# Polarized Infrared Absorption Spectra of Matrix-Isolated Allyl Radicals

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We have measured the polarized infrared absorption spectrum of the allyl radical,  $CH_2CHCH_2$  ( $\tilde{X}$ )  $^2A_2$ , in an argon matrix at 10 K. The experimental  $CH_2CHCH_2$  frequencies (cm $^{-1}$ ) and polarizations follow:  $a_1$  modes, 3109, 3052, 3027, 1478, and 1242;  $b_1$  modes, 983, 801, and 510;  $b_2$  modes, 3107, 3020, 1464, 1390, and 1182. Two modes ( $\nu_6$  and  $\nu_{18}$ ) are very weak and could not be detected; the lowest frequency  $a_1$  mode (the  $CH_2-CH-CH_2$  bending mode  $\nu_7$ ) is estimated to be beyond the wavelength range of our MCT infrared detector. Infrared absorption spectra of two deuterated isotopomers,  $CH_2CDCH_2$  and  $CD_2CDCD_2$ , were recorded in order to compare experimental frequency shifts with calculated [UB3LYP/6-311-G(d,p)] harmonic frequencies. Linear dichroism spectra were measured with photooriented samples in order to establish experimental polarizations of most vibrational bands. True gas-phase vibrational frequencies were estimated by considering the gas-to-matrix shifts and matrix inhomogeneous line broadening. The allyl radical matrix frequencies listed above are within  $\pm 1\%$  of the gas-phase vibrational frequencies. A final experimental set of all the vibrational frequencies for the allyl radical are recommended. See also: http://ellison.colorado.edu/allyl.

### I. Introduction

The allyl radical, CH<sub>2</sub>CHCH<sub>2</sub>, is one of the simplest conjugated hydrocarbon radicals. It is commonly depicted as  $[CH_2-CH=CH_2 \leftrightarrow CH_2=CH-CH_2]$  and is used as an example of resonance. 1,2 The stabilization of the allyl radical is generally reckoned<sup>3,4</sup> to be roughly 40 kJ mol<sup>-1</sup> (10 kcal mol<sup>-1</sup>). As a consequence of the importance of this resonance stabilization, the thermochemistry of CH<sub>2</sub>CHCH<sub>2</sub> (plus the ions, CH<sub>2</sub>CHCH<sub>2</sub><sup>+</sup> and CH<sub>2</sub>CHCH<sub>2</sub><sup>-</sup>) has been carefully studied. The adiabatic ionization potential was originally has been measured<sup>5</sup> via classical photoelectron spectroscopy as  $IP(CH_2CHCH_2) = 8.13$  $\pm$  0.02 eV. The allyl's ionization potential has recently been redetermined via a Rydberg extrapolation<sup>6</sup> to be *IP*(CH<sub>2</sub>CHCH<sub>2</sub>)  $= 65.638 \pm 18 \text{ cm}^{-1} \text{ [or } 8.138 \pm 0.002 \text{ eV]}$  which differs slighly from a ZEKE measurement<sup>7,8</sup> of  $IP(CH_2CHCH_2) =$  $65\,762\pm5~\text{cm}^{-1}$  [8.1535  $\pm$  0.0006 eV]. The adiabatic electron affinity<sup>9,10</sup> is  $EA(CH_2CHCH_2) = 0.581 \pm 0.008$  eV. The bond energy for propene has been carefully measured to be D<sub>0</sub>(CH<sub>2</sub>- $CHCH_2-H) = 365.7 \pm 1.7 \text{ kJ mol}^{-1} (87.4 \pm 0.4 \text{ kcal mol}^{-1}),$ and the bond enthalpy<sup>3</sup> is  $DH_{298}(CH_2CHCH_2-H) = 371.5 \pm 10^{-2}$  $1.7~\rm kJ~mol^{-1}~(88.4\pm0.4~\rm kcal~mol^{-1})$ . These bond dissociation energies imply  $\Delta_f H_0(CH_2CHCH_2) = 184.5 \pm 2.1 \text{ kJ mol}^{-1}$  and  $\Delta_{\rm f} H_{300}({\rm CH_2CHCH_2}) = 173.2 \pm 1.7 \ {\rm kJ \ mol^{-1}}$ . Since the bond energy of cyclopropane, 11 c-C<sub>3</sub>H<sub>5</sub>, has been reported to be  $DH_{298}(c-C_3H_5-H) = 444 \pm 13 \text{ kJ mol}^{-1} (106 \pm 3 \text{ kcal mol}^{-1}),$ one can estimate that the rearrangement of the cyclopropyl radical to allyl radical will be exothermic by roughly 1.4 eV.

$$C = -1.4 \text{ eV}$$

$$H \qquad H \qquad H \qquad H$$

$$C + CH_2 \qquad H \qquad H \qquad H$$

$$C + C_3H_5 \quad \widetilde{\chi}^2 A' \qquad CH_2CHCH_2 \quad \widetilde{\chi}^2 A_2$$

In this paper we report the infrared absorption spectrum of  $CH_2CHCH_2$  ( $\tilde{X}$ )  $^2A_2$ . There have been many previous studies of the vibrational spectrum of allyl but they are incomplete and somewhat contradictory. Over the past few years we have developed a convenient hyperthermal nozzle to prepare matrix isolated organic radicals.  $^{12,13}$  We have used this intense radical source to prepare matrix isolated samples of the three isotopic variants of the allyl:  $CH_2CHCH_2$ ,  $CH_2CDCH_2$ , and  $CD_2CDCD_2$ . Following preparation of the target allylic radicals, we have used a Fourier transform infrared spectrometer (FTIR) to measure the infrared absorption spectra of these species. With the help of a polarized laser, the linear dichroism spectrum was measured with photooriented samples in order to establish experimental polarizations of most of the vibrational bands of the allyl radical

## **II. Experimental Section**

**A. Radical Beam and Infrared Spectroscopy.** Most matrixisolated radicals are prepared by photodissociation of an appropriate precursor (usually by a laser). A drawback of this procedure is that the target radical is almost always photochemically active and suffers photofragmentation. Consequently, as fast at the laser generates the target radical, it may also destroy it. Thus, we have resorted to high-temperature flash pyrolysis (850 °C) of thermally labile precursors. We produce the allyl

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radical in the gas phase by pyrolysis of a suitable precursor molecule in a hyperthermal, supersonic jet expansion. The hyperthermal dosing nozzle is interfaced with a commercial APD two stage closed-cycle helium cryostat that is maintained at 25 K. CH<sub>2</sub>CHCH<sub>2</sub> radicals are deposited onto a cold CsI window and detected using a Nicolet Magna 550 FTIR. The IR detectors are mercury-cadmium-telluride sensors; the MCT-A which has a detection range from 4000 to 600 cm<sup>-1</sup> and while the MCT-B has a detection range from 4000 to 400 cm<sup>-1</sup> and a much lower sensitivity. Polarized laser light for photoorientation experiments is generated at 351 nm from a Coherent Innova 200/25 CW Argon ion laser. Polarized IR light is generated using a Molectron wire grid IR polarizer.

The hyperthermal supersonic nozzle used in our experiments is a modified version of the nozzle developed in the laboratory of Peter Chen. 14 Our version 12,13 consists of a Series 9 Parker General Valve pulsed solenoid valve (0.1 mm diam) followed by a concentric 1 mm silicon carbide tube. The resistively heated silicon carbide tube is connected through two carbon electrodes to the output of a Love Control microprocessor-based temperature controller (model 16A3051). The temperature is monitored using a type C thermocouple attached to the center of the silicon carbide tube, which continuously supplies feedback to the controller unit; typical operating temperatures<sup>13</sup> are roughly 1150 K. This control system maintains the temperature steady within typically  $\pm 1$  K over the course of an experiment. The length of the heated region of the silicon carbide tube is approximately 2 cm. The residence times in the heated region of the hyperthermal nozzle are estimated to be less than 100  $\mu$ s. Residence times are believed to increase as pyrolysis temperature increases.<sup>15</sup> The output of the nozzle is directed toward a cryogenic matrix window and the radicals entrained in the molecular beam are frozen onto the cryogenic substrate. Dosing times are typically 3-4 h.

This work focuses on the infrared spectrum of the allyl radical parent and two of its deuterated isotopomers, CH2CDCH2 and CD<sub>2</sub>CDCD<sub>2</sub>. Precursors used to produce the allyl radical included CH2=CHCH2I, CH2=CHCH2Br, CH2=CHCH2S2CH2-CH=CH<sub>2</sub>, and CH<sub>2</sub>=CHCH<sub>2</sub>CH=CH<sub>2</sub> which were supplied by Aldrich Chemical Co. The appropriate deuterated allyl iodides, CH<sub>2</sub>=CDCH<sub>2</sub>I and CD<sub>2</sub>=CDCD<sub>2</sub>I, were synthesized as described below. All precursors afforded us samples of the allyl radical but some were better than others. We observed near complete depletion (roughly 90-95%) of the 1,5-hexadiene and allyl iodide precursors in the hyperthermal nozzle. Vibrational frequencies and intensities were matched for the different precursors used. Spectra shown in this work are from the allyl iodide precursor.

Gas mixtures were primarily argon seeded with a suitable radical precursor. These gas mixtures were made by sampling the vapor of the de-gassed precursor liquid at ambient temperature. Higher vapor pressure contaminants such as CH<sub>3</sub>OCH<sub>3</sub> or CH<sub>3</sub>I became enriched in the dosing mixture. CH<sub>2</sub>=CHCH<sub>2</sub>S<sub>2</sub>-CH<sub>2</sub>CH=CH<sub>2</sub> and CH<sub>2</sub>=CHCH<sub>2</sub>CH=CH<sub>2</sub> will decompose to produce two allyl radical and were used in approximately half the concentration compared to the other precursors. Gas mixtures were made in a 300 mL or a 1.3 L reservoir; deposition quantities and dosing rates were determined by monitoring pressure drop in the reservoir. We pulsed the dosing nozzle at a repetition rate of 30 to 10 Hz in order not to exceed the pumping speed of our system. Valve-opening times were 150-300  $\mu$ s. A hyperthermal nozzle firing at 30 Hz for 150  $\mu$ s produces about 10<sup>18</sup> Ar atoms s<sup>-1</sup> and roughly 10<sup>15</sup> CH<sub>2</sub>CHCH<sub>2</sub> radicals s<sup>-1</sup> at the 25 K CsI window of the cryostat as estimated from choked flow at the 0.1 mm orifice.

B. PIMS Spectroscopy as a Radical Diagnostic. One of the experimental difficulties of matrix-isolated radical spectroscopy is to devise and optimize a clean source of radicals. The hyperthermal nozzle has been shown by Chen<sup>14</sup> to be an excellent radical source which he developed using a photoionization mass spectrometer (PIMS) to monitor and adjust the radical beam. We have adopted a PIMS as well in order to optimize our allyl radical source.

Prior to matrix deposition for an FTIR study, the optimal parameters for clean CH2CHCH2 radical production in the dosing nozzles was determined by monitoring products in a reflectron time-of-flight photoionization mass spectrometer. The mass spectrometer used in this experiment is a commercial RM Jordan reflectron TOF instrument that has been described in detail previously. 16 The output of the hyperthermal dosing jet is allowed to expand for approximately 1 cm and is skimmed before the molecular beam enters a laser interaction chamber. In the laser interaction region, an orthogonally orientated beam of 118.2 nm light (10.487 eV) crosses the molecular beam, ionizing the molecules entrained in the beam with ionization potential less than 10.5 eV. Positive ions are deflected by a positively biased deflector plate and injected through a pinhole into a time-of-flight mass analyzer. Ions are mass selected by a reflectron analyzer<sup>17</sup> and detected at off-axis microchannel plates.

Tripling<sup>18</sup> the third harmonic of a Nd:YAG laser in a xenon tripling cell generates the  $(3 \times 3 \times 1.064 \,\mu\text{m}^3)$  118.2 nm laser pulse. The 118.2 nm light is focused at the interaction region while the nontripled 532 nm light diverges. The laser has a fixed repetition rate of 10 Hz and provides the timing trigger for the experiment.

Before introduction of the thermally dissociated gas mixture, the instrument was calibrated with propylene (m/z, 42) or other simple hydrocarbons mixed with an inert gas. The parameters of the hyperthermal nozzle were then adjusted to the point where only the desired radical ion and leaving group ion are detected. Note that the peak intensities are not a reflection of the relative amounts of species present. Peak heights depend on both detection efficiencies and photoionization cross sections. Detection efficiencies are a function of particle mass and momentum. Photoionization cross sections are dependent<sup>19</sup> on the species' ionization potential and vibrational excitation.

C. Chemical Synthesis of Allyl Radical Precursors. To produce samples of CH2CDCH2 and CD2CDCD2, the corresponding allylic iodides were required; these compounds were synthesized. A dry 250 mL round-bottomed, three-necked flask was equipped with a condenser, an addition funnel, and a stir bar. The flask was charged with 1.20 g (28.6 mmol) of lithium aluminum deuteride, 3.09 g (57.2 mmol) of sodium methoxide, and 90 mL of freshly distilled diethyl ether. The mixture was cooled in an ice bath while propargyl alcohol, HC≡CCH<sub>2</sub>OH (1.5 mL, 25.8 mmol), in 60 mL of ether was added dropwise from the addition funnel over 1 h. The reaction mixture was heated to reflux for 52 h, then cooled in an ice bath and quenched by addition of 5 mL of water. After stirring for 4 h, the reaction mixture was filtered through a glass frit. Ether was distilled from the filtrate at atmospheric pressure with a shortpath distillation head. The remaining material was then fractionally distilled. Allyl-2-d<sub>1</sub> alcohol, CH<sub>2</sub>=CDCH<sub>2</sub>OH, was collected as a colorless oil at 96-105 °C (45% yield). <sup>1</sup>H NMR (400 MHz,  $C_6D_6$ )  $\delta$  3.80 (s), 4.95 (m), 5.14 (m).

TABLE 1: Previously Measured CH2CHCH2 Vibrational Fundamentals

		review <sup>55</sup>	matrix FTIR <sup>41</sup>	matrix FTIR <sup>43</sup>	matrix FTIR <sup>39</sup>		$MPI^{34}$	Rydberg <sup>6</sup>	resonance Raman <sup>35</sup>	laser absorption <sup>27</sup>	laser absorption <sup>25</sup>	laser absorption <sup>26</sup>	negative ion PES <sup>10</sup>
$\overline{a_1}$	$\nu_1$		3105	3107	3109			3212			3113.98488	3113.9779	
	$\nu_2$	3048	3048	3051, 3040	3051			3109			$\pm 0.00089$	$\pm 0.0003$	
	$\nu_3$		3016	3019	3019			3104					
	$\nu_4$	1488, 1477	1463	1463	1478			1504	1487				
	$\nu_5$	1245, 242	1403	1242, 1182	1242			1268	1246				
	$\nu_6$	1066	1242	972.8	1183			1032	1068				990
	$\nu_7$	427	511	510			426	423	429				425
$a_2$	$\nu_8$	5.40					550	705	5.40				
1	$\nu_9$	549	005	000.5	004	002.6	558	552	549				
$b_1$		968, 983.6	985	808.5	984	983.6		1001	968 (2 $\nu_{10}$ )	801.71909			800
	$\nu_{11}$	801.7	810	801.2	802	801.1		721		$\pm 0.00037$			800
	11.0	518, 510.1	802		511	510.1	$508 \pm 12$	516	518 (2\nu_{12})	±0.00037			
h <sub>2</sub>		3105	3105	3107	3109	510.1	300 ± 12	3210	310 (21/12)		3110.59857	3110.5	
02	13	3103	3103	5107	310)			3210			$\pm 0.00036$	5110.5	
	$\nu_{14}$	3016	3016	3019	3019			3100			_0.00000		
		1463	1477	1477	1464			1467					
	$\nu_{16}$	1389	1463	1389, 1317	1389			1382					
	$\nu_{17}$	1182	1389	1284	1284			1104					
	$\nu_{18}$		1285	983.2	810			918					
					Pre	eviously !	Measured (	CD <sub>2</sub> CDCD <sub>2</sub>	Fundamenta	ıls			
$a_1$	$\nu_1$	2205		2205									
	$\nu_2$	2285		2285									
	$\nu_3$	2215 1272, 1263		2214 1018									
	$v_4$ $v_5$	1020, 1018,		1016						1020			
	, ,	1007								1020			
	$\nu_6$	844							1272				835
	$\nu_7$								350, 357				345
$a_2$	$\nu_8$												
	$\nu_9$								$372 (2\nu_9)$				
$b_1$									$762 (2\nu_{10})$				
		647	646			646.5			202 (2				678
L	$\nu_{12}$	2272							$383 (2\nu_{12})$				
$D_2$	$\nu_{13}$	2372 2209	2209										
	$\nu_{14}$	1387	1062										
	$\nu_{15} = \nu_{16}$	1062	1002										
	$\nu_{16}$ $\nu_{17}$	900	1263										
	$\nu_{18}$	689											

The following procedure, workup, and purification were carried out in the dark. Allyl-2- $d_1$  alcohol (1.0 g, 16.9 mmol) and methyltriphenoxyphosphonium iodide (9.95 g, 22.0 mmol) were combined in 25 mL of diethyl ether at room temperature and stirred under Ar for 44 h. The reaction mixture was then washed with saturated, aqueous sodium thiosulfate and saturated aqueous sodium chloride. The aqueous layers were extracted three times with diethyl ether and the combined ether layers were dried over magnesium sulfate. Ether was distilled off at atmospheric pressure with gentle heating. When the solvent level became low, all of the volatile components were removed by distillation under reduced pressure, leaving a viscous orange residue behind. The volatile components were then placed in the smallest flask possible, and all remaining ether was distilled under atmospheric pressure. The allyl iodide was then distilled under aspirator vacuum, yielding 1.54 g (54% yield) of a light yellow oil. The product CH<sub>2</sub>=CDCH<sub>2</sub>I is 95% pure, with methyl iodide being the primary contaminant. The allyl iodide was stored in a freezer over copper metal for stability. The product is heat and light sensitive. <sup>1</sup>H NMR (200 MHz, CDCl<sub>3</sub>) δ 3.87 (s), 4.98 (m), 5.24 (m). <sup>2</sup>H NMR (61.47 MHz, C<sub>6</sub>H<sub>6</sub>, referenced to  $C_6D_6$ )  $\delta$  5.66. <sup>13</sup>C NMR (100 MHz,  $C_6D_6$ )  $\delta$  5.47 (s), 117.51 (s), 135.98 (t).

Allyl- $d_5$  iodide was prepared from allyl- $d_5$  alcohol (Aldrich) in the same manner as allyl-2- $d_1$  iodide (56% yield). Gas

chromatographic analysis shows the product  $CD_2$ = $CDCD_2$ I to be 94% pure.  $^1$ H NMR (400 MHz) shows methyl iodide as the primary contaminant, along with minor amounts of nondeuterated allyl iodide, diethyl ether, and phenol.  $^2$ H NMR (61.47 MHz,  $C_6H_6$ , referenced to  $C_6D_6$ )  $\delta$  3.22, 4.53, 4.72, 5.63.  $^{13}C$  NMR (100 MHz,  $C_6D_6$ )  $\delta$  5.21 (m), 116.93 (m), 135.58 (t).

**D.** Computational Methods. The vibrational lines in this work were assigned using harmonic frequencies from electronic structure calculations as a guide. We used the density functional method of Becke, <sup>20,21</sup> UB3LYP with a 6-311-G (p,d) basis set and the Gaussian 98<sup>22</sup> computational suite to calculate harmonic frequencies of allyl radical and its isotopomers. These computational values for allyl radical are listed in Table 2.

#### III. Results and Discussion

**A. PIMS Optimization of the Allyl Radical Source.** Prior to spectroscopic analysis of allyl radical, pyrolysis conditions were established for the precursors. The bond enthalpies of the possible precursors are all known:  $^{3,23,24}$   $DH_{298}(CH_2CHCH_2-Br) = 251 \pm 5$  kJ mol<sup>-1</sup> [ $60.0 \pm 1.1$  kcal mol<sup>-1</sup>],  $DH_{298}(CH_2-CHCH_2-I) = 200 \pm 4$  kJ mol<sup>-1</sup> [ $47.7 \pm 1.0$  kcal mol<sup>-1</sup>], and  $DH_{298}(CH_2-CH_2-CH_2-CH_2-I) = 285 \pm 4$  kJ mol<sup>-1</sup> [ $68.1 \pm 1.0$  kcal mol<sup>-1</sup>]. Allyl iodide was found to be the best precursor; PIMS traces of the  $CH_2$ = $CHCH_2I$  are shown in

TABLE 2: Calculated UB3LYP/6-311 G(p,d) Harmonic Frequencies ( $\omega$ /cm <sup>-1</sup> ) and Infrared Intensities ( $A$ /km mol <sup>-1</sup> ) for the $\tilde{X}$
<sup>2</sup> A <sub>2</sub> Allyl Radical (All Values Unscaled). To View the Animated Vibrational Modes, See http://ellison.colorado.edu/allyl

			CH <sub>2</sub> CHCH <sub>2</sub>		СН	2CDCH <sub>2</sub>	$CD_2CDCD_2$	
ω		local mode description	$\omega/\text{cm}^{-1}$	A/km mol <sup>-1</sup>	$\omega/\text{cm}^{-1}$	A/km mol <sup>-1</sup>	$\omega/\text{cm}^{-1}$	A/km mol <sup>-1</sup>
1	$a_1$	CH <sub>2</sub> CHCH <sub>2</sub> asymmetric CH <sub>2</sub> st	3235	21.0	3233	16.8	2410	11.2
2		CH <sub>2</sub> CHCH <sub>2</sub> symmetric CH <sub>2</sub> st	3141	4.6	3140	7.4	2316	2.9
3		CH <sub>2</sub> (C-H)CH <sub>2</sub> stetch	3128	15.8	2311	8.9	2278	7.7
4		symmetric CH <sub>2</sub> scissors	1517	2.9	1514	2.8	1291	0.3
5		$CH_2$ - $CH$ - $CH_2$ stretch + $CH_2$ scis	1270	0.9	1246	1.1	1040	1.4
6		CH <sub>2</sub> -CH-CH <sub>2</sub> stretch +CH <sub>2</sub> rock	1037	0.1	1036	0.1	856	0.3
7		CH <sub>2</sub> -CH-CH <sub>2</sub> bend	429	0.0	425	0.0	350	0.0
8	$a_2$	CH <sub>2</sub> CHCH <sub>2</sub> out-of-phase umbrella	786	0.0	786	0.0	622	0.0
9		CH <sub>2</sub> -CH-CH <sub>2</sub> out-of-phase twist	553	0.0	553	0.0	393	0.0
10	$b_1$	CH <sub>2</sub> (C-H)CH <sub>2</sub> out-of-plane bend	1014	21.4	816	9.0	784	1.4
11		CH <sub>2</sub> CHCH <sub>2</sub> in-phase umbrella	811	78.0	808	80.0	657	52.0
12		CH <sub>2</sub> -CH-CH <sub>2</sub> in-phase twist	532	16.5	531	17.0	408	12.5
13	$b_2$	CH <sub>2</sub> CHCH <sub>2</sub> asymmetric CH <sub>2</sub> stretch	3232	5.8	3232	3.1	2408	3.0
14		CH <sub>2</sub> CHCH <sub>2</sub> symmetric CH <sub>2</sub> stretch	3134	8.7	3134	4.3	2273	4.2
15		asymmetric CH/CH <sub>2</sub> scissors	1511	1.0	1494	0.6	1375	0.0
16		asymmetric CH <sub>2</sub> -CH-CH <sub>2</sub> stretch + CH/CH <sub>2</sub> scissors	1423	6.7	1348	3.5	1075	4.4
17		asymmetric CH <sub>2</sub> -CH-CH <sub>2</sub> stretch + CH bend	1207	0.3	1036	1.5	939	0.1
18		asymmetric CH/CH <sub>2</sub> rock	936	0.0	851	0.0	699	0.1

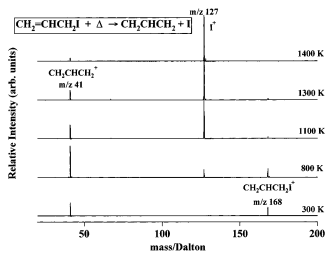


Figure 1. Photoionization mass spectra of the output of the hyperthermal nozzle. The nozzle temperature varies from 300 to 1400 K. The molecular beam is photoionized by 118.2 nm laser light (10.487 eV), and the ions are analyzed with a reflectron TOF mass spectrometer. The appearance of I<sup>+</sup> at a nozzle temperature of 1100 K shows the threshold for the thermal dissociation:  $CH_2$ = $CHCH_2I + 1100 K \rightarrow$  $CH_2CHCH_2 + I$ .

Figure 1 and demonstrates that the parent ion,  $[CH_2=CHCH_2I]^+$ , undergoes a large amount of photoionization fragmentation:  $CH_2CHCH_2I^+ \rightarrow CH_2CHCH_2^+ + I$ . The simultaneous presence of I<sup>+</sup> and CH<sub>2</sub>CHCH<sub>2</sub><sup>+</sup> is a clear signal that we are pyrolyzing CH<sub>2</sub>=CHCH<sub>2</sub>I and producing allyl radicals. On the basis of this figure, we opted to dose at 1150 K, where CH<sub>2</sub>=C=CH<sub>2</sub> production is minimal and to subtract the allyl iodide precursor spectra from the final spectra shown in this work.

B. Vibrational Spectroscopy of Allyl Radical. Highresolution, gas-phase studies of the infrared absorption spectrum of the CH2CHCH2 radical have been reported. Allyl radicals were studied using high-resolution IR absorption techniques<sup>25,26</sup> in a cooled jet. In these works the in-phase and out-of-phase asymmetric C-H stretches  $v_1$  (3113.98488 cm<sup>-1</sup>) and  $v_{13}$ (3110.59857 cm<sup>-1</sup>) were detected with rotational resolution. The  $C_{2v}$  symmetry of the allyl radical was also confirmed by studying the nuclear spin statistics in rotationally resolved spectra. Hirota<sup>27</sup> studied  $\nu_{11}$ , the most intense vibration in allyl radical. The lines in the  $v_{11}$  rotationally resolved manifold were fitted using a vibration-rotation Hamiltonian and this fundamental

was shown to be polarized along the molecular c-axis and to be a b<sub>1</sub> transition. The rotational constants were used to calculate C-C bond lengths of 1.3869  $\pm$  0.0033 Å and the CCC angle of 123.96  $\pm$  0.48 $^{\circ}$  . The change in the centrifugal distortion constants induced by the coupling of  $v_{11}$  and  $v_7$  in allyl radical has also been described.<sup>28</sup>

Earlier EPR measurements indicated that allyl radical contains only 3 types of hydrogens.<sup>29,30</sup> This is consistent with CH<sub>2</sub>-CHCH<sub>2</sub> being planar and having  $C_{2\nu}$  symmetry. Equation 2 shows the allyl radical's symmetry axis; the x axis is out of the plane and defines the  $B_1$  symmetry axis or the molecular c axis.

$$\angle (CCC) = 123.96^{\circ} \pm 0.48^{\circ}$$

$$r_{e}(CH) = 1.3869 \pm 0.0033 \text{ Å}$$
H

C

H

y (a axis, B<sub>2</sub> sym)

The ground electronic state of the allyl radical is  $(\tilde{X})$   $^2A_2$ , and this molecule has 18 vibrational modes. The irreducible representations of the allylic modes are:  $\Gamma_{VIB}(CH_2CHCH_2) =$  $7 \ a_1 \oplus 2 \ a_2 \oplus 3 \ b_1 \oplus 6 \ b_2$ . The two  $a_2$  modes are not IR active so an infrared study should only find 16 separate fundamentals.

The negative ion photoelectron spectrum<sup>9,10</sup> of the allylic anion has been observed;  $CH_2CHCH_2^- + \hbar\omega_0 \rightarrow CH_2CHCH_2$ + e<sup>-</sup>. Vibrational progressions observed in the spectrum correspond to the  $a_1$  modes  $\nu_6$ , the CCC stretch, and  $\nu_7$  the CCC bend. The overtone of the nontotally symmetric  $b_1$  mode  $v_{11}$ was (weakly) observed as well. A number of multiphoton ionization (MPI) studies  $^{31-34}$  investigated the  $\tilde{B}~^2A_1 \leftarrow \tilde{X}~^2A_2$ transition in the 3s Rydberg state of CH<sub>2</sub>CHCH<sub>2</sub>. In these studies Sappey et al.<sup>34</sup> used electronic structure calculations to assign vibrational modes  $\nu_7$ ,  $\nu_{12}$ , and  $\nu_9$  of the ground-state CH<sub>2</sub>-CHCH<sub>2</sub>,  $\tilde{X}$  <sup>2</sup>A<sub>2</sub>. Minsek et al.<sup>33</sup> used rotational contours to fit their MPI data and re-assigned these same modes. From the resonance Raman spectra of Getty et al. 35-37 and Liu et al., 38 many of the totally symmetric modes of allyl radical and allyl $d_5$  were assigned. In Raman studies the most intense lines in the Stokes spectrum are attributed to totally symmetric vibrations. Even overtones of nontotally symmetric vibrations, which reflect large changes between the excited-state geometry and the ground-state such as  $\nu_9$ ,  $\nu_{10}$ , and  $\nu_{12}$ , were also detected.

The CH<sub>2</sub>CHCH<sub>2</sub> radical has been the subject of several previous matrix IR studies.<sup>39–43</sup> Many of the vibrational modes in these studies are consistent with modes identified in the present work. Allyl radical lines were confirmed through performing annealing experiments where the matrices were heated to temperature of 30–40 K. Lines that were depleted were believed to be due to allyl radical. In these earlier studies, the best ab initio electronic structure calculations (at that time) were used as a guide for assignments. There were no mechanisms to assign symmetries of the separate vibrational modes.

All of the previous experimental spectroscopic work on the CH<sub>2</sub>CHCH<sub>2</sub> radical is summarized in Table 1.

C. Electronic Structure of Allyl Radical and Photoorientation Experiments. To anticipate the later photoorientation experiments, the electronic structure of allyl radical is reviewed. The electronic configuration of the ground state is ...  $(1b_1)^2(1a_2)^1$  leading to a  $\tilde{X}$   $^2A_2$  symmetry. In a first-order approximation, the first low-lying excited electronic state of  $CH_2CHCH_2$  will be a superposition of the two configurations,  $(1b_1)^1(1a_2)^2$  and  $(1b_1)^2(2b_1)^1$ , leading to  $^2B_1$  symmetry for the  $\tilde{A}$   $CH_2CHCH_2$  state.

The electronic spectrum of allyl radical was first detected by Currie and Ramsey<sup>44</sup> who studied the lower electronic transition,  $\tilde{A} \leftarrow \tilde{X}$ , centered at 408.3 nm. From the diffuseness of the band of the transitions and lack of rotational structure they concluded that allyl radical predissociates in the lowest excited state. Callear and Lee<sup>45</sup> studied the upper excited states of allyl radical in the 210-250 nm region. They assigned their observed spectrum to transitions in an upper state of <sup>2</sup>B<sub>1</sub> symmetry. These studies were confirmed by Maier<sup>41</sup> who also observed a strong structureless UV transition centered at 213 nm and a weak transition centered at 408.5 nm indicating that the allyl radical dissociates in both excited-state manifolds. The upper excitedstate potential energy surface was calculated by Ha et al. 46 and subsequently investigated using MPI by Blush et al.47 Blush studied the  $\tilde{D}$   $^2B_2 \leftarrow \tilde{X}$   $^2A_2$ ,  $\tilde{C}$   $^2B_1 \leftarrow \tilde{X}$   $^2A_2$ , and the dipole-forbidden,  $\tilde{B}$   $^2A_1 \leftarrow \tilde{X}$   $^2A_2$ , transitions in the 238–250 nm region. They found substantial vibronic mixing between electronic levels in the excited-state manifold. They further conjecture that allyl radical loses its  $C_{2v}$  symmetry in making the forbidden  $\tilde{B}^2A_1 \leftarrow \tilde{X}^2A_2$  transition allowed.

In photoorientation experiments we excite a dissociative electronic transition of CH<sub>2</sub>CHCH<sub>2</sub> with known symmetry in order to selectively deplete the matrix-isolated allyl species and generate a preferentially aligned distribution of radicals. We use the polarized 351 nm line from an Ar II laser to excite the blue edge of the  $\tilde{\rm A}$   $^2{\rm B}_1\leftarrow \tilde{\rm X}$   $^2{\rm A}_2$  transition. For the allyl transition dipole moment,  $\langle ^2{\rm B}_1|\hat{\mu}|^2{\rm A}_2\rangle$ , to be nonzero, the  $\tilde{\rm A}$   $^2{\rm B}_1\leftarrow \tilde{\rm X}$   $^2{\rm A}_2$  transition must be polarized along  $\mu_{\rm y}$  (or  ${\rm B}_2$ ) and is aligned parallel to the main a molecular axis of allyl radical, see eq 2.

When the matrix is dosed the cryogenically trapped CH<sub>2</sub>-CHCH<sub>2</sub> radicals have a random distribution. Irradiation of these allyl radicals with polarized UV light at 351 nm dissociates molecules that have a substantial projection of their transition dipole moments parallel to the depleting laser light.<sup>48</sup> The probability of photon absorption and subsequent dissociation for a molecule i is proportional to  $\langle \cos^2(\theta_i) \rangle$  where  $\theta_i$  is the angle between the transition dipole moment  $\hat{\mu}_i$  and  $\mathbf{E}$ . The molecules remaining after the photodepletion process have their transition moments  $\hat{\mu}_i$  preferentially oriented perpendicular with respect to the  $\mathbf{E}$  vector of the depleting laser light.

This preferentially oriented ensemble of radicals is prepared using laser light, which is horizontally oriented with respect to the laboratory frame or in the Z direction. The allyl molecules are preferentially oriented with their molecular a axes perpendicular to this depleting laser light and thus vertically oriented with respect to the laboratory frame. The ensemble of molecules are then measured using two linearly independent polarized IR beams in the Z and Y directions, parallel and perpendicular to the depleting laser light. The IR absorption intensity for the a<sub>1</sub> (parallel to molecular b axis) and the  $b_1$  modes (parallel to the molecular c axis) will be greater using IR light polarized in the Z direction and be less using IR light polarized in the Y direction. That is  $I_Z - I_Y \ge 0$  (exhibit positive linear dichroism or LD) for modes of a<sub>1</sub> and b<sub>1</sub> symmetry. Correspondingly absorption intensities for b2 modes will be lower using horizontally oriented (Z direction) IR light for measurement; that is  $I_Z - I_Y < 0$ (exhibit negative linear dichroism or LD) for modes of b2 symmetry. In this work the symmetry of the b<sub>2</sub> modes is determined unambiguously. Unfortunately, because the excited state manifold has a high degree of mixing, there is no convenient method to preferentially orient the ensemble of trapped allyl radicals in order to separately assign the a<sub>1</sub> and b<sub>1</sub> modes.

**D.** Assignments of the Allyl Radical IR Spectra. Matrix-isolated allyl radicals are reactive. Three contaminants that we repeatedly encounter in this work are allene, propene, and 1,5-hexadiene. The sequential bond enthalpies of the allyl radicals are known.<sup>4</sup>

$$CH_2$$
= $CHCH_3 \rightarrow CH_2CHCH_2 + H$   
 $DH_{298} = 372 \pm 2 \text{ kJ mol}^{-1} (3a)$ 

$$CH_2CHCH_2 \rightarrow CH_2 = C = CH_2 + H$$
  
 $DH_{298} = 109 \pm 3 \text{ kJ mol}^{-1} \text{ (3b)}$ 

Consequently a pair of allyl radicals can disproportionate in a 263 kJ mol<sup>-1</sup> (62.7  $\pm$  0.7 kcal mol<sup>-1</sup>) exothermic process: CH<sub>2</sub>-CHCH<sub>2</sub> + CH<sub>2</sub>CHCH<sub>2</sub>  $\rightarrow$  CH<sub>2</sub>=C=CH<sub>2</sub> + CH<sub>3</sub>CH=CH<sub>2</sub>. Likewise a pair of allyl radicals can simply dimerize in a 285 kJ mol<sup>-1</sup> (68.1  $\pm$  1.0 kcal mol<sup>-1</sup>) exothermic process to produce CH<sub>2</sub>=CHCH<sub>2</sub>-CH<sub>2</sub>CH=CH<sub>2</sub>. Peaks marked as CH<sub>2</sub>=C=CH<sub>2</sub> are the most intense bands of allene, <sup>49-51</sup>  $\nu_6$ (CH<sub>2</sub>=C=CH<sub>2</sub>) = 1957 cm<sup>-1</sup> and  $\nu_{10}$ (CH<sub>2</sub>=C=CH<sub>2</sub>) = 837 cm<sup>-1</sup>. The feature marked with **H** in the parent allyl radical spectra at 914 cm<sup>-1</sup> (Figure 2) is the most intense band of 1,5-hexadiene, the allylallyl recombination product. Smaller peaks (not marked) attributed to 1,5-hexadiene are found at 996 and 1361 cm<sup>-1</sup>.

We have measured the infrared absorption spectrum of three  $C_{2v}$  isotopomers of the allyl radical: allyl- $d_0$  (CH<sub>2</sub>CHCH<sub>2</sub>), allyl- $d_1$  (CH<sub>2</sub>CDCH<sub>2</sub>), and allyl- $d_5$  (CD<sub>2</sub>CDCD<sub>2</sub>). The results are displayed in Figures 2–13. The final experimental vibrational frequencies for the matrix-isolated allyl radicals are collected together in Table 3.

Figure 2 is an overview of the entire infrared absorption spectrum of the parent allyl radical,  $CH_2CHCH_2$ ,  $\tilde{X}$   $^2A_2$ . The experimental IR spectrum is the black trace and each of the assigned allyl radical fundamentals is marked by a bullet (•). Colored red and offset above the IR absorption spectrum are the DFT calculated harmonic predictions  $\{\omega\}_i$  [UB3LYP/6-311-G(d, p)] from Table 2. In the construction of Figure 2, we have ignored the two IR inactive harmonic modes,  $\omega_8$  and  $\omega_9$ , and only plotted the active modes in order for an uncluttered comparison with our experiment. The lowest frequency CCC bending  $a_1$  mode  $\omega_7$  is not shown because it is predicted to appear at 429 cm $^{-1}$ , below the range of our MCT detector. The irreducible representations of allyl radical's vibrations are:

TABLE 3: Experimental Vibrational Frequencies ( $\nu$ /cm<sup>-1</sup>) and Relative Infrared Intensities (A/km mol<sup>-1</sup> as a Percent of the Base Peak) for the Matrix-Isolated X <sup>2</sup>A<sub>2</sub> Allyl Radical<sup>a</sup>

			C	H <sub>2</sub> CHC	CH <sub>2</sub>	CH <sub>2</sub> CDCH <sub>2</sub>			CD <sub>2</sub> CDCD <sub>2</sub>		
$\nu$		local mode description	$\nu$ /cm <sup>-1</sup>	A/%	polarity	$\nu$ /cm <sup>-1</sup>	A/%	polarity	$\nu$ /cm <sup>-1</sup>	A/%	polarity
1	$a_1$	CH <sub>2</sub> CHCH <sub>2</sub> asymmetric CH <sub>2</sub> st	3109	18	+	3106	38	+			
2		CH <sub>2</sub> CHCH <sub>2</sub> symmetric C CH <sub>2</sub> stretch	3052	5	+	3022	4		2285	4	+
3		CH <sub>2</sub> (C-H)CH <sub>2</sub> stretch	3027	2	+	2268	8	+	2215	4	+
4		symmetric CH <sub>2</sub> scissors	1478	6	+	1474	15	+	1263		
5		CH <sub>2</sub> -CH-CH <sub>2</sub> stretch + CH <sub>2</sub> scissors	1242	1	+	1220	4	+	1018	4	+
6		$CH_2$ - $CH$ - $CH_2$ stretch + $CH_2$ rock	na			1012	3	+	844	1	
7		CH <sub>2</sub> -CH-CH <sub>2</sub> bend	na			na			na		
8	$a_2$	CH <sub>2</sub> CHCH <sub>2</sub> out-of-phase umbrella	775 ?			na			na		
9		CH <sub>2</sub> -CH-CH <sub>2</sub> out- of-phase twist	na			na			na		
10	$b_1$	CH <sub>2</sub> (C-H)CH <sub>2</sub> out-of-plane bend	983	22	+	802	141	+	767	3	
11		CH <sub>2</sub> CHCH <sub>2</sub> in-phase umbrella	801	100	+	787	100	+	647	100	+
12		CH <sub>2</sub> -CH-CH <sub>2</sub> in-phase twist	510	14	+	511	22	+			
13	$b_2$	CH <sub>2</sub> CHCH <sub>2</sub> asymmetric CH <sub>2</sub> stretch	3107	4	_	3106	20	_	2372	2	_
14		CH <sub>2</sub> CHCH <sub>2</sub> symmetric CH <sub>2</sub> st	3020	3	_	3013	7	_	2209	6	_
15		asymmetric CH/CH <sub>2</sub> scissors	1464	17	_	1452	22	_	1387	13	_
16		asymmetric CH <sub>2</sub> -CH-CH <sub>2</sub> stretch + CH/CH <sub>2</sub> scissors	1390	1	_	1352	5	_	1062	7	_
17		asymmetric CH <sub>2</sub> −CH−CH <sub>2</sub> stretch ⊕ CH bend	1182		_	1010	1		900	7	_
18		asymmetric CH/CH <sub>2</sub> rock	na			845			689		

<sup>&</sup>lt;sup>a</sup> See Figures 2-13 for details. Na means not available.

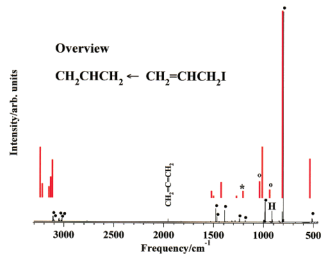


Figure 2. An overview of the infrared absorption spectrum of matrix isolated allyl radical. Allyl radicals are produced by the dissociation of CH<sub>2</sub>=CHCH<sub>2</sub>I in a hyperthermal nozzle at 1150 K. The solid, black trace is the experimental IR spectrum and the fundamental infrared modes  $\{\nu\}$  are marked by bullets (•). The DFT calculated [UB3LYP/ 6-311-G(d,p)] harmonic frequencies  $\{\omega\}$  are plotted as red sticks. Several of the harmonic modes are predicted to be very weak and have been scaled by a factor of 10 and are marked by a star (\*); weak  $\omega$ modes scaled by 100 are marked by ( ).

 $\Gamma_{\text{VIB}}(\text{CH}_2\text{CHCH}_2) = 7 \ a_1 \oplus 2 \ a_2 \oplus 3 \ b_1 \oplus 6 \ b_2$ . Consequently there are 15 red peaks marked on the spectrum in Figure 2 (18 modes of CH<sub>2</sub>CHCH<sub>2</sub>, less the 2 a<sub>2</sub> modes, less the lowfrequency  $a_1$  mode  $\omega_7$ ). Three of the harmonic modes ( $\omega_5$ ,  $\omega_6$ , and  $\omega_{17}$ ) are predicted to be very weak ( $A_i \leq 0.1 \text{ km mol}^{-1}$ ) and will be difficult to detect. Consequently we have scaled these three modes by factors of 10 (marked with a star, \*) and 100 (marked with a o) in Figure 2.

Qualitatively the comparison of the predicted harmonic modes and our experimental spectrum in Figure 2 is quite good. All the harmonic frequencies are predicted close to our assigned fundamentals although the computed harmonic infrared intensities are very erratic.

A matrix of allyl radicals was depleted by irradiation at 351 nm, exciting the blue edge of the (dissociative)  $\tilde{A}^2B_1 \leftarrow \tilde{X}^2A_2$ transition. Several sharp features are depleted in the IR spectrum

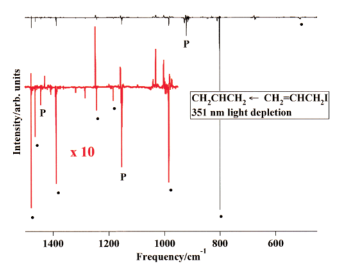
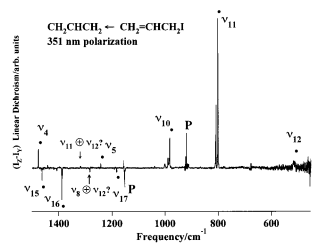


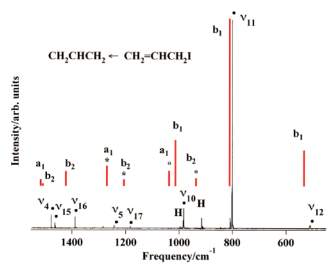
Figure 3. An infrared spectrum showing the depletion of matrix isolated CH2CHCH2 radicals "fingerprint region" upon bombardment by 351 nm laser light. Excitation of the A state of CH<sub>2</sub>CHCH<sub>2</sub> is known to dissociate the radical. Infrared fundamentals of the CH<sub>2</sub>CHCH<sub>2</sub> radical are marked by bullets (•). The weak portion of the spectrum between 950 - 1450 cm<sup>-1</sup> has been scaled by a factor of 10 and is plotted in red. An intense absorption of the allyl radical precursor, CH<sub>2</sub>= CHCH<sub>2</sub>I, is marked by P.

and are likely part of the CH2CHCH2 spectrum; these depletions are marked with a bullet (•) in Figure 3. The weak portion of Figure 3 between 1000 and 1400 cm<sup>-1</sup> has been expanded (in red) by a factor of 10 to clearly show the CH<sub>2</sub>CHCH<sub>2</sub> depletions.

Excitation with a polarized laser at 351 nm produces a linear dichroism spectrum  $(I_Z - I_Y)$  of allyl, Figure 4. The negatively polarized features must have b2 symmetry (vide supra) while the positive peaks can be polarized either  $a_1$  or  $b_1$ . With reference to the harmonic frequencies of Figure 2 and Table 2 and by matching the depletion and polarization spectra, we have assigned the CH<sub>2</sub>CHCH<sub>2</sub> vibrations in Figure 4. There are five features that have negative polarization. The bands at 1152 and 920 cm<sup>-1</sup> are strong features of CH<sub>2</sub>=CHCH<sub>2</sub>I and marked as P (for precursor). The negatively polarized bands at 1464, 1390, and 1182 cm<sup>-1</sup> are assigned as  $b_2$  fundamentals ( $\nu_{15}$ ,  $\nu_{16}$ , and  $\nu_{17}$ ). Since the linear dichroism spectra cannot distinguish between the a<sub>1</sub> and b<sub>1</sub> modes, we must rely on the computed

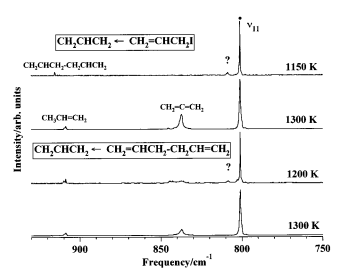


**Figure 4.** A linear dichroism spectrum of the CH<sub>2</sub>CHCH<sub>2</sub> radical following matrix-depletion by polarized 351 nm light. IR bands of CH<sub>2</sub>-CHCH<sub>2</sub> that are b<sub>2</sub> polarized will have a negative dichroism while IR features with a<sub>1</sub> and b<sub>1</sub> polarization will have a positive dichroism. Two weak features are attributed to combination bands of  $\nu_{12}$  of allyl radical; these are marked by  $(\nu_{11} \oplus \nu_{12})$  and  $(\nu_8 \oplus \nu_{12})$ . These absorptions are more clearly evident in Figure 5. Allyl fundamentals are marked by a bullet (•), while **P** is a precursor band.



**Figure 5.** A final comparison of allyl radical's fingerprint region. The DFT harmonic frequencies  $\{\omega\}$  in red reproduce the experimental fundamentals  $\{\nu\}$  marked by bullets  $(\bullet)$ . The intensities (\*) of the harmonic modes,  $\omega_5$  and  $\omega_{17}$ , have been scaled by a factor of 10. Two of the harmonic modes,  $\omega_6$  and  $\omega_{18}$ , are exceptionally weak and have had their intensities ( ) scaled by a factor of 100. We could not observe these two fundamentals,  $\nu_6$  and  $\nu_{18}$ .

harmonic frequencies in Table 2 to assign the spectra in Figure 4. The two bands at 1478 and 1242 cm<sup>-1</sup> are assigned as a<sub>1</sub> modes ( $\nu_4$  and  $\nu_5$ ) but the features at 983, 801, and 510 cm<sup>-1</sup> are clearly  $b_1$  modes ( $\nu_{10}$ ,  $\nu_{11}$ , and  $\nu_{12}$ ). We conjecture that the weak band at 1317 cm<sup>-1</sup> is likely a combination band, ( $\nu_{11} \oplus$  $\nu_{12}$ ). Such a transition would be polarized ( $b_1 \otimes b_1$ ) or  $a_1$  which is compatible with the LD spectrum, Figure 4. The weak, negatively polarized feature at 1285 cm<sup>-1</sup> could be possibly be a combination band of  $(\nu_8 \oplus \nu_{12})$ . Such a band would be polarized ( $a_2 \otimes b_1$ ) or  $b_2$  with a negatively polarized absorption in Figure 4. This implies an assignment of 775 cm<sup>-1</sup> for the a<sub>2</sub> mode  $\nu_8$ , which has never been assigned before. The predicted harmonic mode  $\omega_8$  from Table 2 is 786 cm<sup>-1</sup>. Figure 5 is a summary of the entire fingerprint region of the CH<sub>2</sub>CHCH<sub>2</sub> radical; this figure is a color-coded comparison of our experimental spectra (in black) and the red sticks which locate the



**Figure 6.** The feature at 809 cm<sup> $^{-1}$ </sup> in Figures 2–4 is not a fundamental of allyl radical. This band appears in matrices produced by thermal dissociation (1150 K) of CH<sub>2</sub>=CHCH<sub>2</sub>I or CH<sub>2</sub>=CHCH<sub>2</sub>-CH<sub>2</sub>CH=CH<sub>2</sub>. If the hyperthermal nozzle is heated to higher temperatures (1300 K), the 809 cm<sup> $^{-1}$ </sup> feature disappears and signals from the radical/radical disproportionation appear (CH<sub>2</sub>=C=CH<sub>2</sub> and CH<sub>2</sub>=CHCH<sub>3</sub>). However the allyl fundamental at 801 cm<sup> $^{-1}$ </sup> ( $\nu_{11}$ ) persists.

harmonic frequencies. The experimental values are collected together in Table 3. In so far as we can measure, the ordering of the harmonic frequencies,  $\{\omega\}$  in red, match our experimental assignments exactly. We cannot detect the CH<sub>2</sub>CHCH<sub>2</sub> modes  $\nu_6$  and  $\nu_{18}$  which are predicted (Table 2) to be extremely weak. There are two absorptions in the fingerprint region that clearly belong to CH<sub>2</sub>=CHCH<sub>2</sub>-CH<sub>2</sub>CH=CH<sub>2</sub> (**H**).

There is an intense feature near at 809 cm<sup>-1</sup> near  $\nu_{11}$  in Figure 4. Because this line has been consistently measured in many allyl radical studies, we believe that it must be attributed to either the CH2CHCH2 radical or species associated with it. Although we could not determine the carrier of the 809 cm<sup>-1</sup> line, we believe that it is not a CH<sub>2</sub>CHCH<sub>2</sub> radical fundamental nor combination band as has been suggested in earlier studies. We produced CH<sub>2</sub>CHCH<sub>2</sub> radical from both CH<sub>2</sub>=CHCH<sub>2</sub>I at 1150 and 1300 K and from CH<sub>2</sub>=CHCH<sub>2</sub>-CH<sub>2</sub>CH=CH<sub>2</sub> at 1200 and 1300 K. Spectra of the region are shown in Figure 6. In the higher temperature production of CH<sub>2</sub>CHCH<sub>2</sub> radical, the 809 cm<sup>-1</sup> is no longer present although other allyl radical features persist. IR signals of the disproportionation products, CH<sub>3</sub>CH=CH<sub>2</sub> and CH<sub>2</sub>=C=CH<sub>2</sub>, are clearly identified. This 809 cm<sup>-1</sup> is present in both lower temperature spectra. Also note that the 809 cm<sup>-1</sup> has a dramatically different intensity pattern in polarization spectrum in Figure 4 than it had in the depletion spectrum in Figure 3. In earlier studies, researchers produced CH<sub>2</sub>CHCH<sub>2</sub> radical in low-pressure pyrolysis at 1100 to 1200 K only. We believe that this line is a mode of some intermediate species in the thermal production of CH<sub>2</sub>CHCH<sub>2</sub> radical.

Figure 7 shows the depletion of the CH portion of the IR spectrum of CH<sub>2</sub>CHCH<sub>2</sub>. We expect to find 5 CH absorption features and Figure 7 clearly shows five bands. The (eventual) fundamentals are shown as bullets (•). Figure 8 is the LD spectrum of allyl that results from polarization of the matrix with 351 nm light. By symmetry we know that the five CH stretches transform like (3  $a_1 \oplus 2 b_2$ ) modes. Consequently we assign the three  $a_1$  CH stretches  $\nu_1$ ,  $\nu_2$ , and  $\nu_3$  (see Table 3). Earlier high-resolution IR absorption studies<sup>25,26</sup> of the highest ( $a_1$ ,  $b_2$ ) CH stretching pair,  $\nu_1$  and  $\nu_{13}$ , have observed these features at 3113.98488  $\pm$  0.00089 cm<sup>-1</sup> and 3110.59857  $\pm$ 

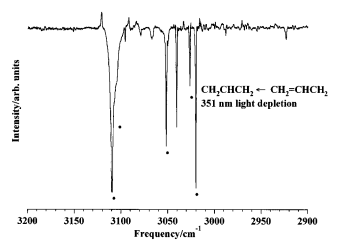


Figure 7. An infrared spectrum showing the depletion of matrix isolated CH<sub>2</sub>CHCH<sub>2</sub> radicals "CH region" upon bombardment by 351 nm laser light. Infrared fundamentals of the CH2CHCH2 radical are marked by bullets (•).

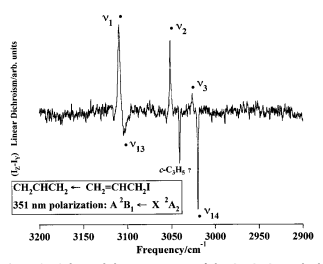


Figure 8. A linear dichroism spectrum of the CH2CHCH2 radical following matrix-depletion by polarized 351 nm light. IR bands of CH<sub>2</sub>-CHCH<sub>2</sub> that are b<sub>2</sub> polarized will have a negative dichroism while IR features with a<sub>1</sub> and b<sub>1</sub> polarization will have a positive dichroism. Allyl fundamentals are marked by a bullet (•); the strong feature at 3040 cm<sup>-1</sup> is not an allyl radical mode.

0.00036 cm<sup>-1</sup>, respectively (see Table 1). Rotational analysis clearly established that  $v_1$  is a b-type band consistent with an  $a_1$  transition while  $v_{13}$  is an a-type band or  $b_2$  mode. The harmonic frequencies (Table 2) predict the  $(\omega_1, \omega_{13})$  pair at 3235 and 3232 cm<sup>-1</sup>. Consequently we assign the first two matrix bands in Figure 8 as  $v_1 = 3109 \text{ cm}^{-1}$  and  $v_{13} = 3107 \text{ cm}^{-1}$ . We have now assigned all of the CH stretches but for  $\nu_{14}$ . We appeal to the predicted harmonic frequencies in Table 2 where  $\omega_{14}$  is estimated to be 3134 cm<sup>-1</sup>; consequently we assign the intense, negatively polarized band at 3020 cm $^{-1}$  as  $\nu_{14}$ . This leaves us with the strong, negatively polarized feature at 3040 cm<sup>-1</sup> in Figure 8.

The line at 3040 cm<sup>-1</sup> is not consistent with being a combination band of allyl radical fundamentals. There are no combinations of two fundamentals which result in a mode at 3040 cm<sup>-1</sup> and that would exhibit b<sub>2</sub> type LD. Yet this line is consistently found all matrix CH2CHCH2 radical spectra. Consequently we consider an isomer of CH2CHCH2 radical, the cyclopropyl radical (c-C<sub>3</sub>H<sub>5</sub>). We conjecture that this mode is one of the most intense a" modes of the c-C<sub>3</sub>H<sub>5</sub> radical,  $\nu_{12}$ . Recently Davis<sup>52</sup> studied the high-resolution IR absorption

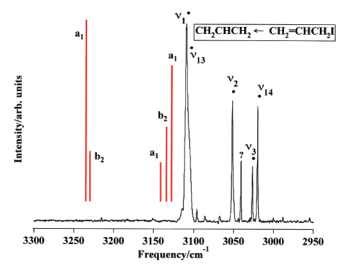


Figure 9. A final comparison of allyl radical's CH stretching region. The DFT harmonic frequencies  $\{\omega\}$  in red reproduce the experimental fundamentals  $\{v\}$  marked by bullets  $(\bullet)$ .

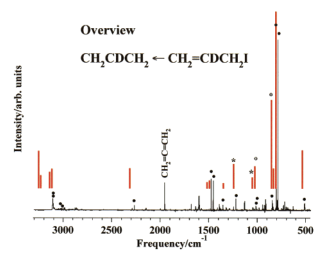
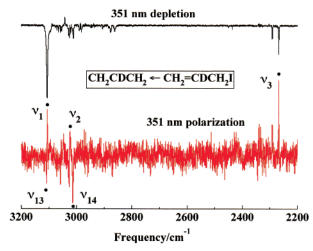


Figure 10. An overview of the infrared absorption spectrum of matrix isolated allyl radical. Allyl radicals are produced by the dissociation of CH<sub>2</sub>=CDCH<sub>2</sub>I in a hyperthermal nozzle at 1150 K. The solid, black trace is the experimental IR spectrum and the fundamental infrared modes  $\{\nu\}$  are marked by bullets (•). The DFT calculated [UB3LYP/ 6-311-G(d,p)] harmonic frequencies  $\{\omega\}$  are plotted as red sticks. Several of the harmonic modes are predicted to be very weak and have been scaled by a factor of 10; these weak  $\{\omega\}$  transitions are marked by a star (\*). Two harmonic modes are exceptionally weak and have been scaled by a factor of 100 are marked by ().

spectra of a pulsed electric discharge of cyclopropyl bromide wherein he observed a cyclopropyl C-H stretch at 3040.8458 cm<sup>-1</sup>; this he identified as a c-type band or a" mode. Our unassigned matrix band is also at 3040 cm<sup>-1</sup> and this is a significantly smaller gas-to-matrix shift than the 4 or 5 cm<sup>-1</sup> shift that has been observed for allyl radical C-H stretches. However, different species could certainly have different shifts.

Figure 9 is a direct comparison of the UB3LYP/6-311-G(d,p) harmonic modes (red sticks) with our assigned CH fundamentals for CH<sub>2</sub>CHCH<sub>2</sub>. The harmonic predictions are very close to our experimental findings except for the  $(\nu_2, \nu_3, \text{ and } \nu_{14})$  order.

In addition to the parent allyl radical, we have studied its  $d_1$ isotopomer, CH<sub>2</sub>CDCH<sub>2</sub>, and the d<sub>5</sub> isotopomer, CD<sub>2</sub>CDCD<sub>2</sub>. Figure 10 is a overview of the CH<sub>2</sub>CDCH<sub>2</sub> infrared absorption spectrum. The black trace is the experimental spectrum and the assigned fundamentals are marked by bullets (•). The red sticks are the harmonic frequencies that result from a UB3LYP/6-



**Figure 11.** An overplot of the 351 nm depletion spectrum of matrix isolated  $CH_2CDCH_2$  radical (black, on top) with the linear dichroism spectrum (red, on bottom). The  $CH_2CDCH_2$  fundamentals  $\{\nu\}$  are marked by bullets  $(\bullet)$ .

311-G(d,p) electronic structure calculation. Four of the harmonic frequencies ( $\omega_5$ ,  $\omega_6$ ,  $\omega_{17}$ , and  $\omega_{18}$ ) have very weak intensities and have been scaled; those bands scaled by a factor of 10 have been marked by an star (\*) while those by a factor of 100 by (). The final, assigned CH<sub>2</sub>CDCH<sub>2</sub> fundamentals are collected in Table 3.

Figure 11 shows the CH/D stretching region of the  $d_1$  isotopomer. The matrix-isolated CH<sub>2</sub>CDCH<sub>2</sub> radical is photodissociated with 351 nm radiation in the top of Figure 11; the depleted fundamentals are marked with bullets (•). To confirm the assignment of these CH<sub>2</sub>CDCH<sub>2</sub> fundamentals, the matrix was partially depleted with polarized 351 nm light. The linear dichroism spectrum of the resulting polarized matrix is shown at the bottom of Figure 11. The CH<sub>2</sub>CDCH<sub>2</sub> radial has four CH stretches, 2  $a_1 \oplus 2 b_2$ , and one CD stretch of  $a_1$  symmetry. Figure 11 clearly demonstrates the identity of the CH  $a_1$  modes,  $\nu_1$  and  $\nu_2$ , as well as the  $b_2$  pair,  $\nu_{13}$  and  $\nu_{14}$ . Simple harmonic reasoning would predict that the CH<sub>2</sub>(C-D)CH<sub>2</sub> stretching frequency would be shifted to a lower frequency, 3027 cm<sup>-1</sup>/ $\sqrt{2}$  or roughly 2140 cm<sup>-1</sup>. Figure 11 identifies the CD stretch as the  $a_1$  mode,  $\nu_3$ , shifted down to 2268 cm<sup>-1</sup>.

The IR spectrum of the  $d_5$  isotopomer, CD<sub>2</sub>CDCD<sub>2</sub>, is displayed in Figure 12. The CD stretching modes are assigned in Figure 13. The 351 nm photodepleted CD fundamentals are on the top of Figure 13 while the bottom is the linear dichroism spectrum of a 351 nm polarized matrix sample. We can only identify four of the CD fundamentals. One of the  $a_1$  modes,  $\nu_1$ , is clearly missing. It is surprising that we cannot observe  $\nu_1$  of CD<sub>2</sub>CDCD<sub>2</sub> since we compute  $\omega_1$ (CD<sub>2</sub>CDCD<sub>2</sub>) to be a strong band (Table 2).

## **IV. Conclusions**

The set of experimental frequencies for the allyl radical (Table 3) are reasonably reproduced by the UB3LYP/6-311-G(d,p) harmonic frequencies in Table 2. Figures 2, 5, 9, 10, and 12 provide a direct, visual comparison of set of experimental frequencies  $\{\nu\}$  with the corresponding harmonic ones  $\{\omega\}$ . A useful test of the computed  $\{\omega\}$  values is to check the predicted isotope shifts. A direct comparison of the experimental frequency shifts of the  $d_0$ ,  $d_1$ , and  $d_5$  isotopomers  $\{\Delta\nu\}$  with the corresponding harmonic shifts  $\{\Delta\omega\}$  is interesting. For example, Table 3 indicates that the intense,  $b_1$  CH<sub>2</sub> in-phase-twisting mode  $\nu_{11}$  shifts by 14 cm<sup>-1</sup> in the  $d_1$  isotopomer [801–787 cm<sup>-1</sup>]

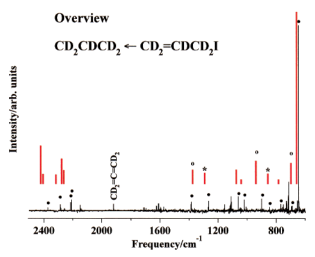
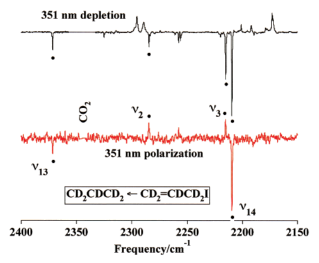


Figure 12. An overview of the infrared absorption spectrum of matrix isolated allyl radical. Allyl radicals are produced by the dissociation of CD<sub>2</sub>=CDCD<sub>2</sub>I in a hyperthermal nozzle at 1150 K. The solid, black trace is the experimental IR spectrum and the fundamental infrared modes  $\{\nu\}$  are marked by bullets (•). The DFT calculated [UB3LYP/6-311-G(d,p)] harmonic frequencies  $\{\omega\}$  are plotted as red sticks. Several of the harmonic modes are predicted to be very weak and have been scaled by a factor of 10; these weak  $\{\omega\}$  transitions are marked by a star (\*). Three harmonic modes are exceptionally weak and have been scaled by a factor of 100 are marked by ().



**Figure 13.** An overplot of the 351 nm depletion spectrum of matrix isolated  $CD_2CDCD_2$  radical (black, on top) with the linear dichroism spectrum (red, on bottom). The  $CD_2CDCD_2$  fundamentals  $\{\nu\}$  are marked by bullets ( $\bullet$ ). The  $a_1$  fundamental of allyl- $d_5$   $\nu_1$  could not be detected.

and by  $154 \text{ cm}^{-1}$  in the  $d_5$  isotopomer [ $801-647 \text{ cm}^{-1}$ ]; Table 2 indicates that the predicted shifts in  $\omega_{11}$  are 3 and  $154 \text{ cm}^{-1}$ , respectively. Table 4 has such a comparison for all of the infrared fundamentals and the results are generally satisfactory. Although the *absolute* harmonic frequencies are off by roughly 2%-4%, Table 4 indicates that the relative isotopic shifts are faithfully reproduced. However the infrared intensities  $\{A\}$  for the harmonic modes predicted by UB3LYP/6-311-G(d,p) do not correlate very well with the observed intensities. This may be the fault of the small basis set that we have used to compute  $\{\omega\}$ .

Before we can make a final recommendation for the gasphase vibrational frequencies of allyl, we must estimate the magnitude of the gas-to-matrix frequency shifts. Jacox has reviewed the matrix shifts for a large number of diatomic and small polyatomic free radicals and ions trapped in Ne and Ar

TABLE 4: Isotopic Shifts of Allyl Radicals (Tables 2 and 3)

			$\Delta(d_0-d_1)$		$\Delta(d_0)$	$-d_5$ )
ν		local mode description	$\Delta \nu$ /cm <sup>-1</sup>	${\Delta\omega}/{\rm cm}^{-1}$	$\Delta \nu$ /cm <sup>-1</sup>	${\Delta\omega}/{\rm cm}^{-1}$
1	$a_1$	CH <sub>2</sub> CHCH <sub>2</sub> asymmetric CH <sub>2</sub> stretch	3	2		825
2		CH <sub>2</sub> CHCH <sub>2</sub> symmetric CH <sub>2</sub> stretch	30	1	767	825
3		$CH_2(C-H)CH_2$ stretch	759	817	812	850
4		symmetric CH <sub>2</sub> scissors	4	3	215	226
5		$CH_2$ - $CH$ - $CH_2$ stretch + $CH_2$ scissors	22	24	224	230
6		$CH_2-CH-CH_2$ stretch $+$ $CH_2$ rock		1		181
7		CH <sub>2</sub> -CH-CH <sub>2</sub> bend		4		79
8	$a_2$	CH <sub>2</sub> CHCH <sub>2</sub> out-of-phase umbrella				
9		CH <sub>2</sub> -CH-CH <sub>2</sub> out-of-phase twist				
10	$b_1$	$CH_2(C-H)CH_2$ out-of-plane bend	181	198	216	230
11		CH <sub>2</sub> CHCH <sub>2</sub> in-phase umbrella	14	3	154	154
12		CH <sub>2</sub> -CH-CH <sub>2</sub> in-phase twist	-1	1		124
13	$b_2$	CH <sub>2</sub> CHCH <sub>2</sub> asymmetric CH <sub>2</sub> stretch	1	0	735	824
14		CH <sub>2</sub> CHCH <sub>2</sub> symmetric CH <sub>2</sub> stretch	7	0	811	861
15		asymmetric CH/CH <sub>2</sub> scissors	12	17	77	136
16		asymmetric CH <sub>2</sub> -CH-CH <sub>2</sub> stretch + CH/CH <sub>2</sub> scissors	38	75	328	348
17		asymmetric CH <sub>2</sub> -CH-CH <sub>2</sub> stretch + CH bend	172	171	282	268
18		asymmetric CH/CH <sub>2</sub> rock		85		237

TABLE 5: Recommended Vibrational Frequencies (v/cm<sup>-1</sup>) for  $\tilde{X}^2A_2$  Allyl Radical

ν		local mode description	$\nu/\mathrm{cm}^{-1}$	ref
1	$a_1$	CH <sub>2</sub> CHCH <sub>2</sub> asymmetric CH <sub>2</sub> stretch	3114	25 and 26
2		CH <sub>2</sub> CHCH <sub>2</sub> symmetric CH <sub>2</sub> stretch	3052	this work
3		CH <sub>2</sub> (C-H)CH <sub>2</sub> stretch	3027	this work
4		symmetric CH <sub>2</sub> scissors	1478	this work
5		$CH_2$ - $CH$ - $CH_2$ stretch + $CH_2$ scissors	1242	this work
6		$CH_2$ - $CH$ - $CH_2$ stretch + $CH_2$ rock	1068	35
7		CH <sub>2</sub> -CH-CH <sub>2</sub> bend	425	9 and 10
8	$a_2$	CH <sub>2</sub> CHCH <sub>2</sub> out-of-phase umbrella	775	this work
9		CH <sub>2</sub> -CH-CH <sub>2</sub> out-of-phase twist	549	35
10	$b_1$	CH <sub>2</sub> (C-H)CH <sub>2</sub> out-of-plane bend	983	this work
11		CH <sub>2</sub> CHCH <sub>2</sub> in-phase umbrella	802	27
12		CH <sub>2</sub> -CH-CH <sub>2</sub> in-phase twist	510	this work
13	$b_2$	CH <sub>2</sub> CHCH <sub>2</sub> asymmetric CH <sub>2</sub> stretch	3111	25 and 26
14		CH <sub>2</sub> CHCH <sub>2</sub> symmetric CH <sub>2</sub> stretch	3020	this work
15		asymmetric CH/CH <sub>2</sub> scissors	1464	this work
16		asymmetric CH <sub>2</sub> -CH-CH <sub>2</sub>	1390	this work
		stretch + CH/CH <sub>2</sub> scissors		
17		asymmetric CH <sub>2</sub> -CH-CH <sub>2</sub>	1182	this work
		stretch + CH bend		
18		asymmetric CH/CH <sub>2</sub> rock		

matrices.53,54 She concluded that for polyatomic free radicals in Ar matrices the frequency shift is generally less than 1% and usually to the red. Rotationally resolved frequencies are available<sup>25–27</sup> for three of allyl's fundamentals:  $\nu_1$  (3113.98488  $\pm$  0.00089 cm<sup>-1</sup>),  $\nu_{11}$  (801.71909  $\pm$ 0.00037 cm<sup>-1</sup>), and  $\nu_{13}$  $(3110.59857 \pm 0.00036 \text{ cm}^{-1})$ . The matrix values (from this study) are  $\nu_1$  (3109 cm<sup>-1</sup>),  $\nu_{11}$  (801 cm<sup>-1</sup>), and  $\nu_{13}$  (3107 cm<sup>-1</sup>). So for the allyl radical, the gas-to-matrix shifts for these three modes are  $\Delta \nu_1$  (4 cm<sup>-1</sup>),  $\Delta \nu_{11}$  (0 cm<sup>-1</sup>), and  $\Delta \nu_{13}$  (3 cm<sup>-1</sup>). Consequently we believe that all of the matrix frequencies for the allyl radical are within ≤1% of the true, gas-phase frequencies.

Finally we recommend a set of vibrational frequencies for the CH2CHCH2 radical in Table 5. High-resolution laser absorption spectroscopy<sup>25,26</sup> has established the CH stretching frequencies,  $\nu_1$  and  $\nu_{13}$ , as well as<sup>27</sup> the out-of-plane umbrella mode  $\nu_{11}$ . Resonance Raman studies<sup>35–37</sup> have observed the  $a_1$ CCC stretching mode  $\nu_6$  and the  $a_2$  out-of-phase CH<sub>2</sub> twist  $\nu_9$ . The low-frequency CCC bending mode  $v_7$  has been observed<sup>9,10</sup> in the negative ion photodetachment of the allylic anion:  $CH_2CHCH_2^- \rightarrow CH_2CHCH_2 + e^-$ . All the rest of the frequencies in Table 5 are from this matrix infrared study. Our polarization spectroscopy has provided new information that makes many of CH2CHCH2 radical's fundamental assignments

more definite. The pulsed nozzle techniques of this paper permit us to produce clean samples of the target allyl radicals. We observe narrow IR absorption lines for CH<sub>2</sub>CHCH<sub>2</sub> and these can be measured with high accuracy; consequently, apart from the unknown matrix shifts, the fundamentals are determined to high accuracy.

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#### References and Notes

- (1) Salem, L. Pi Electron Theory of Organic Chemistry; W. A. Benjamin: New York City, 1962.
- (2) Salem, L.; Rowland, C. Angew. Chem., Int. Ed. Engl. 1972, 11,
- (3) Ellison, G. B.; Davico, G. E.; Bierbaum, V. M.; DePuy, C. H. Int. J. Mass Spectrosc. Ion Processes 1996, 156, 109.
- (4) The precise formulation of allyl's resonance energy  $E_{Res}(CH_2-$ CHCH<sub>2</sub>) is a subtle issue. This matter is discussed in Section 5 of Ellison, Davico, Bierbaum, and DePuy. One measure of  $E_{Res}(CH_2CHCH_2)$  is the rotational barrier of the allylic radical, CH2CHCHD, which yields a value of  $64 \pm 4$  kJ mol<sup>-1</sup>. The difference between the bond energies for propane and propene is  $[(97 \pm 2) - (87.4 \pm 0.4)] = 10 \pm 2 \text{ kcal mol}^{-1} \text{ or } 40 \pm 9$ kJ mol-
- (5) Houle, F. A.; Beauchamp, J. L. J. Am. Chem. Soc. 1978, 100, 3290. (6) Wu, J. C.; Li, R. H.; Chang, J. L.; Chen, Y. T. J. Chem. Phys. 2000, 113, 7286.
- (7) Schultz, T.; Clarke, J. S.; Gilbert, T.; Deyerl, H. J.; Fischer, I. Faraday Discuss. 2000, 17.
  - (8) Gilbert, T.; Fischer, I.; Chen, P. J. Chem. Phys. 2000, 113, 561.
- (9) Oakes, J. M.; Ellison, G. B. J. Am. Chem. Soc. 1984, 106, 7734. (10) Wenthold, P. G.; Polak, M. L.; Lineberger, W. C. J. Phys. Chem.
- 1996, 100, 6920. (11) Baghol-Vayjooee, M. H.; Benson, S. W. J. Am. Chem. Soc. 1979, 101, 2838.

- (12) Friderichsen, A. V.; Radziszewski, J. G.; Nimlos, M. R.; Winter, P. R.; Dayton, D. C.; David, D. E.; Ellison, G. B. *J. Am. Chem. Soc.* **2001**, *123*, 1977.
- (13) Friderichsen, A. V. Infrared Spectroscopy of Matrix-Isolated Aromatic Hydrocarbon Radicals. Ph.D. Thesis, Chemistry, University of Colorado, Boulder, CO, 2001.
- (14) Kohn, D. W.; Clausberg, H.; Chen, P. Rev. Sci. Instrum. 1992, 63,
- (15) Chen, P. The Spectroscopy and Photochemistry in Supersonic Jets. Ph.D. Thesis, Chemistry, Yale, 1987.
- (16) Brown, A. L.; Dayton, D. C.; Nimlos, M. R.; Daily, J. W. Chemosphere 2001, 42, 663.
- (17) Mamyrin, B. A.; Karataev, V. I.; Shmikk, D. V.; Zagulin, V. A. Sov. Phys.-JEPT 1973, 37, 45.
  - (18) Kung, A. H.; Young, J. F.; Harris, S. E. Appl. Phys. Lett. 1973.
- (19) Berkowitz, J. *Photoabsorption, Photoionization and Photoelectron Spectroscopy*; Academic Press: New York, 1979.
  - (20) Becke, A. D. J. Phys. Chem. 1993, 98, 5648
  - (21) Becke, A. D. J. Chem. Phys. 2000, 112, 4020.
- (22) Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Zakrzewski, V. G.; Montgomery, J. J. A.; Stratmann, R. E.; Burant, J. C.; Dapprich, S.; Millam, J. M.; Daniels, A. D.; Kudin, K. N.; Strain, M. C.; Farkas, O.; Tomasi, J.; Barone, V.; Cossi, M.; Cammi, R.; Mennucci, B.; Pomelli, C.; Adamo, C.; Clifford, S.; Ochterski, J.; Petersson, G. A.; Ayala, P. Y.; Cui, Q.; Morokuma, K.; Malick, D. K.; Rabuck, A. D.; Raghavachari, K.; Foresman, J. B.; Cioslowski, J.; Ortiz, J. V.; Baboul, A. G.; Stefanov, B. B.; Liu, G.; Liashenko, A.; Piskorz, P.; Komaromi, I.; Gomperts, R.; Martin, R. L.; Fox, D. J.; Keith, T.; Al-Laham, M. A.; Peng, C. Y.; Nanayakkara, A.; Gonzalez, C.; Challacombe, M.; Gill, P. M. W.; Johnson, B.; Chen, W.; Wong, M. W.; Andres, J. L.; Gonzalez, C.; Head-Gordon, M.; Replogle, E. S.; Pople, J. A. GAUSSIAN98, revsion A.7, Gaussian, Inc.: Pittsburgh, PA, 1998.
- (23) Pedley, J. B.; Naylor, R. D.; Kirby, S. P. Thermochemical Data of Organic Compounds, 3rd ed.; University of Sussex Press: Cambridge, U.K., 1986.
- (24) Berkowitz, J.; Ellison, G. B.; Gutman, D. J. Phys. Chem. 1994, 98, 2744.
  - (25) Uy, D.; Davis, S.; Nesbitt, D. J. J. Chem. Phys. 1998, 109, 7793.
- (26) DeSain, J. D.; Thompson, R. I.; Sharma, S. D.; Curl, R. F. J. Chem. Phys. **1998**, 109, 7803.
- (27) Hirota, E.; Yamada, C.; Okunishi, M. J. Chem. Phys. 1992, 97, 2963.
  - (28) Hirota, E. J. Mol. Struct. 1994, 320, 75.

- (29) Heller, C.; Cole, T. J. Chem. Phys. 1962, 37, 243.
- (30) Krusic, P. J.; Kochi, J. K. J. Am. Chem. Soc. 1968, 90, 7.
- (31) Hudgens, J. W.; Dulcey, C. S. J. Phys. Chem. 1985, 89, 1505.
  (32) Minsek, D. W.; Blush, J. A.; Chen, P. J. Phys. Chem. 1992, 96, 2025.
  - (33) Minsek, D. W.; Chen, P. J. Phys. Chem. 1993, 97, 7.
  - (34) Sappey, A. D.; Weisshaar, J. C.J. Phys. Chem. 1987, 91, 3731.
- (35) Getty, J. D.; Burnmeister, M. J.; Westre, S. G.; Kelly, P. B. J. Am. Chem. Soc. 1991, 113, 801.
  - (36) Getty, J. D.; Kelly, P. B. Chem. Phys. 1992, 168, 357.
- (37) Getty, J. D.; Liu, X.; Kelly, P. B. Chem. Phys. Lett. 1993, 201, 236.
  - (38) Liu, X.; Getty, J. D.; Kelly, P. B. J. Chem. Phys. 1993, 99, 1522.
- (39) Holtzhauer, K.; Cometta-Morini, C.; Oth, J. F. M. J. Phys. Org. Chem. 1990, 3, 219.
  - (40) Huang, J. W.; Graham, W. R. M. J. Chem. Phys. 1990, 93, 1583.
- (41) Maier, G.; Reisenauer, H. P.; Rohde, B.; Dehnicke, K. Chem. Ber. 1983, 116, 732.
- (42) Mal'tsev, A. K.; Korolov, V. A.; Nefedov, O. M. Bull. Acad. Sci. USSR, Chem. Ser. 1983, 31, 2131.
- (43) Mal'tsev, A. K.; Korolov, V. A.; Nefedov, O. M. Bull. Acad. Sci. USSR, Chem. Ser. 1984, 33, 510.
  - (44) Currie, C. L.; Ramsay, D. A. J. Chem. Phys. 1966, 45, 488.
  - (45) Callear, A. B.; Lee, H. K. Trans. Faraday Soc. 1968, 64, 308.
- (46) Ha, T. K.; Baumann, H.; Oth, J. F. M. J. Chem. Phys. 1986, 85, 1438.
- (47) Blush, J. A.; Minsek, D. A.; Chen, P. J. Phys. Chem. 1992, 96, 10150.
- (48) Michl, J.; Thulstrup, E. W. Spectroscopy with Polarized Light; VCH Publishers, Inc.: New York, 1996.
- (49) Ball, D. W.; Pong, R. G. S.; Kafafi, Z. H. J. Am. Chem. Soc. 1993, 115, 2864.
- (50) Cyvin, S. J. J. Chem. Phys. 1958, 29, 583.
- (51) Woodfin, E. S.; Fletcher, W. H. J. Mol. Spectrosc. 1957, 1, 95.
- (52) Davis, S. High-Resolution Infrared Spectroscopy of Slit-Jet Cooled Transient Molecules: From van der Waals clusters, to Hydrogen Bound Dimers to Small Organic Radicals. Ph.D. Thesis, Chemistry, University of Colorado, Boulder, CO, 1999.
  - (53) Jacox, M. E. J. Mol. Spectrosc. 1985, 113, 286.
  - (54) Jacox, M. E. Chem. Phys. 1994, 189, 149.
- (55) Jacox, M. E. Vibrational and Electronic Energy Levels of Polyatomic Transient Molecules; American Chemical Society: Gaithersburg, Maryland, 1994.