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The reactions of chlorine and hydrogen atoms with trans-1,3-butadiene in solid para-hydrogen (p-H2) were investigated with infrared (IR) absorption spectra. When a p-H2 matrix containing Cℓ2 and trans-1,3-butadiene was irradiated with ultraviolet light at 356 nm, intense lines at 650.3, 809.0, 962.2, 1240.6 cm−1, and several weaker ones due to the trans-1-chloromethylallyl radical, •(CH2CHCH)CH2Cℓ, appeared. Observed wavenumbers and relative intensities agree with the anharmonic vibrational wavenumbers and IR intensities predicted with the B3PW91/6-311++g(2d, 2p) method. That the Cℓ atom adds primarily to the terminal carbon atom of trans-1,3-butadiene is in agreement with the path of minimum energy predicted theoretically, but in contrast to the reaction of Cℓ + propene in solid p-H2 [J. Amicangelo and Y.-P. Lee, J. Phys. Chem. Lett. 1, 2956 (2010)] in which the addition of Cℓ to the central C atom is favored, likely through steric effects in a p-H2 matrix. A second set of lines, intense at 781.6, 957.9, 1433.6, 2968.8, 3023.5, 3107.3 cm−1, were observed when the UV-irradiated Cℓ2/trans-1,3-butadiene/p-H2 matrix was further irradiated with IR light from a SiC source. These lines are assigned to the trans-1-methylallyl radical, •(CH2CHCH)CH3, produced from reaction of 1,3-butadiene with a H atom resulted from the reaction of Cℓ atoms with solid p-H2 exposed to IR radiation. © 2012 American Institute of Physics.

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I. INTRODUCTION

Unsaturated hydrocarbon radicals are important intermediates in combustion reactions; their reaction kinetics have been investigated extensively to understand combustion processes.1–4 The dissociation and mechanism of isomerization of C3H5 and C4H7 radicals, for example, have been studied both experimentally5–12 and theoretically13–17 to understand the branching ratios among various dissociation channels as a function of excitation energy. The reactions of alkynes (CnH2n) with atomic chlorine (Cℓ) play important roles in the chemistry of the polar troposphere;18–20 these reactions have also been studied extensively with theory.21–23 The two most significant initial processes of Cℓ + CnH2n reactions are the addition of a Cℓ atom to form a chloroalkyl radical (CℓCnH2nCℓ) and the abstraction of a H atom by the Cℓ atom to form an alkyl radical (CnH2n−1) and HCl.24 In most cases, the infrared (IR) spectra of the chloroalkyl radical remain unidentified because of the instability of these species and the final products are generally the dichloroalkanes when molecular chlorine reacts with alkenes in a solution or in the gaseous phase.25,26

Photochemical reactions in solid para-hydrogen (p-H2) at low temperatures have generated considerable interest in recent years.27,28 The p-H2 matrix is considered to be a quantum solid as the amplitude of the zero-point lattice vibrations is a large fraction of the lattice spacing.29 One unique feature of the p-H2 matrix is that it is softer than noble-gas matrices, such as Ar and Ne, such that the matrix cage effect is diminished. A recent study of matrix-isolated water dimers indicated that, for a given monomer concentration, the dimer concentration is significantly higher in solid Ne (or Ar) than in solid p-H2.30 However, because the photofragments can diffuse in solid p-H2, formation of complexes prior to photolysis might not be the critical criterion for bimolecular reaction to take place in a p-H2 matrix; the more critical property is the diminished cage effect so that the fragments can escape from the original cage to become isolated or react with other guest molecules. Furthermore, one can always perform moderate annealing after deposition to enhance the formation of complexes. As a consequence, photoproducts such as Cℓ atom and CH3 radical produced on ultraviolet (UV) irradiation of Cℓ2 (Ref. 31) and CH3I,32–34 respectively, become excellent sources for studying bimolecular reactions in solid p-H2. We have employed this method to investigate the reactions of Cℓ or CH3 with several small molecules in solid p-H2. For instance, the reaction Cℓ + CS2 in p-H2 produced CℓCS, CℓCS, and CℓSC,35 Cℓ + propene (C3H6 or CH2=CHCH3) produced the 2-chloropropyl radical (•CH2CHCℓCH3),36 and CH3 + SO2 produced CH3SO2Cℓ.34

In the reaction Cℓ + C3H6 in solid p-H2, we observed IR lines ascribable to only the 2-chloropropyl radical, not the 1-chloropropyl radical, •CH(CH2Cℓ)CH3. This result is in contrast to the reported gaseous experiments in which

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formation of 1-chloropropyl is favored over 2-chloropropyl at least six fold.\textsuperscript{37} High-level quantum-chemical computations on the reaction $\text{C}^{\ell} + \text{C}_2\text{H}_4$, however, predicted that the addition of the $\text{C}^{\ell}$ atom to either carbon atom involving the double bond is barrierless, and the formation of 1-chloropropyl is energetically favored by less than 1 kJ mol$^{-1}$ over that of 2-chloropropyl, indicating little preference for the addition of $\text{C}^{\ell}$ to the terminal versus the central carbon atom.\textsuperscript{23} A proposed explanation of this unique selectivity observed in the reaction of $\text{C}^{\ell} + \text{C}_2\text{H}_4$ in solid $p$-$\text{H}_2$ involves the steric effects within the solid $p$-$\text{H}_2$ matrix that cause the reacting $\text{C}^{\ell}$ atom to be guided to the central C atom of $\text{C}_2\text{H}_4$ embedded in a single substitutional matrix site.\textsuperscript{36}

In this work, we extend the study of $\text{C}^{\ell} + \text{C}_2\text{H}_4$ to $\text{C}^{\ell} + \text{trans-1,3-butadiene}$ ($\text{H}_2\text{C}=$CH–CH$=$CH$_2$, hereafter indicated as $\text{C}_4\text{H}_6$), which has two double bonds and a molecular size larger than $\text{C}_2\text{H}_4$. We observed absorption lines ascribable to the trans-1-chloromethylallyl radical, $\text{•}(\text{CH}_2\text{CHCH})\text{CH}_2\text{C}^{\ell}$, indicating that $\text{C}^{\ell}$ was added to the terminal carbon of $\text{C}_2\text{H}_4$. When the UV-irradiated matrix was further irradiated with IR light, we observed lines in another set ascribable to the trans-1-methylallyl radical, $\text{•}(\text{CH}_2\text{CHCH})\text{CH}_3$, indicating the addition of H atom to the terminal carbon of $\text{C}_4\text{H}_6$.

II. EXPERIMENTS

The experimental setup has been described previously.\textsuperscript{15,36,38} In these experiments, a flow of a mixture of trans-1,3-butadiene seeded in $p$-$\text{H}_2$ at a rate of 0.010 mol h$^{-1}$ was co-deposited for 3–4 h with a small flow of pure $\text{C}_2\text{H}_2$ onto a gold-plated copper flat, maintained at 3.3 K with a closed-cycle helium refrigerator (Janis, SHI-415). Typically, the mixing ratio of $\text{C}_4\text{H}_6$/trans/2-$\text{C}_4\text{H}_6$ thus deposited was estimated to be approximately 1.0/5/1500.

Infrared absorption spectra were recorded with a Fourier-transform infrared (FTIR) spectrometer (Bomem, DA8) equipped with a KBr beam splitter and a HgCdTe detector cooled to 77 K to cover the spectral range of 500–5000 cm$^{-1}$. The gold-plated copper flat, on which the mixture of $\text{C}_2\text{H}_4$/trans/2-$\text{C}_4\text{H}_6$ was deposited, served also as a mirror to reflect the incident IR light to the detector. Typically, 600 scans were coadded at each stage of experiment with resolution 0.25 cm$^{-1}$.

After deposition, the matrix was irradiated with light at 365 ± 10 nm from a light-emitting diode (Honle UV Technology, 375 mW) for several hours to produce $\text{C}^{\ell}$ atoms for reactions with $\text{C}_2\text{H}_4$. In some experiments, after deposition, the matrix was annealed at 4.3 K for 1 h to enhance the formation of the molecular complex $\text{C}_2\text{H}_4$/trans/2-$\text{C}_4\text{H}_6$. Because excitation of a $p$-$\text{H}_2$ matrix containing $\text{C}^{\ell}$ atoms with IR light above 4000 cm$^{-1}$ induces the reaction $\text{C}^{\ell} + \text{H}_2 \rightarrow \text{H} + \text{H}\text{C}^{\ell}$,\textsuperscript{39,40} we placed a 2.4-$\mu$m cutoff filter (Andover) before the IR-beam entrance window of the matrix system during spectral data acquisition so as to avoid the reaction of $\text{C}^{\ell}$ atoms with $p$-$\text{H}_2$. In experiments on H + $\text{C}_2\text{H}_4$, to promote the formation of H atoms from the reaction of $\text{C}^{\ell}$ atoms with $p$-$\text{H}_2$ matrix, we removed the 2.4-$\mu$m filter and utilized the IR source of the FTIR spectrometer to excite either the UV-irradiated matrix or the matrix concurrently with the 365-nm light. To distinguish various groups of IR lines observed upon photolysis, secondary photolysis was performed with a low-pressure Hg lamp (Pen-Ray lamp, UVP) with a bandpass filter (254 ± 10 nm, ESCO Products).

$\text{C}_2\text{H}_6$ (99.5%, Scientific Gas Products) and $\text{C}_2\text{H}_2$ (99.9%, Air Products) were used without further purification. The $p$-$\text{H}_2$ was prepared by catalytic conversion at low temperature. Normal $\text{H}_2$ gas (99.9999%, Scott Specialty Gases) passed through a trap at 77 K and a copper coil filled with hydrated iron (III) oxide (30–50 mesh, Sigma-Aldrich) that was cooled with a helium refrigerator (Advanced Research Systems, DE204AF). At the temperature of the catalyst, typically 11–13 K, the concentration of o-$\text{H}_2$ is estimated to be less than 100 ppm.

III. QUANTUM-CHEMICAL CALCULATIONS

Density-functional theory (DFT) calculations with the B3PW91 hybrid functional\textsuperscript{41–43} and the 6-311++G(2d, 2p) basis set\textsuperscript{44} were performed to predict equilibrium geometries and vibrational wavenumbers of possible products of the reactions $\text{C}^{\ell} + \text{C}_2\text{H}_4$ and H + $\text{C}_2\text{H}_4$. The PW91 functional has shown advantages in the prediction of binding energies and geometries of weakly bound complexes,\textsuperscript{35–47} and B3PW91 is one of the family of Becke three parameter hybrid functionals with non-local correlation provided by the PW91 functional.\textsuperscript{41} Analytic first and second derivatives were employed for geometry optimization and vibrational wavenumbers at each stationary point. The anharmonic effects were calculated with a second-order perturbation approach using effective finite-difference evaluation of the third and semi-diagonal fourth derivatives. The GAUSSIAN 09 program was used for the calculations.\textsuperscript{48} Unless specified, the calculations were performed for the isotope $^{35}\text{C}^{\ell}$.

A. $\text{C}_4\text{H}_6$ and the $\text{C}^{\ell}$_2-$\text{C}_4\text{H}_6$ complex

The geometries of trans-1,3-butadiene $\text{C}_4\text{H}_6$ and the $\text{C}^{\ell}$_2-$\text{C}_4\text{H}_6$ complex optimized with the B3PW91/6-311++G(2d, 2p) calculations are shown in Fig. S1 of the supplementary material;\textsuperscript{49} the atomic numbering is indicated in the figure and the $\text{C}^{\ell}$_2 is attaching to the C4 and C3 atoms. $\text{C}_4\text{H}_6$ has a planar C=C–C=C geometry; the calculated C=C and C–C bond lengths (1.334 and 1.451 Å) agree with the experimental values (1.338 and 1.455 Å).\textsuperscript{30} In $\text{C}^{\ell}$_2-$\text{C}_4\text{H}_6$, $\text{C}^{\ell}$_2 is attached to one C=C bond of $\text{C}_4\text{H}_6$ nearly perpendicular to the molecular plane of $\text{C}_4\text{H}_6$. The distance between $\text{C}^{\ell}$ and the terminal carbon (C=C–C, 2.741 Å) is smaller than that between $\text{C}^{\ell}$ and the inner carbon (C=C–C, 2.970 Å); the angle $\angle$C4C/C/C is 168.4°.

Harmonic and anharmonic vibrational wavenumbers of $\text{C}_4\text{H}_6$ were calculated with the B3PW91/6-311++G(2d, 2p) method. Vibrational modes $\nu_{11}$ (CH$_2$ wag, $a_u$, 925 cm$^{-1}$), $\nu_{10}$ (out-of-plane CH bend, $a_u$, 1030 cm$^{-1}$), and $\nu_{20}$ (C=C stretch, $b_u$, 1616 cm$^{-1}$) were predicted to have IR intensities greater than 20 km mol$^{-1}$; the listed values are anharmonic vibrational wavenumbers.

Harmonic and anharmonic vibrational wavenumbers and IR intensities of $\text{C}^{\ell}$_2-$\text{C}_4\text{H}_6$ predicted with the
B3PW91/6-311+g(2d, 2p) method are listed in Table I; for comparison, the mode numbers and symmetry corresponding to C2H6 are also listed. The predicted shifts of anharmonic vibrational wavenumbers from C4H6 are also listed. Wavenumbers of lines of the C2ℓ−3-butadiene (C3H5) complex derived from experiments and calculations using the B3PW91/6-311+g(2d, 2p) method.

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<th>Description</th>
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<th>Anharmonic (cm⁻¹)</th>
<th>Shift (cm⁻¹)</th>
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TABLE I. Comparison of vibrational wavenumbers and IR intensities of the C2ℓ−3-butadiene (C3H5) complex derived from experiments and calculations using the B3PW91/6-311+g(2d, 2p) method.

B. Reaction products of Cℓ + C₄H₆

The optimized geometries of three possible products of the reaction Cℓ + C₄H₆ are presented in Fig. 1; the atomic numbering of each species is indicated. The trans-1-chloromethylallyl radical, (CH₃C=CHCH=CH₂Cℓ, has a planar CCCCH structure with the unpaired electron delocalized over C1, C2, and C3. Two conformers are stable: conformer (1a) with the Cℓ atom located nearly perpendicular to the allyl molecular plane with a dihedral angle of φH4CCCℓ = 70.5° is most stable, and conformer (1b) with the Cℓ atom in the allyl molecular plane has energy ~8 kJ mol⁻¹ above (1a). The lengths of bonds C1−C2 and C2−C3 in (1a), 1.375 and 1.385 Å, respectively, are slightly greater than the predicted C=C bond length of 1.334 Å in C2H6.

In contrast, the unpaired electron of the 2-chloro-3-buten-1-yl radical, (CH₂C=CHCH=CH₂, is localized on the terminal carbon C4. Two conformers are stable: the more stable conformer (2a) has the chlorine atom located off the C1C2C3 plane with an angle of ζC2C3Cℓ = 107.3° and the C1=C2 and C2=C3 bond lengths of 1.327 and 1.485 Å, respectively. Its energy is ~65 kJ mol⁻¹ above trans-1-chloromethylallyl (1a), rationalized as due to the lack of delocalization of the unpaired electron. The conformer (2b) with the chlorine atom in the C1C2C3 molecular plane has energy ~7 kJ mol⁻¹ greater than that of (2a).
The radical has one stable conformer (3), with the in-plane H atom of the methyl moiety trans to the C\text{\textepsilon} atom. It is more stable than trans-1-chloromethylallyl (1a) by 18 kJ mol\(^{-1}\). This radical has the unpaired electron delocalized over C1C2C3 with the C1–C2 and C2–C3 bond lengths of 1.373 and 1.386 Å, respectively.

The predicted harmonic and anharmonic vibrational wavenumbers and IR intensities for conformers (1a) and (1b) of the trans-1-chloromethylallyl radical are listed in Table II; the mode order and approximate mode descriptions follow those of (1a). Modes \(\nu_{21}\) (C\text{\textepsilon}C\text{\textepsilon} stretch, 610 cm\(^{-1}\)), \(\nu_{19}\) (CH\text{\textepsilon} wag, 824 cm\(^{-1}\)), \(\nu_{16}\) (out-of-plane CH bend, 966 cm\(^{-1}\)), and \(\nu_{12}\) (CH\text{\textepsilon}H\text{\textepsilon} wag, 1228 cm\(^{-1}\)) for (1a) have IR intensities greater than 20 km mol\(^{-1}\) and those of \(\nu_{21}\) (697 cm\(^{-1}\)), \(\nu_{19}\) (982 cm\(^{-1}\)), \(\nu_{12}\) (1266 cm\(^{-1}\)), \(\nu_{11}\) (sym. C1C2C3 stretch, 1252 cm\(^{-1}\)), and \(\nu_{6}\) (C4H\text{\textepsilon} sym. stretch, 2947 cm\(^{-1}\)) for (1b) have IR intensities greater than 15 km mol\(^{-1}\); the listed values are anharmonic vibrational wavenumbers predicted with the B3PW91/6-311++g(2d,2p) method.

A C4–C\text{\textepsilon} bond length of 1.867 Å for trans-1-chloromethylallyl radical (1a) is predicted with the B3LYP/6-311++g(2d,2p) method, 0.028 Å greater than that predicted with the B3PW91/6-311++g(2d,2p) method, resulting to discrepancies in predicted harmonic/anharmonic vibrational wavenumbers of the C\text{\textepsilon}C\text{\textepsilon} stretching (\(\nu_{21}\)) mode of 592/569 cm\(^{-1}\) for B3LYP and 636/610 cm\(^{-1}\) for B3PW91. As will be discussed in Sec. VB, observed experimental value for this mode is 650.3 cm\(^{-1}\), supporting the adequacy of using the B3PW91 method.

The predicted harmonic and anharmonic vibrational wavenumbers and IR intensities of 2-chloro-3-buten-1-yl ((2a) and (2b)) and 1-chloro-1-methylallyl (3) radicals are listed in Table SI of the supplementary material. Lines of 2-chloro-3-buten-1-yl (2a) predicted for \(\nu_{22}\) (404 cm\(^{-1}\)), \(\nu_{20}\) (653 cm\(^{-1}\)), \(\nu_{19}\) (708 cm\(^{-1}\)), and \(\nu_{17}\) (947 cm\(^{-1}\)) have IR intensities greater than 30 km mol\(^{-1}\) and those of 2-chloro-3-buten-1-yl (2b) predicted for \(\nu_{22}\) (452 cm\(^{-1}\)), \(\nu_{20}\) (645 cm\(^{-1}\)), \(\nu_{19}\) (688 cm\(^{-1}\)), \(\nu_{18}\) (830 cm\(^{-1}\)), and \(\nu_{17}\) (948 cm\(^{-1}\)) have IR intensities greater than 15 km mol\(^{-1}\); the vibrational modes for these molecules are ordered by wavenumber. Those of trans-1-chloro-1-methylallyl (3) predicted for \(\nu_{20}\) (618 cm\(^{-1}\)), \(\nu_{19}\) (767 cm\(^{-1}\)), \(\nu_{14}\) (1134 cm\(^{-1}\)), \(\nu_{12}\) (1281 cm\(^{-1}\)), \(\nu_{10}\) (1424 cm\(^{-1}\)), and \(\nu_{6}\) (2900 cm\(^{-1}\)) have IR intensities greater than 19 km mol\(^{-1}\). All listed values are anharmonic vibrational wavenumbers.

C. Reaction products of H + C\text{\textepsilon}H\text{\textepsilon}

According to the calculations of Miller using the G3/B3LYP/6-31G(d) method, five structural isomers of the straight chain C\text{\textepsilon}H\text{\textepsilon} radical are stable. Among these, each of the four isomers 1-methylallyl, 3-buten-1-yl, 2-buten-2-yl, and 1-buten-2-yl has two conformers, cis and trans, according to the structure of the CCCC skeleton, whereas the 1-buten-1-yl radical has four conformers (cc, ct, tc, and tt) because the terminal CCCH chain also has either a cis or trans configuration. The most stable structural isomer is the 1-methylallyl radical, (\textbullet(CH_{2}CHCHCH_{3}), rationalized as due to the delocalization of the unpaired electron, and the other four structural isomers have energy at least 67 kJ mol\(^{-1}\) greater than that of trans-1-methylallyl, which is more stable than cis-1-methylallyl radical. The isomerization between various structural isomers of C\text{\textepsilon}H\text{\textepsilon} is unlikely to occur because of large barriers.

The geometries of two most likely reaction products of H + trans-1,3-butadiene, 1-methylallyl and 3-buten-1-yl radicals, predicted with the B3PW91/6-311++g(2d,2p) method are illustrated in Fig. 2; the atomic numbering is indicated. The 1-methylallyl radical, (\textbullet(CH_{2}CHCHCH_{3}), has a planar CCCC skeletal geometry with the unpaired electron delocalized over C1,C2, and C3. Of the two isomers, trans-1-methylallyl (4a) is more stable than cis-1-methylallyl (4b) by ~4 kJ mol\(^{-1}\). For isomer (4a), the C1–C2 and C2–C3 bond lengths are 1.379 and 1.382 Å, respectively, similar to those of the allyl radical (1.387 Å). The H6 atom of the methyl group is located in the molecular plane to have C\text{\textepsilon} symmetry. In contrast, 3-buten-1-yl (\textbullet(CH_{2}CH_{2}CH_{2}CH_{3}) has no conjugated structure and the CCCC skeleton is non-planar. For trans-3-buten-1-yl (5a), the CCCC dihedral angle is 119.6° and the C4H\text{\textepsilon} group, on which the unpaired electron is located, is rotated about 27° from the C2C3C4 plane. The
lengths of the C1–C2 and C2–C3 bonds are 1.327 and 1.501 Å, respectively. For cis-3-buten-1-yl (5b), the four carbon atoms are nearly coplanar, with a dihedral angle of 9.3°.

The predicted harmonic and anharmonic vibrational wavenumbers and IR intensities of trans-1-methylallyl (4a) and cis-1-methylallyl (4b) radicals are listed in Table III; the mode order and approximate descriptions follow those of (4a). Lines of trans-1-methylallyl (4a) for ν23 (C1H2 wag, 786 cm⁻¹), ν22 (out-of-plane CH bend, 954 cm⁻¹), ν19 (C4H2 antisym. stretch, 2902 cm⁻¹), ν19 (C4H2 scissor, 1427 cm⁻¹), ν6 (C3H sym. stretch, 2877 cm⁻¹), ν3 (C3H stretch, 3016 cm⁻¹), and ν2 (C1H2 sym. stretch, 3033 cm⁻¹) are predicted to have IR intensities greater than 20 km mol⁻¹. Those of cis-1-methylallyl (4b) for ν24 (out-of-plane C3H bend, 677 cm⁻¹), ν23 (755 cm⁻¹), ν19 (2910 cm⁻¹), ν9 (1415 cm⁻¹), ν6 (2885 cm⁻¹), ν3 (3054 cm⁻¹), and ν2 (3040 cm⁻¹) are predicted to have IR intensities greater than 20 km mol⁻¹. All listed values are anharmonic vibrational wavenumbers predicted with the B3PW91/6-311+++g(2d, 2p) method.

The predicted harmonic and anharmonic vibrational wavenumbers and IR intensities of 3-buten-1-yl radical are listed in Table II of the supplementary material.49 Lines of trans-3-buten-1-yl (5a) for ν23 (526 cm⁻¹), ν19 (932 cm⁻¹), ν8 (1662 cm⁻¹), ν7 (2831 cm⁻¹), ν5 (3032 cm⁻¹), and those of cis-3-buten-1-yl (5b) for ν24 (524 cm⁻¹), ν23 (528 cm⁻¹), ν19 (925 cm⁻¹), ν8 (1657 cm⁻¹), ν7 (2792 cm⁻¹), ν5 (2880 cm⁻¹), and ν2 (2989 cm⁻¹) are predicted to have IR intensities greater than 15 km mol⁻¹; the listed values are anharmonic vibrational wavenumbers predicted with the B3PW91/6-311+++g(2d, 2p) method.

### IV. EXPERIMENTAL RESULTS

#### A. Complex of C₄H₆

A partial IR absorption spectrum of C₄H₆ in solid p-H₂ (1/1400) at 3.3 K is shown in Fig. 3(a). Intense lines

---

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---

*Approximate mode description and mode order follow those of (1a): v: stretch, δ: bend or deformation, δC: scissor, aC: wag, ρ: rock, τ: torsion, t: twist, a: antisymmetric, σ: symmetric, oop: out-of-plane, ip: in-phase, op: out-of-phase. Numbering of C atoms is according to Fig. 1.

*Numbers in parentheses are IR intensities normalized to the most intense line. The intensities are 52.4 (ν19) and 69.7 (ν23) km mol⁻¹ for conformers (1b) and (1a), respectively.

*Numbers in parentheses are integrated intensities normalized to the most intense one.

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observed at 528.6, 907.9, 1015.2, 1378.9, and 1595.1 cm\(^{-1}\), indicated with arrows in Fig. 3(a). agree with the literature values 524.5 (\(v_{12}\)), 908.1 (\(v_{11}\)), 1013.8 (\(v_{10}\)), 1380.8 (\(v_{11}\)), and 1596.4 cm\(^{-1}\) (\(v_{10}\)) of gaseous trans-C\(_2\)H\(_6\) (Ref. 53–55) and values 536, 925, 1030, 1383, and 1616 cm\(^{-1}\), indicated with “o” in Fig. 3(a), are due to combination or overtone transitions. The weak lines, observed at 916.2 and 994.5/995.4 cm\(^{-1}\) and indicated with cis in Fig. 3(a), are due to cis-1,3-butadiene (\(\sim 1.6\%\)); corresponding lines were observed at 914.6 and 996 cm\(^{-1}\) in a Ar matrix.\(^{57}\) The difference spectrum of a \(\text{C}_2\text{H}_4/\text{C}_2\text{H}_6/p-H_2\) (1/0.1/1.5/1500) matrix sample with lines of C\(_2\)H\(_6\) stripped by subtraction of the scaled spectrum in Fig. 3(a) is shown in Fig. 3(b); some intense lines of C\(_2\)H\(_6\) might have been stripped incompletely. The original intense lines of C\(_2\)H\(_6\) listed above shifted to 534.2, 914.5, 1016.2, 1378.9, and 1591.8 cm\(^{-1}\), and new features at 517.3/519.0, 524.4, 755.3, 918.9, 1281.8, and 1640.4 cm\(^{-1}\) appeared. These lines are indicated as asterisks * in Fig. 3(b). These new features are assigned to absorption of the \(\text{C}_2\text{H}_4/\text{C}_2\text{H}_6\) complex, to be discussed in Sec. V A. The observed wavenumbers and relative intensities are presented in Table I. The ratio of the \(\text{C}_2\text{H}_4/\text{C}_2\text{H}_6\) complex to C\(_2\)H\(_6\) is estimated to be approximately 1:10 according to observed integrated intensities of lines of C\(_2\)H\(_6\) at 907.9 and 1595.1 cm\(^{-1}\) and those of C\(_2\)H\(_4\) at 914.5 and 1591.8 cm\(^{-1}\) and the corresponding IR intensities predicted with theory.

B. Products of the reaction \(\text{Cl} + \text{C}_4\text{H}_6\)

Figure 4(a) shows the absorption spectrum of the \(\text{C}_4\text{H}_6/\text{C}_2\text{H}_2/p-H_2\) (1/1.5/1500) matrix recorded after deposition. Figure 4(b) shows a difference spectrum after irradiation of the matrix at 365 nm for 1.5 h. The difference spectrum was obtained by subtracting the spectrum recorded in the preceding step from that recorded after this step; features pointing upward thus indicate production, whereas those pointing downward indicate destruction. Lines due to the \(\text{C}_2\text{H}_2/\text{C}_4\text{H}_6\) complex decreased in intensity, those of C\(_2\)H\(_6\) increased in intensity, and several new lines appeared; among them the most prominent ones are at 809.0, 962.2, and 1240.6 cm\(^{-1}\). These lines and several weaker lines, showing similar relative intensities in disparate stages of various experiments, are indicated with arrows and wavenumbers in Fig. 4(b); a list is given in Table II. These lines are designated as group A and assigned to the trans-1-chloromethylallyl (1a) radical in Sec. V B. We estimated that about 40% of \(\text{C}_2\text{H}_2/\text{C}_4\text{H}_6\) was reacted and \(\sim 11\%\) of C\(_2\)H\(_6\) was reproduced according to the observed variation in line intensities.

After we continued irradiation at 365 nm for an additional 3 h, the difference spectrum is illustrated in Fig. 4(c). The intensities of lines in group A decreased, whereas those of several weak lines at 738.6/735.3, 933.6, 983.5, 1181.0, 1426.9, and 1447.9 cm\(^{-1}\), indicated as * in Fig. 4(c), increased. These increasing weak features are readily assigned to 3,4-dichloro-1-butene, H\(_2\)C=CHCHC≡CH\(_2\), with assignments made according to the observed spectrum of 3,4-dichloro-1-butene isolated in gaseous p-H\(_2\) recorded in a separate experiment. Weak lines at 967.8 and 1250.4 cm\(^{-1}\), designated as # in Fig. 4(c), are tentatively assigned to 1,4-dichloro-2-butene according to the literature spectrum of gaseous 1,4-dichloro-2-butene.\(^{58}\) These features became much more prominent when we employed a greater mixing ratio of C\(_2\)H\(_2\) to C\(_4\)H\(_6\), consistent with the expectation for dichloro compounds.

The intensities of lines of \(\text{C}_2\text{H}_2/\text{C}_4\text{H}_6\) decreased continuously, those in group A increased in the initial stage, but diminished after further irradiation, whereas the intensities of lines of 3,4-dichloro-1-butene increased continuously. The integrated intensities of lines at 809.0 cm\(^{-1}\) (group A, trans-1-chloromethylallyl), 738.9 cm\(^{-1}\) (3,4-dichloro-2-butene), and 914.5 cm\(^{-1}\) (\(\text{C}_2\text{H}_2/\text{C}_4\text{H}_6\)) are shown in Fig. S2 of the supplementary material.\(^{49}\) In some experiments, the UV irradiated matrix was further irradiated for 1.5 h with light near 254 nm from a low-pressure Hg lamp. The intensities of lines in group A increased significantly, whereas those of 3,4-dichloro-1-butene decreased only slightly.

C. Products of the reaction \(\text{H} + \text{C}_4\text{H}_6\)

Anderson and co-workers reported that Cl atoms might react with \(p-H_2\) to form HCl \(\ell\) when the matrix is subjected to IR irradiation during acquisition of IR absorption spectra.\(^{39,40}\) Utilizing the reaction \(\text{Cl} + H_2 \rightarrow HC\ell + H\) is thus an
excellent method to produce H atoms in solid p-H₂. We have employed this method to investigate reactions of H + C₂H₅/CH₂CH₃Cl (Ref. 59) and H + CH₂ClCH₃.⁶⁰

To promote the reaction of Cl atoms with p-H₂ matrix, we removed the 2.4-µm filter and increased the aperture to 10 mm to utilize more light from the IR source of the FTIR spectrometer to excite the UV-irradiated matrix. Figure 4(d) shows the difference spectrum after irradiation of the UV-irradiated matrix with IR light for 1 h. In addition to the much enhanced line of HCℓ at 2894.1 cm⁻¹ and several weak lines of (HCℓ)₂ complexes in the region 2750–2900 cm⁻¹, several new lines appeared; they are distinct from those observed in group A shown in Fig. 4(b) and those of 3,4-dichloro-1-butene indicated as * in Fig. 4(c). Prominent features were observed at 781.6, 957.9, 1433.6, 3023.5, and 3107.1 cm⁻¹. These lines and several weaker lines showing similar relative intensities in various experiments, designated as group B and indicated with arrows and wavenumbers in Fig. 4(d), are assigned to the trans-1-methallyl radical in Sec. V C. A list of observed lines is given in Table III. Other weak lines observed at 966.5, 1381.2, 1443.7, 1456.7, 2971.8, and 2944.8 cm⁻¹, marked with V symbols in Fig. 4(d), likely correspond to trans-but-2-ene (C₄H₈),⁶¹ even though we could not positively eliminate the possibility of a small contributions from trans-but-1-ene.

V. DISCUSSION

A. Assignments of the Cℓ₂-C₄H₆ complex

New features observed in the Cℓ₂/C₄H₆/p-H₂ matrix as compared with lines of C₄H₆ are indicated with * in Fig. 3(b). The stick spectra of C₄H₈ and Cℓ₂-C₄H₆ according to predicted anharmonic vibrational wavenumbers and IR intensities are shown in Figs. 3(c) and 3(d), respectively. The observed features are listed in Table I and compared with theoretical predictions for Cℓ₂-C₄H₆. The observed and predicted spectral shifts of Cℓ₂-C₄H₆ from features of C₄H₆, also listed in Table I, agree each other satisfactorily. For instance, lines at 534.2 and 914.5 cm⁻¹ of Cℓ₂-C₄H₆ are blueshifted from those

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Raman lines at 1900 (C=C), 1492 (νs), 1265 (νs), 782 (νs), 499 (νs), 289 (νs), and 205 (νs) cm⁻¹ for the gaseous species were reported.⁶²

Numbers in parentheses are IR intensities normalized to the most intense line. The intensities are 38.3 (νs) and 53.4 (νs) km mol⁻¹ for conformers (4b) and (4a), respectively.

Numbers in parentheses are integrated intensities normalized to the most intense one.
FIG. 3. (a) Absorption spectrum of a C$_4$H$_6$/p-H$_2$ (1/1400) matrix deposited at 3.3 K for 5 h. Fundamental lines are indicated with arrows and overtones or combination lines are indicated with o. (b) Absorption spectra of a C$_4$H$_6$/p-H$_2$ (1/1.5/1500) matrix, deposited for 5 h and annealed at 4.3 K for 0.5 h, with lines of C$_4$H$_6$ stripped by subtraction of the scaled spectrum in (a); lines of C$_2$C$_2$H$_6$ are indicated with *. Both spectra were recorded at resolution of 0.25 cm$^{-1}$. (c) IR stick spectrum simulated according to anharmonic vibrational wavenumbers and IR intensities of C$_4$H$_6$ predicted with the B3PW91/6-311++g(2d, 2p) method. (d) simulated stick spectrum of C$_2$C$_2$H$_6$. Some intense lines were decreased and shifted downward for viewing convenience.

FIG. 4. (a) Absorption spectra of a C$_2$/C$_4$H$_6$/p-H$_2$ (1/1.5/1500) matrix, deposited for 5 h and annealed at 4.3 K for 0.5 h. (b) Difference spectrum of the matrix after irradiation with light at 365 nm for 1.5 h. Lines in group A (trans-1-chloromethylallyl) are indicated with arrows and wavenumbers. (c) Difference spectrum of the matrix after further irradiation at 365 nm for additional 3 h. Lines of C$_2$C$_2$H$_6$ are marked with * (3,4-dichloro-1-butene) and # (1,4-dichloro-2-butene). (d) Difference spectrum of the matrix after further irradiation with IR light for 1 h. Lines in group B (trans-1-methylallyl) are marked with arrows and wavenumbers. The intense line at 781.6 cm$^{-1}$ is halved and shifted downward for viewing convenience. Lines marked with ∇ likely correspond to trans-but-2-ene (C$_4$H$_8$). All spectra were recorded at resolution 0.25 cm$^{-1}$.

Lines at 524.4 and 519.0/517.3 cm$^{-1}$ are assigned to the C/Cl stretching mode of $^{35}$Cl$_2$-C$_4$H$_6$ and $^{35}$Cl$^{37}$Cl$-C_4$H$_6$/$^{37}$Cl$^{35}$Cl-C$_4$H$_6$, respectively. Observed isotopic shifts of $-5.4$ and $-7.1$ cm$^{-1}$ are consistent with theoretical predictions of $-5.5$ and $-6.7$ cm$^{-1}$, respectively. An extremely weak line at 510.3 cm$^{-1}$ might be assigned to $^{37}$Cl$_2$-C$_4$H$_6$ with an observed shift of $-14.1$ cm$^{-1}$ near a predicted value of $-12.3$ cm$^{-1}$, but the poor signal-to-noise ratio precludes a definitive assignment. The observed intensity ratios of $\sim$9:3:4 for these lines are also consistent with the natural abundance of $^{35}$Cl : $^{37}$Cl $\approx$ 3:1. This mode was predicted to have an anharmonic wavenumber of 460 cm$^{-1}$, much smaller than our observation. A similar trend was also observed for C$_2$H$_2$: the $^{35}$C/Cl$^{37}$Cl stretching mode was observed at 531.2 cm$^{-1}$, whereas the MP2/aug-cc-pVDZ method predicts...
471 cm$^{-1}$. The discrepancy between observed and predicted vibrational wavenumbers might indicate that the interaction between C$_6$ and C$_2$H$_6$ was overestimated by theory. The vibrational wavenumber of C$_2$ is about 554 cm$^{-1}$.

In the C–H stretching region, lines at 3098.2, 3052.2, 3029.9, 3006.7, 2990.5, and 2980.4 cm$^{-1}$ were observed. The pattern follows that of C$_4$H$_6$, but we hesitate to provide definite mode assignments except for the line at 3098.2 cm$^{-1}$ ($v_{17}$) because of the complication due to the presence of combination and overtone bands.

### B. Assignment of lines in group A to trans-1-chloromethylallyl (1a)

Upon photolysis of the C$_4$H$_6$/C$_6$/p-H$_2$ matrix at 365 nm, we observed new IR lines in group A, indicated with arrows and wavenumbers in Fig. 4(b), and several weak lines of 3,4-dichloro-1-butene, indicated with * in Fig. 4(c). A spectrum recorded at an early stage of photolysis at 365 nm showing negligible contribution of 3,4-dichloro-1-butene is reproduced in Fig. 5(a). We expect that these features are due to the product of the reaction CCl + C$_4$H$_6$. The stick IR spectra of possible products trans-1-chloromethylallyl ((1a) and (1b)) and 2-chloro-3-buten-1-yl (1b) simulated according to anharmonic vibrational wavenumbers and IR intensities predicted with the B3PW91/6-311++g(2d, 2p) method are shown in Figs. 5(b)–5(e) for comparison.

The best agreement between experiment and theory in both wavenumbers and relative intensities is clearly with the spectrum predicted for trans-1-chloromethylallyl (1a) shown in Fig. 5(b). The observed wavenumbers and relative intensities are compared with theoretical predictions for (1a) and (1b) in Table II. Observed intense lines at 650.3, 809.0, 962.2, and 1240.6 cm$^{-1}$ agree satisfactorily with the calculated anharmonic vibrational wavenumbers and relative IR intensities for $v_{13}$ (CC stretch, 610 cm$^{-1}$), $v_{19}$ (C$H_2$ wag, 824 cm$^{-1}$), $v_{16}$ (out-of-plane CH bend, 966 cm$^{-1}$), and $v_{12}$ (C$H_2$ wag, 1228 cm$^{-1}$). A weaker line near 646.7 cm$^{-1}$ might be assigned to the C(CL stretching ($v_{21}$) mode of ($\text{CH}_2\text{CHCH})\text{CH}_2\text{CCl}$; the observed ($^{37}\text{Cl}$ isotopic shift of $-3.6$ cm$^{-1}$ is consistent with values $-3.3$ and $-3.5$ cm$^{-1}$ derived from predicted harmonic and anharmonic vibrational wavenumbers for configuration (1a). In the CH stretching region, weaker lines observed at 2962.3 and 3112.8 cm$^{-1}$ are assigned to the $v_6$ (C$H_2$ sym. stretch, 2973 cm$^{-1}$) and $v_1$ (C$H_2$ antisym. stretch, 3111 cm$^{-1}$) modes; other modes in this region suffer from interference due to absorption of the parent. The deviations between experiments and calculations are less than 15 cm$^{-1}$ (1.8%) except for $v_{21}$ (CC stretch) for which predicted harmonic and anharmonic vibrational wavenumbers are 636 and 610 cm$^{-1}$, respectively, but the observed line is at 646.7 cm$^{-1}$; the observed IR intensity is also smaller than predicted.

Although the other conformer trans-1-chloromethylallyl (1b) has a similar allyl structure, its predicted IR spectrum (Fig. 5(c)) differs distinctively from that of trans-1-chloromethylallyl (1a) and matches much less satisfactorily the observed spectrum in both wavenumbers and relative intensities. Similarly, the spectra of the two conformers of 2-chloro-3-buten-1-yl shown in Figs. 5(d) and 5(e) do not match the observed spectrum. The predicted vibrational wavenumbers and relative IR intensities of 1-chloro-1-methylallyl (not shown) also agree poorly with the observed lines in group A. Furthermore, it is unlikely that this species can be formed from a direct reaction of CCl with C$_4$H$_6$. We exclude also the cis-1-chloromethylallyl from our assignments because the barrier height for trans-cis isomerization is large and the predicted spectral pattern matches poorly with our observation. Hence, we assign the observed features in group A to trans-1-chloromethylallyl.

### C. Assignment of lines in group B to trans-1-methylallyl

As shown in Fig. 4(d), lines in group B appeared upon further irradiation of the UV-irradiated C$_4$H$_6$/C$_6$/p-H$_2$ matrix with IR light from the spectrometer. We reproduce this trace in Fig. 6(a) and indicate lines in group B with arrows to compare with stick spectra, shown in Figs. 6(b)–6(e), of possible products trans-1-methylallyl (4a), cis-1-methylallyl (4b), trans-3-buten-1-yl (5a), and cis-3-buten-1-yl (5b) simulated according to anharmonic vibrational wavenumbers and IR intensities predicted with the B3PW91/6-311++g(2d, 2p) method. Lines marked with symbols $\nabla$ likely correspond to trans-but-2-ene (C$_4$H$_6$)$_2$. The best agreement in both wavenumbers and relative intensities is clearly with the spectrum predicted for trans-1-methylallyl (4a) shown in Fig. 6(b). The observed wavenumbers and relative intensities are compared with theoretical predictions for (4a) and (4b) in Table III.

The intense lines observed at 781.6, 957.9, 1433.6, 2968.8, 3023.5, and 3107.3 cm$^{-1}$ agree with anharmonic vibrational wavenumbers predicted for the $v_{23}$ (C$H_2$ wag, 786 cm$^{-1}$), $v_{22}$ (out-of-plane CH wag, 954 cm$^{-1}$), $v_9$ (C$H_2$ scissor, 1427 cm$^{-1}$), $v_5$ (C$H_4$ stretch, 2964 cm$^{-1}$), $v_3$ (CH$_2$ sym. stretch, 3016 cm$^{-1}$) or $v_2$ (CH$_2$ sym. stretch, 3033 cm$^{-1}$), and $v_1$ (C$H_2$ antisym. stretch, 3105 cm$^{-1}$).
modes of trans-1-methylallyl; some CH stretching lines could not be identified because of interference from the parent absorption. Lines at 1459.1 and 1520.5 cm$^{-1}$, indicated with symbols o in Fig. 6(a), might be due to combination or overtone transitions. The deviations between experiments and calculations are less than 10 cm$^{-1}$ except $v_6$ (25 cm$^{-1}$, 0.9%) and $v_{20}$ ($-19$ cm$^{-1}$, 1.3%). Several lines ($v_7$ at 1499.2, $v_{12}$ at 1257.8 and $v_{16}$ at 879.0 cm$^{-1}$) agree with those (1492, 1265, and 872 cm$^{-1}$) reported in the Raman spectrum of gaseous trans-1-methylallyl.$^{62}$

Although the other conformer cis-1-methylallyl (4b) has a similar allyl structure, its predicted IR spectrum (Fig. 6(c)) differs significantly from that of trans-1-methylallyl (4a), especially in the region 600–1000 cm$^{-1}$, and matches poorly the observed spectrum in both wavenumbers and relative intensities. Similarly, as shown in Figs. 6(d) and 6(e), trans-3-buten-1-yl (5a) and cis-3-buten-1-yl (5b) are predicted to have spectral patterns significantly different from the observed spectrum. We hence conclude that the carrier of the observed new features in group B is trans-1-methylallyl.

**D. Reaction mechanism**

Lines in group A are assigned to trans-1-chloromethylallyl (1a) and no spectral lines of 2-chloro-3-buten-1-yl were identified. We thus conclude that in a solid $p$-$H_2$ matrix, the $Cl$ atom reacts selectively at the terminal carbon atom of trans-1,3-butadiene to generate the trans-1-chloromethylallyl radical rather than at the central carbon atom to form the 2-chloro-3-buten-1-yl radical.

To understand possible reaction paths, we performed DFT calculations using the B3PW91/6-311++g(2d, 2p) method. According to the energy diagram in Fig. 7, only one accessible entrance channel is identified, in which the reacting $Cl$ atom attacks the terminal C atom of C$_4$H$_6$ to form 1-chloromethylallyl with no barrier. We failed to identify a reaction channel for the $Cl$ atom to attack directly the central C atom to form 2-chloro-3-buten-1-yl. The trans-1-chloromethylallyl radical (1a) is more stable by $\sim65$ kJ mol$^{-1}$ than 2-chloro-3-buten-1-yl (2a), and the barrier height for isomerization from (1a) to (2a) is $\sim79$ kJ mol$^{-1}$. Although the energy of this barrier is still smaller than that of the reactant $Cl$ + C$_4$H$_6$, we expect rapid quenching of excess energy in solid $p$-$H_2$ after the addition reaction occurs, so the isomerization (1a) to (2a) unlikely occurs. Although trans-1-chloromethylallyl (3) is more stable than trans-1-chloromethylallyl (1a) by $\sim18$ kJ mol$^{-1}$, we identified no direct reaction path from $Cl$ + C$_4$H$_6$. Isomerization from 2-chloro-3-buten-1-yl (2b) to trans-1-chloro-1-methylallyl (3) has a large barrier ($\sim128$ kJ mol$^{-1}$). Our observation of only trans-1-chloromethylallyl (1a) agrees with the theoretical prediction.

The barrier to convert conformer (1a) of 1-chloromethylallyl to conformer (1b) was calculated to be $\sim9$ kJ mol$^{-1}$. The fact that we observed only trans-1-chloromethylallyl (1a), not conformer (1b), indicates that either the internal rotation of the chloromethyl moiety around the C3–C4 bond is hindered in solid $p$-$H_2$ or the quenching is so rapid that conversion from (1a) to (1b) is infeasible. We previously reported that internal rotation of CH$_3$OH persists in solid $p$-$H_2$, but the large size of the chlorine atom and the geometry of trans-1-chloromethylallyl in solid $p$-$H_2$ might hinder the internal rotation.

The intensities of lines of 3,4-dichloro-1-buten increased continuously with period of irradiation at 365-nm, and also as the mixing ratio of $C_2$ was increased. At the later period of irradiation, the rate of growth of $C_2$C$_3$H$_6$ is nearly equal to the rate of decrease of $C_2$C$_4$H$_6$, as shown in Fig. S2 of the supplementary material, indicating that the secondary reaction of a chlorine atom with C$_2$C$_4$H$_6$ produces C$_2$ClC$_2$H$_4$. The observation that the second chlorine atom attaches to the C1 and C3 positions of trans-1-chloromethylallyl (1a) is consistent with a radical center delocalized among C1, C2, and
C3. That observed lines of 3,4-dichloro-1-butene are more intense than those of 1,4-dichloro-2-butene might indicate that the secondary addition of chlorine to the carbon (C3) adjacent to the chloromethyl moiety of trans-1-chloromethylallyl (1a) is favored if both Cl atoms are from the same ClH.

Lines in group B are assigned to absorption of trans-1-methylallyl (4a); no spectral lines of trans-3-buten-1-yl (5a) are identified. We thus conclude that in a solid p-H2 matrix, the addition of the H atom occurs at the terminal carbon of trans-1,3-butadiene to yield trans-1-methylallyl rather than at the central carbon to yield 3-buten-1-yl. A theoretical calculation reported by Miller predicted the same behavior.34 The reaction channel of H + trans-1,3-butadiene to form trans-1-methylallyl (Fig. 4 of Ref. 51) is nearly barrierless, whereas the transition state for formation of 3-buten-1-yl has energy ~21 kJ mol⁻¹ greater than that of the reactants. The primary product trans-1-methylallyl radical is more stable than trans-3-buten-1-yl by ~67 kJ mol⁻¹, and the isomerization from (4a) to (5a) following migration of a hydrogen atom from the terminal carbon to the central carbon has a large barrier of 205 kJ mol⁻¹. Although the energy of cis-1-methylallyl is predicted to be greater than that of trans-1-methylallyl by only ~4 kJ mol⁻¹, the barrier for trans-cis isomerization in the gaseous phase is predicted to be ~59 kJ mol⁻¹,51 in the environment of a cold p-H2 matrix, this isomerization is unlikely to occur. According to the calculations, formation of other isomers – 2-buten-2-yl, 1-buten-2-yl, and 1-buten-1-yl – cannot occur directly from H + C3H6; isomerization from 1-methylallyl or 3-buten-1-yl also encounters large barriers.51 Our observation of trans-1-methylallyl as the major product is consistent with theoretical predictions.

E. Comparison with the reaction Cℓ + C3H6 in the p-H2 matrix

The present experiments clearly demonstrate that, for the reaction Cℓ + trans-1,3-butadiene in a p-H2 matrix, the reacting Cℓ atom attacks selectively the terminal carbon atom of the C=C double bond to form the trans-1-chloromethylallyl radical, in agreement with theoretical predictions of the path of minimum energy. This result is in contrast to the reaction Cℓ + C3H6 in a p-H2 matrix in which the reacting Cℓ atom attacks the central carbon atom of C3H6 so that 2-chloropropyl radical was observed as the major product.36 In the gaseous phase the formation of 1-chloropropyl has been reported to be much favored over 2-chloropropyl.37

The size restriction and steric effects within the solid p-H2 matrix were proposed to be responsible for this unique selectivity of the reaction Cℓ + C3H6 in p-H2.36 These authors performed DFT calculations in which the geometry of the Cℓ2-C3H6 complex was optimized within the p-H2 lattice at the B3LYP/aug-cc-pVQZ level while holding the p-H2 lattice fixed in space. At 3.2 K, solid p-H2 has a hexagonally close-packed (hcp) crystal structure; within a single hcp layer each p-H2 molecule is separated by 3.76 Å.25 The constraints of the p-H2 lattice produce substantial changes in the geometry of the Cℓ2-C3H6 complex. In the gaseous phase, the distances between the reacting chlorine atom (the one adjacent to the C3H6) and the terminal and central carbon atoms of C3H6 are predicted to be 2.822 Å and 2.925 Å, respectively, whereas in the p-H2 lattice the corresponding distances were predicted to be 3.107 and 2.847 Å, respectively. The calculations thus indicate that in a p-H2 lattice the reacting Cℓ atom will be much nearer the central carbon atom than the terminal carbon atom, as opposed to the gaseous phase in which the Cℓ atom is nearer the terminal carbon. Upon photolysis of Cℓ2, the p-H2 lattice guides the Cℓ atom to the central carbon atom of C3H6, located in a single substitutional site of the p-H2 lattice, through the steric effect.

To examine the Cℓ + C4H6 reaction in solid p-H2, we performed a similar calculation for Cℓ2-C4H6 embedded in the p-H2 lattice with the B3PW91/6-31++G(2d, 2p) method. As C4H6 is larger than C3H6, a double substitutional site is required to accommodate the C4H6 molecule. With an initial geometry of Cℓ2-C4H6 predicted in the gaseous phase, we placed the Cℓ2-C4H6 complex in the p-H2 lattice such that the C=C double bonds of the Cℓ2-C4H6 complex is nearly parallel to the bc plane of the hcp lattice and Cℓ2 is nearly parallel to the a axis. The geometry of the Cℓ2-C4H6 complex was then optimized with the p-H2 lattice fixed in space. The structure of Cℓ2-C4H6 resulted from this optimization is compared with that of gaseous Cℓ2-C4H6 in Fig. 8. The p-H2 molecules are represented with single blue spheres and some p-H2 molecules are removed to reveal the structure of Cℓ2-C4H6. At the same level of calculation, the distances between the reacting chlorine atom and the terminal and middle carbon atoms of C4H6 in gaseous Cℓ2-C4H6 are predicted to be 2.773 Å and 3.007 Å, respectively, whereas in the p-H2 lattice the corresponding distances were predicted to be 2.697 and 3.040 Å, respectively; the reacting Cℓ atom moves toward the terminal carbon atom slightly in solid p-H2. Because C4H6 is located in a double substitutional site, the steric effect of the p-H2 lattice pushes the reacting chlorine atom toward the
terminal carbon slightly as compared with gaseous Cℓ2-C4H6. The reaction Cℓ + trans-1,3-butadiene hence follows the path of minimum energy as predicted in the gaseous phase to form trans-1-chloromethylallyl radicals.

VI. CONCLUSION

The reaction of Cℓ and H atoms with trans-1,3-butadiene in a p-H2 matrix has been studied with IR spectroscopy. When the Cℓ2/C4H6/p-H2 matrix was irradiated with UV light at 365 nm, a group of new lines with intense ones at 646.7, 809.0, 962.2, and 1240.6 cm⁻¹ are assigned to the trans-1-chloromethylallyl (1a) radical according to the expected reaction, the anharmonic vibrational frequencies and IR intensities predicted with the B3PW91/6-311+g(2d, 2p) method. We conclude that the addition of the Cℓ atom to C4H6 in the p-H2 matrix occurs primarily at the terminal carbon atom, consistent with theoretical predictions but in contrast to the reaction of Cℓ + propene in which the addition of Cℓ to the central carbon atom is favored because of the steric effects in the p-H2 matrix. These results demonstrate further that the cage effect of solid p-H2 is diminished so that an isolated Cℓ atom can be produced via photodissociation of Cℓ2 in situ, and subsequently reacts with C4H6 to form the 1-chloromethylallyl radical.

When the UV-irradiated Cℓ2/C4H6/p-H2 matrix was further irradiated with IR light from the spectrometer, a group of new lines with intense ones at 781.6, 957.9, 1433.6, 2851.3, 2968.8, and 3107.3 cm⁻¹ are assigned to the trans-1-methylallyl (4a) radical. The trans-1-methylallyl radical was produced because of the site-selective reaction of H atom with C4H6, as predicted with theory; 31 H atoms were produced from the reaction of Cℓ2 atom with IR-irradiated p-H2 matrix.

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41See supplementary material at http://dx.doi.org/10.1063/1.4745075 for geometries of trans-1,3-butadiene (C4H6) and Cℓ2-C4H6 complex, integrated line intensities of observed species as a function of the UV irradiation period, and vibrational wavenumbers and IR intensities of 2-chloro-3-buten-1-yl, 1-chloro-1-methylallyl, trans-3-buten-1-yl and cis-3-buten-1-yl radicals.