, as will be discussed at the end of this subsection, the PFI-ZEKE photoelectron spectra shown in Fig. 9 have a dominant contribution from c-CD₂(CH)₂ and weaker contributions from c-CD₂CDCH and c-C₃D₄.

The partially resolved rotational structure observed in these PFI-ZEKE photoelectron spectra was analyzed using an iterative procedure. First, the contribution of c-CD₂(CH)₂ to the spectrum obtained from Sample 3 (left panel of Fig. 9, Trace (a)) was calculated. The rotational constants of the neutral species were kept fixed to the values calculated ab initio at the MP2/6-311G(2d,p) level (see Table IV) and the ionization energy and rotational constants of the cationic species were optimized. The rotational temperature of the sample was kept fixed at 7 K. Once a satisfactory agreement between the observed and calculated spectra was reached, the calculated spectrum of c-CD₂(CH)₂ was subtracted from both experimental spectra, enabling us to analyze the rotational structure of the photoelectron spectrum of c-CD₂CDCH and c-C₃D₄. The intensities of the transitions of c-CD₂(CH)₂, which has one pair of identical protons and one pair of identical deuterons, were weighted by nuclear-spin statistical factors of 5 and 7 for even and odd values of \( K'' \), respectively. For c-C₃D₄, which has two pairs of identical deuterons, the intensities of the transitions were weighted by nuclear-spin statistical factors of 8 or 10 depending on the odd or even value of \( K'' \), respectively. Several iteration steps yielded the molecular constants listed in Table IV and to the decomposition of the overall spectra into contributions from selected isotopomers presented at the bottom of Fig. 9. The PFI-ZEKE photoelectron spectra calculated for the two mixtures (Trace (b) in each panel of Fig. 9) are in good agreement with the experimental spectra, which makes us confident that our procedure to decompose the spectra into contributions from the isotopomers constituting the mixtures and to determine the isotopic shifts is reliable.

![FIG. 7. Trace (a): High-resolution PFI-ZEKE photoelectron spectrum of the origin band of the \( \tilde{X}^+ \)→ \( \tilde{X} \) transition of \( c\text{-C}_3\text{H}_4 \). The asterisk placed at about 77 934 cm\(^{-1}\) above trace (a) marks a discontinuity in the experimental spectrum caused by a mode hop of the tunable ring dye laser during the measurement. Trace (b): Calculated spectrum obtained by convolution of a stick spectrum (not shown) using a Gaussian function of full width at half maximum of 0.15 cm\(^{-1}\). Traces (c)–(f): Calculated contributions to the overall intensity distribution associated with rotational branches of specific values of \( K'' \) and \( K''' \) as indicated above each trace.](image)

| TABLE IV. Molecular constants derived from the rotationally resolved PFI-ZEKE photoelectron spectra of cyclopropene and its deuterated isotopomers. |
|-----------------|-----------------|-----------------|-----------------|-----------------|
|                 | \( \tilde{B}^+(\text{cm}^{-1}) \) | \( A'^+(\text{cm}^{-1}) \) | \( B'^+(\text{cm}^{-1}) \) | \( C'^+(\text{cm}^{-1}) \) |
| \( c\text{-C}_3\text{H}_4 \) | 1.00281709 | 0.728023652 | 0.460175019 | 6 |
| \( c\text{-C}_3\text{D}_4 \) | 77 931.8(7)\(^a\) | 0.960 | 0.774 | 0.459 | This work |
| \( c\text{-CD}_2\text{(CH)}_2 \) | 0.960484470 | 0.64566341 | 0.43020258 | 6 |
| \( c\text{-CD}_2\text{(CH)}_2 \) | 78 008.3(20)\(^a\) | 0.9494\(^b\) | 0.6738\(^b\) | 0.4303\(^b\) | This work\(^*,\) Ref. 22\(^f\) |
| \( c\text{-CD}_2\text{(CD)}_3 \) | 78 078.3(10) | 0.872 | 0.619 | 0.413 | This work\(^d\) |
| \( c\text{-CD}_2\text{CHCD} \) | 0.817 | 0.553 | 0.380 | This work\(^d\) |
| \( c\text{-CD}_2\text{CHCD} \) | 78 084.2(15) | 0.781 | 0.589 | 0.379 | This work |
| \( c\text{-C}_3\text{D}_4 \) | 0.717 | 0.538 | 0.350 | This work\(^d\) |
| \( c\text{-C}_3\text{D}_4 \) | 78 086.3(20) | 0.685 | 0.573 | 0.350 | This work |

\(^a\)The values of the adiabatic ionization energy of the different isotopomers of cyclopropene have been corrected for the field-induced shift of the ionization energy. 
\(^b\)The uncertainties in the values of the \( A'^+ \), \( B'^+ \), and \( C'^+ \) rotational constants are estimated to be ±0.020 cm\(^{-1}\), ±0.015 cm\(^{-1}\), and ±0.010 cm\(^{-1}\), respectively.
\(^c\)The adiabatic ionization energy of \( c\text{-C}_3\text{H}_4 \) was determined to be 77 994(240) cm\(^{-1}\) from the He I photoelectron spectrum. 
\(^d\)Calculated ab initio at the MP2/6-311G(2d,p) level.
Whereas deuteration of the ethylenic sites of the molecule does not significantly shift the adiabatic ionization energy of cyclopropene, successive deuteration of the methylenic site lowers the adiabatic ionization energy in steps of about 75 cm$^{-1}$. The PFI-ZEKE photoelectron spectrum of the origin band of c-CHD(CH)$_2$ is therefore located below the range depicted in Fig. 9. Because of the low concentration of this isotopomer in the mixture, a large extraction pulse...
of −831 mV/cm had to be applied to achieve a sufficiently high signal-to-noise ratio. From the spectrum (not shown) the value of the adiabatic ionization energy of c-CHD(CH)2 was determined to be 78 008.3(20) cm⁻¹.

C. Photoionization mass spectrometry and spectroscopy

The different isotopomers present in the deuterated samples pose a problem in the identification of specific bands of the survey photoelectron spectra to a given isotopomer. To overcome this problem, the photoelectron spectra were complemented by photoionization spectra. In photoionization spectroscopy, the yield of ions of a certain mass is recorded as a function of the wave number of the ionizing radiation. Each time an ionization threshold is crossed, an ionization channel opens and the ion signal is expected, ideally, to rise in a stepwise manner. However, interactions between different ionization channels, which lead to autoionization resonances in photoionization spectra, can make the steps difficult to observe. In cases where the Rydberg states are predisso- cative, however, the contributions of autoionizing resonances to photoionization spectra can be strongly suppressed. The combination of mass-selective photoionization spectroscopy with photoelectron spectroscopy, which is not mass selective, then represents a powerful tool for the spectroscopic investigation of samples that are mixtures of several species.

In the upper panel of Fig. 10, traces (b), (c), and (d) correspond to the photoionization spectra of c-CD₂(CH)₂, c-CD₄CDCH, and c-C₂D₄, respectively, obtained with Sample 2 (see Table I). The intensity scale was corrected for the variation of the intensity of the VUV radiation, and the relative intensities of the different spectra reflect the relative abundances of the corresponding isotopomers. The photoionization spectra of c-C₃H₄ and c-CHD(CH)₂ are not shown, because the signal-to-noise ratio was insufficient to extract useful information. For comparison, the PFI-ZEKE photoelectron spectrum of this sample recorded over the same wave-number range is displayed as Trace (a) in Fig. 10. In all three photoionization spectra, the signal rises sharply from a background level of nearly zero, forming a step near 78 083 cm⁻¹ at the position of the first adiabatic ionization energy. Additional steps are visible in the photoionization spectra, the first one located at approximately 78 255 cm⁻¹ and common to all three isotopomers, and the others at around 78 340 cm⁻¹ (Trace (d)), 78 360 cm⁻¹ (Trace (c)), and 78 380 cm⁻¹ (Trace (b)) for c-C₂D₂⁺, c-CD₂CDCH⁺, and c-CD₄CH⁺, respectively. Every stepwise increase in the photoionization signal corresponds to a line in the photoelectron spectrum and vice versa, as highlighted by the dashed vertical lines that connect the lines of the PFI-ZEKE spectrum with the steps observed in the photoionization spectra. The photoionization spectra therefore exhibit an almost ideal behavior in this wave-number range, which allows the decomposition of the PFI-ZEKE photoelectron spectrum of the mixture into contributions from selected isotopomers and the reconstruction of the vibrational energy-level structure of the corresponding cations. The absence of autoionization resonances in the photoionization spectrum of cyclopropene further indicates that the Rydberg states of cyclopropene are subject to rapid predissociation in the region of the adiabatic ionization threshold.

The PFI-ZEKE photoelectron spectra obtained using Samples 1 and 4 (see Table I for their compositions) are displayed in Traces (a'), and in the bottom panel of Fig. 10, respectively. The lines observed in these spectra can be assigned to specific isotopomers by comparison with the corresponding photoionization spectra (not shown) using the same procedure. For the identification, we use a short notation in which the numbers separated by a comma indicate the carbon atoms to which the deuterium atoms are attached and the suffix “−dₓ” with i = 0–4 gives the number of deuterium atoms. The analysis confirms the adiabatic ionization energies determined from the decomposition of the high-resolution photoelectron spectra presented in Sec. IV B. Their values are primarily determined by the degree of deuteration of the methylenic group: The adiabatic ionization energies of c-C₃H₄, c-CH₂CHCD, c-CH₂CD₂ are located around 77 930 cm⁻¹, those of c-CHD(CH)₂ and c-CHD(CD)₂ around

FIG. 10. Upper panel, Traces (a) and (a'): PFI-ZEKE photoelectron spectrum of the X ← X photoionizing transition of c-CD₂(CH)₂, c-CD₄CDCH, and c-C₂D₄ recorded with the deuterated Samples 2 and 1, respectively (see Table I for their compositions). Upper panel, Traces (b), (c), and (d): Photoionization spectra of c-CD₂(CH)₂, c-CD₄CDCH, and c-C₂D₄, respectively, obtained from Sample 2. The vertical dashed lines connect the steps in the photoionization spectra with the peaks of the corresponding PFI-ZEKE photoelectron spectrum (Trace (a)). Lower panel: PFI-ZEKE photoelectron spectrum of the X ← X photoionizing transition of c-C₃H₄, c-CH₂CHCD(CD)₂, c-CH₂CD₂, and c-CHD(CD)₂ recorded with Sample 4 (see Table I for its composition).
78 010 cm\(^{-1}\), and those of c-CD\(_2\)(CH\(_2\))\(_2\), c-CD\(_2\)CDCH, and c-C\(_3\)D\(_4\) around 78 085 cm\(^{-1}\) (see Tables III and IV).

Below the adiabatic ionization thresholds, the photoionization spectra of c-CD\(_2\)(CH\(_2\))\(_2\), c-CD\(_2\)CDCH and c-C\(_3\)D\(_4\) reveal a weak step at \(\sim 78\,045\,\text{cm}^{-1}\) at the same position as a weak line in the PFI-ZEKE photoelectron spectrum of sample 2 (Trace (a)). These weak features must correspond to a sequence band. The fundamental wave numbers of the lowest-frequency vibration in the neutral species, the CH (or CD) out-of-plane bend (\(v_{\text{15}}^+\)) amounts to 562 cm\(^{-1}\), 471 cm\(^{-1}\), and 424 cm\(^{-1}\) for the doubly, triply, and the fully deuterated cyclopropene isotomers, respectively. These sequence bands may therefore correspond to transitions from the fundamental out-of-plane bending levels of the neutral species to the fundamental out-of-plane bending levels of the cation (14\(^1\) for c-CD\(_2\)(CH\(_2\))\(_2\) and c-C\(_3\)D\(_4\) and 13\(^1\) for c-CD\(_2\)CDCH\(^+\), see Table II). From this very tentative assignment, one would deduce out-of-plane bending fundamental wave numbers of \(\approx 525\,\text{cm}^{-1}(14\(^1\))\) for c-CD\(_2\)(CH\(_2\))\(_2\), \(\approx 430\,\text{cm}^{-1}(13\(^1\))\) for c-CD\(_2\)CDCH\(^+\), and \(\approx 385\,\text{cm}^{-1}(14\(^1\))\) for c-C\(_3\)D\(_4\). The PFI-ZEKE photoelectron spectrum of the fourth sample displayed in the lower panel of Fig. 10 is more congested than the photoelectron spectra presented as traces (a) and (a’) because the sample contains isotopomers with different degrees of deuteration of the methylenic group which have different origins, at about 77 930 cm\(^{-1}\) for the isotopomers containing an undeuterated methylenic group and at about 78 010 cm\(^{-1}\) for the isotopomers containing a singly deuterated methylenic group. Above 78 200 cm\(^{-1}\), the spectral density is too high for unambiguous assignments to be made.

The positions of the ionization thresholds of the various deuterated isotopomers of cyclopropene determined from photoelectron and photoionization spectra are listed in Table III. The observed isotopic shifts served as a basis for the spectral assignments presented in Subsection IV D.

### D. Assignment of the vibrational structure of c-C\(_3\)H\(_4\), c-CH\(_2\)CHCD\(^+\), c-CH\(_2\)CD\(_2\)H\(^+\), c-CD\(_2\)(CH\(_2\))\(_2\), c-CHD(cD\(_2\))\(_2\), c-CD\(_2\)CDCH\(^+\), and c-C\(_3\)D\(_4\)\(^+\) at low energies

The photoelectron spectra of c-C\(_3\)H\(_4\)\(^+\) (see Fig. 6) and deuterated isotopomers (see Fig. 10) consist of three dominant features at low energies, the first one corresponding to the adiabatic ionization energies, and the other two to two low-lying vibrational levels of the cations. These levels behave differently upon deuteration. The vibrational wave number of the first level (230(2) cm\(^{-1}\) in c-C\(_3\)H\(_4\)\(^+\)) is insensitive to deuteration at the ethylenic sites but shifts by about \(-30\,\text{cm}^{-1}\) upon each successive deuteration at the methylenic site, which indicates that the vibrational motion is localized at the methylenic group. The vibrational wave number of the second level (414(2) cm\(^{-1}\) in c-C\(_3\)H\(_4\)\(^+\)) is reduced by \(\approx 110\,\text{cm}^{-1}\) upon double deuteration at the methylenic site and by \(\approx 20\,\text{cm}^{-1}\) upon each deuteration at the ethylenic sites. This behavior is indicative of a complex vibrational motion of the cation involving primarily the CH\(_2\) group and, to a lesser extent, also the ethylenic hydrogen atoms.

We conclude from the observed isotopic shifts that the two main bands associated with excited vibrational levels of the cyclopropene radical cation must correspond to two different low-frequency vibrational modes. According to \textit{ab initio} calculations, the two lowest vibrational modes of c-C\(_3\)H\(_4\)\(^+\) are the CH\(_2\) torsional mode and a mode corresponding to a combination of CH\(_2\)-rocking and CH-out-of-plane-bending motion. Because neither of these modes is totally symmetric in the C\(_3v\)(\(M\)) group, one would only expect to observe transitions to levels with an even number of vibrational quanta. Moreover, the analysis of the rotationally resolved photoelectron spectra presented in Subsection IV B indicated that these vibrational levels of the cation are totally symmetric (A\(_i\)), from which one must conclude that they are overtones: overtone of the CH\(_2\) torsion (i.e., \(8^\text{th}\)) for the level of c-C\(_3\)H\(_4\)\(^+\) observed 230(2) cm\(^{-1}\) above the origin, and overtone of the combined CH\(_2\)-rocking and the CH-out-of-plane motion (i.e., \(15^\text{th}\)) for the level observed 414(2) cm\(^{-1}\) above the origin. This interpretation further enables the assignments of (1) the very weak band at 78 043 cm\(^{-1}\) in the photoelectron spectrum of c-C\(_3\)H\(_4\) (see Fig. 6) to a nominally “forbidden” transition to the fundamental of the CH\(_2\) torsional mode, \(8^\text{th}\), (2) the band at 78 503 cm\(^{-1}\) to the \(8^\text{th}\) transition, and (3) part of the broader structure observed at \(\approx 78\,820\,\text{cm}^{-1}\) to the \(15^\text{th}\) transition. All vibrational assignments are summarized in Table III.

The fundamental wave numbers obtained from our photoelectron spectra are compared to those calculated \textit{ab initio} for the C\(_2\) equilibrium structure and the C\(_2\)\(_s\) saddle point in Table V. Comparing our results with the results of these \textit{ab initio} calculations leads to three main observations: (i) Whereas our results indicate a C\(_2\)\(_s\) (\(R_\text{s}\)) structure for the X\(^+\) state of c-C\(_3\)H\(_4\)\(^+\), the calculations predict two equivalent C\(_2\) equilibrium structure (\(R\)) separated by a potential barrier of 490 cm\(^{-1}\) along the torsional coordinate, with maximum (saddle point) at the C\(_2\)\(_v\) structure. The height of the potential barrier is predicted to rise to 690 cm\(^{-1}\) upon inclusion of zero-point-energy corrections. (ii) The fundamental wave number we determined for the CH\(_2\)-torsional mode (\(v_{15}\)) is significantly smaller than that determined \textit{ab initio} for the C\(_2\) structure, which is a direct consequence of (i). (iii) The fundamental wave number we determined for the CH\(_2\)-rocking mode (\(v_{13}\)) is significantly smaller than that determined \textit{ab initio} for the C\(_2\)\(_s\) saddle point, but similar to that calculated for the C\(_2\) equilibrium structure.

Our spectroscopic observations indicate a pronounced anharmonicity of the low-frequency vibrational modes, which may be the reason for these discrepancies. The calculations

<table>
<thead>
<tr>
<th>Character</th>
<th>Vibr. wave numbers (exp)</th>
<th>CCSD(T)/cc-pVTZ ((C_2) sym.)(^{22})</th>
<th>MP2/cc-pVQZ ((C_2) sym.)(^{22})</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH(<em>2) torsion ((v</em>{15}))</td>
<td>111 cm(^{-1}) (A(_2))</td>
<td>452 cm(^{-1}) (A(_2))</td>
<td>260 cm(^{-1}) (A(_2))</td>
</tr>
<tr>
<td>CH(<em>2) rocking ((v</em>{13}))</td>
<td>(\sim 207) cm(^{-1}) (B(_2))</td>
<td>287 cm(^{-1}) (B(_2))</td>
<td>541 cm(^{-1}) (B(_2))</td>
</tr>
</tbody>
</table>

\(^{a}\)Estimated as half the overtone wave number.

\(^{b}\)The corresponding vibrational motion is delocalized over the whole molecule and the description as “CH\(_2\)-rocking” is simplified.
might indeed have overestimated the zero-point-energy corrections. The potential energy along the CH$_2$ torsional mode appears to be flat near the C$_2v$ structure so that the torsional wave function extends over a wide range of torsional angles, beyond the angle corresponding to the \textit{ab initio} C$_2$ equilibrium structure. We therefore expect the CH$_2$-rocking wave number calculated \textit{ab initio} for the C$_2$ structure to correspond more closely to our experimental result than that calculated for the C$_2v$ saddle point, in agreement with the data presented in Table V.

V. CONCLUSIONS

The rovibrational energy level structures of the cyclopropene radical cation and several deuterated isotopomers have been determined from high-resolution photoelectron and photoionization spectra of the parent neutral molecules in the vicinity of the $\tilde{X}^+ \leftarrow \tilde{X}$ photoionizing transition. The rotational structure of the photoelectron bands indicated that the cation has a C$_2v$ ($R_v$) structure and enabled the derivation of the vibrational symmetry of several low-lying vibrational levels of c-C$_3$H$_4$. The adiabatic ionization energies of c-C$_3$H$_4$, c-CHD(CH)$_2$, c-CH$_2$CHCD, c-CH$_2$(CD)$_2$, c-CD$_2$(CH)$_2$, c-CD$_2$CDCH, and c-C$_2$D$_2$ were determined and so were the rotational constants of the ground state of most of the corresponding radical cations.

The vibrational structure of the photoelectron spectra associated with the $\tilde{X}^+ \leftarrow \tilde{X}$ transition of cyclopropane was assigned by analyzing the isotopic shifts upon deuteration. Two low-frequency vibrational modes of the cyclopropane radical cation were identified, one associated with the CH$_2$ torsional motion ($\nu^t_{15}$), the other with the CH$_2$ rocking motion ($\nu^r_{15}$). A vibrational progression exhibiting positive anharmonicity (i.e., characterized by a negative anharmonicity constant) was observed in the CH$_2$ torsional mode ($\nu^t_{15}$) of c-C$_3$H$_4$.

The excitation, from the neutral ground state, of the fundamentals of these modes is forbidden within the Franck-Condon approximation. However, the $\delta^t_{10}$ band could be observed as a weak band, presumably because of vibronic interactions. The excitation of the first overtones, $\delta^t_{10}$ and $\delta^t_{15}$, is allowed and, in the case of c-C$_3$H$_4^+$, they are observed at 234 cm$^{-1}$ and 417 cm$^{-1}$ above the vibrational ground state, respectively.

The removal of an electron from the highest occupied molecular orbital of cyclopropane, which has a CC $\pi$ bonding character and $b_2$ symmetry in the C$_2v$ point group, has a strong impact on the molecular structure, particularly on the CH$_2$ torsional and rocking modes and on the out-of-plane CH bending modes. The fundamental wave number of the CH$_2$ torsional mode decreases from about $\sim 1000$ cm$^{-1}$ (Ref. 17) in the neutral species to only about 110 cm$^{-1}$ in the radical cation. Similarly, the fundamental wave number of the CH$_2$ rocking mode decreases from $\sim 1090$ cm$^{-1}$ (Ref. 17) in the neutral species to less than 210 cm$^{-1}$ in c-C$_3$H$_4^+$. Our spectroscopic observations indicate a pronounced positive anharmonicity of the low-frequency vibrational modes, which may be the reason for the discrepancies with the results of recent high-level \textit{ab initio} calculations.\textsuperscript{22} One should note, however, that this comparison is complicated by the fact that the observed vibrational structure may be affected by intermode anharmonic couplings and vibronic coupling. The low-frequency modes observed experimentally imply a fluctional behavior of the H atoms which is related to the delocalization of the $\pi$ bonding orbital over the entire ring.\textsuperscript{19}

The occurrence of several low-frequency modes caused significant spectral congestion already at low internal energies of the cation and made the spectra difficult to assign. Systematic deuteration of the different sites, which normally facilitates the assignments did not work as well as we had hoped: All attempts at synthesizing specific deuterated isotopomers resulted in mixtures of several isotopomers. Although the assignment of the dominant spectral structures to individual isotopomers could be made by comparison of the PFI-ZEKE photoelectron spectra with photoionization spectra, several assignments remain uncertain. In future, more selective synthetic routes to produce specific isotopomers, and spectroscopic experiments at higher resolution may help to clarify the discrepancies between our spectroscopic results and the latest \textit{ab initio} calculations.

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We thank Dr. Urs Hollenstein, Dr. Bérenger Gans, and Dr. René Verel (all three ETH Zurich) for experimental help, and Dr. M. Willeke, Dr. L. Horný (both ETH Zurich), Professor H. F. Schaefer III (University of Georgia, Athens), and Professor J. F. Stanton (University of Texas) for useful discussions. This work is supported financially by the Swiss National Science Foundation under Project No. 200020-146759.

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