Title: Gas Phase Detection of Oxirene

**Authors:** Jia Wang1,2, Joshua H. Marks1,2, Andrew M. Turner1,2, Alexander M. Mebel3\*, André K. Eckhardt4\*, and Ralf I. Kaiser1,2\*

**Affiliations**

1W. M. Keck Research Laboratory in Astrochemistry, University of Hawaii at Manoa, Honolulu, HI 96822, USA.

2Department of Chemistry, University of Hawaii at Manoa, Honolulu, HI 96822, USA.

3Department of Chemistry and Biochemistry, Florida International University, Miami, Florida 33199, USA.

4Lehrstuhl für Organische Chemie II, Ruhr-Universität Bochum, Bochum 44801, Germany.

\*Corresponding Author: [mebela@fiu.edu](mailto:mebela@fiu.edu) (A.M.M.), [andre.eckhardt@rub.de](mailto:andre.eckhardt@rub.de) (A.K.E), [ralfk@hawaii.edu](mailto:ralfk@hawaii.edu) (R.I.K.)

**Abstract:** Oxirenes - highly strained 4π Hückel antiaromatic organics – have been recognized as key reactive intermediates in the Wolff rearrangement and in interstellar environments. Predicting short lifetimes and tendency to ring-open, oxirenes denote one of the most ob­scured classes of organic transients with the isolation of oxirene (*c*-C2H2O) having remained elusive. Here, we report on the preparation of oxirene in low-temperature methanol - acetaldehyde matrices upon energetic processing through isomerization of ketene (H2CCO) followed by resonance energy transfer of the internal energy of oxirene to the vibration modes (hydroxyl stretching and bending, methyl deformation and rocking) of methanol. Oxirene was detected upon sublimation in the gas phase exploiting soft photoionization coupled with a reflectron time-of-flight mass spectrometry. These findings advance our fundamental understanding of the chemical bonding and stability of cyclic, strained molecules and afford a versatile strategy for the synthesis of highly ring-strained transients in extreme environments.

**Teaser:** Oxirene was prepared and stabilized in matrices through resonance energy transfer prior to identification in the gas phase via photoionization.

**MAIN TEXT**

**Introduction**

Since the very first postulation of oxirenes as highly strained 4π hetero anti-aromatic molecules by Berthelot more than 150 years ago (*1*), oxirenes have emerged as key reactive intermediates in preparative organic synthesis (*2*), physical organic chemistry (*3*), and theoretical chemistry (*4, 5*). Envisaging the short lifetimes (*6, 7*) and high reactivity (*8, 9*) of these transient molecules along with their critical role in the Wolff rearrangement (*4, 10-12*), oxirenes signify one of the most obscured classes of organic reactive intermediates with its prototype - oxirene (oxacyclopropene, **1**, *c*-C2H2O; Fig. 1) - still being elusive. So far, only four substituted oxirenes have been isolated in low-temperature matrices (dimethyloxirene (*8*); bis-(trifluoromethyl)- and perfluoromethylethyl- oxirene (*13*)) or through neutralization-reionization mass spectrometry (methyloxirene) (*9*); these species represent benchmarks for testing fundamental concepts of Hückel (anti) aromaticity (*14-17*).

Whether oxirene (**1**) represents a true local minimum or a transition state has remained an infa­mous paradigm of an unsolved computational “existence problem” (*17, 18*).Ultimately, functio­nals such as the revision of Becke's hybrid functional (B97-2) and the pure functional of Perdew, Burke, and Ernzerhof (PBE0) predicted oxirene (**1**) as a local minimum, but less stable by 341 kJ mol-1 with respect to ketene (**2**, H2CCO) (*16, 17*); this agrees nicely with the earlier study at the single, double, and perturbative triple excitation coupled-cluster (CCSD(T)) level of theory (*15, 19*).However, despite the computational prediction that oxirene (**c**-C2H2O, **1**) can be ‘made’ in the laboratory, all experimental attempts involving, e.g., photoche­mistry of vinylene carbonate (1,3-dioxol-2-one, C3H2O3) in an argon matrix (*20*), gas phase neutralization of the oxirene radical cation (*21*), the identification of oxirene (**1**) as a reactive intermediate in the isomerization of ketene (**2**) to ethynol (**4,** HCCOH) in an argon matrix (*22*), and strained organic precursors such as diazoacetaldehyde (HCN2CHO) (*23, 24*) have failed so far. This is likely due to the previously insurmountable difficulties in transferring the internal energy from the newly formed, vibrationally excited oxirene (**1**) to the surrounding medium (gas phase third body collider, matrix molecules) on time scales competing with its decomposition and/or isomerization to ketene (**2**) or ethynol (**4**).

Here, we report the first laboratory synthesis and detection of oxirene (*c*-C2H2O, **1**) in low-temperature ices of methanol (CH3OH) and acetaldehyde (CH3CHO) exposed to energetic electrons at 5 K. Utilizing the vacuum ultraviolet (VUV) photoionization reflectron time-of-flight mass spectrometry (PI-ReTOF-MS), oxirene (**1**) and its ketene isomer (**2**) are identified through *isomer-specific* photoionization in the gas phase during the temperature-programmed desorption (TPD) phase of the irradiated ices. Combined with isotopic labeling of the reactants, intimate insights into the formation mechanism of oxirene (**1**) via isomerization of ketene (**2**), which is prepared *in situ* via molecular hydrogen loss from acetaldehyde, are obtained with internally excited oxirene (**1**) stabilized under matrix isolation conditions through resonances with the OH stretching and bending, CH3 deformation, CH3 rocking and/or CO stretching modes of methanol. Upon sublimation of the matrix, oxirene molecules (**1**) are photoionized in the gas-phase *isomer-selectively* exploiting single photon ionization (PI) and detected with a reflectron time-of-flight mass spectrometer (ReTOF-MS) (*25*). The preparation and detection of oxirene (**1**) along with its isotopologues reveals its gas-phase stability with lifetimes of at least 8 ± 2 μs despite a barrier of only 22 kJ mol-1 toward isomerization to ethynol (**4**) *(26)*. These findings advance our fundamental knowledge of the exotic chemistry, chemical bonding, and unusual preparation of highly strained molecules such as oxirene (**1**) thus affording an elegant route for the synthesis of hitherto elusive transients through resonance energy transfer of internal energy from the newly formed molecule to the vibration modes of the surrounding matrix (methanol). The strategy of ‘tuning’ the matrix medium to match the resonance conditions with (high energy) vibrational modes of the target molecule to be synthesized will not only assist in preparing ‘elusive’ molecules such as methanetriol (CH(OH)3) organic chemists have attempted to isolate for decades, but also predicts that oxirene (**1**) may also be formed in methanol-rich interstellar ices in cold molecular clouds from acetaldehyde; hence upon sublimation in star-forming regions, oxirene (**1**) represents a worthwhile astronomical target to be searched for in the gas phase by radio telescopes such as the Atacama Large Millimeter/ submil­limeter Array (*ALMA*).

**Results**

The methanol−acetaldehyde ices were monitored during the electron irradiation at 5 K via Fourier transform infrared (FTIR) spectroscopy (figs. S1 and S2). New absorptions emerged at 2129 cm−1, 1840 cm−1, 1306 cm−1, and 1199 cm−1 (fig. S2, tables S1 and S2), which are linked to the CO stretch (2129 cm–1) of ketene (**2**) and/or carbon monoxide (CO), the acetyl radical (CH3ĊO, ν(C=O)), the deformation mode of methane (CH4; ν4), and the hydroxymethyl radical (ĊH2OH; ν4), respectively (*27, 28*). The calculated absorptions of oxirene (**1**) strongly overlap with those of the precursor molecules (table S3) including the broad O–H stretching (3020 – 3600 cm–1; *v*1) and bending (1420 cm–1; ν6) modes, the C–H bending mode (1455 cm–1; *v*5), and the C–O stretching mode (1030 cm–1; *v*8) ofmethanol (*27*) and the stretching mode of the carbonyl moiety (1718 cm–1; *v*4) of acetaldehyde (*28*); consequently, infrared spectroscopy cannot identify the elusive oxirene (**1**), and an alternative detection technique is imperative.

The PI-ReTOF-MS approach represents an elegant technique for identifying gas-phase molecules *isomer-selectively* during the TPD phase via soft photoionization based on their mass-to-charge ratios (*m/z*) and distinct adiabatic ionization energies (IEs) (*25*). Considering the computed IEs of the C2H2O isomers (Fig. 1 and table S4), a photon energy of 9.70 eV was selected first to ionize oxirene (**1**, IE = 8.63 – 8.64 eV) *and* ketene (**2**, IE = 9.58 – 9.59 eV), but neither oxiranylidene (**3,** CH2(CO); IE = 9.82 – 9.83 eV) *(29)* nor ethynol (**4**, IE = 10.00 – 10.01 eV). Thereafter, we reduced the photon energy to 9.20 eV to only ionize oxirene (**1**),but not ketene (**2**) (Fig. 1). At 9.70 eV, the TPD profile of ion counts at *m/z* = 42 from exposed CH3OH−CH3CHO ice (Fig. 2A) reveals three prominent sublimation events centered at 103 K, 112 K, and 128 K. Accounting for the molecular mass of the reactants, signal at *m/z* = 42 could belong to the molecular formulae C3H6 and C2H2O. The formula C3H6 can account for two isomers: cyclo­propane (*c*-C3H6, IE = 9.86 ± 0.04 eV) and propene (CH3CHCH2, IE = 9.73 ± 0.01 eV) (*30*). To unravel the molecular formula(e), experiments with isotopically-labeled reactants were conducted. The sublimation profile at *m/z* = 42 obtained in the CH3OH–CH3CHO ice (Fig. 2A) was shifted by 2 amu to *m/z* = 44 in CH3OH–CD3CHO (Fig. 2B), CD3OH–CD3CDO ices (Fig. 2C), and CH3OH–13CH313CHO ices (Fig. 2D) supporting the existence of two carbon and two hydrogen atoms along with one oxygen atom cementing the molecular formula C2H2O. Since oxiranylidene (**3**) and ethynol (**4**) cannot be ionized at 9.70 eV, the sublimation events in the CH3OH–CH3CHO ices must be linked to oxirene (**1**) and/or ketene (**2**). Upon tuning to 9.20 eV, which is below the ionization energy of ketene (**2**, IE = 9.58 – 9.59 eV), the first two sublimation events centered at 103 K and 112 K vanish; this indicates that both peaks originate from ketene (**2**). Previous studies demonstrated a sublimation temperature of ketene (**2**) of 100 ± 2 K (*26, 31*), which is consistent with the lowest temperature sublimation event of 103 K. The second sublimation event at 112 K can be associated with the phase transition from amorphous to the crystalline phase of methanol covering a temperature range from 100 to 120 K (*32*). The signal centered at 129 K still prevails in the irradiated CH3OH−CH3CHO (*m/z* = 42) *and* isotopic labeling ices (*m/z* = 44; Figs. 2A-2D, bottom) at 9.20 eV – a photon energy at which only oxirene (**1**, IE = 8.63 – 8.64 eV) can be ionized. Therefore, the signal at the sublimation profile peaking at 128 K can solely be linked to oxirene (**1**), which – once again – is supported through the presence of signal at *m/z* = 44 in all isotopically labelled experiments (Fig. 2B-D, bottom). Note that the broad sublimation events from 145 K to 200 K in the CD3OH–CD3CDO (Fig. 2C) and CH3OH–13CH313CHO ices (Fig. 2D) originate from C2H2DO+ (*m/z* = 44) and 13CCH3O+ (*m/z* = 44), respectively. Lowering the photon energy further to 8.25 eV, which is below the ionization energy of all C2H2O isomers, eliminates all ion counts at 129 K thus providing additional evidence that this event results from oxirene (**1**) (Fig. 2E, top). Most important, blank experiments were also conducted at 9.70 eV and 9.20 eV under identical conditions saving the exposure of the ices to ionizing radiation. No ion counts at *m/z* = 42 were detected (Fig. 2A, solid gray lines), demonstrating that the identified species are connected to the irradiation of the ices but not due to ion-molecule reactions in the gas phase.

To support the formation of oxirene (**1**), we conducted an ultraviolet-visible (UV-Vis) photolysis experiment (*33, 34*) of oxirene (**1**). The CH3OH–CH3CHO ice was first irradiated with energetic electrons to produce oxirene (**1**), which was subsequently photolyzed with 304 nm at 20 mW for 5 hours, an absorption which is uniquely linked to oxirene (**1**), but to none of the other isomers (fig. S3). As evident from the TPD recorded after the photolysis, the sublimation peak at 129 K vanishes, and no ion counts at *m/z* = 42 remain (Fig. 2E, bottom). This finding provides compelling evidence on the radiation induced formation and successive photolytic destruction of oxirene (**1**).

Having demonstrated the detection of oxirene (**1**) and ketene (**2**), we are now shifting our attention to their formation mechanisms. After electron irradiation, the new infrared feature at 1840 cm–1 in the ice at 5 K (figs. S1 and S2) is linked to the acetyl radical (CH3ĊO), which represents the dominant product of the unimolecular decomposition of acetaldehyde via carbon-hydrogen single bond rupture (*35*); this process is endoergic by 377 kJ mol–1 (*28*) and can be supplied by the impinging energetic electrons. Note that the vinoxy radical (ĊH2CHO) can also be produced via an endoergic (411 kJ mol–1) atomic hydrogen loss from acetaldehyde (*36*). Ketene (**2**) may form via the removal of one hydrogen atom from the acetyl (CH3ĊO) or vinoxy radical (ĊH2CHO) with reaction endoergicities of 177 kJ mol–1 or 150 kJ mol–1 (*37*), respectively (fig. S4A). Alternatively, ketene (**2**) can be prepared in one step via the elimination of molecular hydrogen through a transition state located 337 kJ mol–1 above the reactant (acetaldehyde) (*38*). Both mechanism via two successive atomic hydrogen losses and/or molecular hydrogen elimination are also confirmed in ices carrying acetaldehyde-*d*3 (CD3CHO; *m/z* = 47), acetaldehyde-*d*4 (CD3CDO; *m/z* = 48), and acetaldehyde-13C2 (13CH313CHO; *m/z* = 46) through the identification of ketene-*d*2 (D2CCO, *m/z* = 44), ketene-*d*2 (D2CCO, *m/z* = 44), and ketene-13C2 (H213C13CO, *m/z* = 44), respectively (figs. S4B-S4D). Therefore, these findings demonstrate that ketene (**2**) originates from acetaldehyde along with its isotopically substituted species. Oxirene (**1**) can then be formed via electron-induced isomerization of ketene (**2**) through a barrier of 353 kJ mol–1. To aid the stabilization and hence transfer of internal energy from oxirene (**1**), the internal energy can be transferred resonantly from the symmetric and asymmetric CH stretching (ν1,ν7), CH wagging (ν3), and the combination modes of oxirene to the OH stretching (ν1) and bending (ν6), CH3 deformation (ν4 and ν5), CH3 rocking (ν7), and/or CO stretching (ν8) modes of methanol (Fig. 3, table S3). Therefore, the presence of a methanol matrix is essential to the preparation and stabilization of oxirene (**1**) due to their overlapping vibrational modes and hence the possibility of resonance energy transfer. The critical role of methanol as an energy transfer medium is also supported through the lack of any detection of oxirene (**1**) in pure acetaldehyde ices under otherwise identical conditions (fig. S5).

The detection of oxirene could have been thwarted by the existence of a singlet formylcarbene intermediate (HCOCH, **6**) in its vicinity. CCSD(T)/6-311G(df,p) calculations by Scott et al. (*11*) predicted singlet formylcarbene (**6**) as a local minimum on the C2H2O potential energy surface. This species was suggested to be 2.0 kJ mol-1 lower in energy than oxirene (**1**) with a barrier separating oxirene (**1**) and formylcarbene (**6**) of only 1.9 kJ mol-1. Successive calculations of Guan et al. (*39*) suggested that singlet formylcarbene (**6**) is an excited electronic state of formylcarbene, with the ground triplet state lying 8.4 kJ mol-1 lower in energy as computed at the CCSD(T)/cc-pVQZ level of theory. If this were the case, under our experimental conditions in the ice matrix, oxirene would have avoided detection by rapidly isomerizing to singlet formylcarbene (**6**), which in turn would have undergone an intersystem crossing (ISC) to triplet formylcarbene (**5**); ISC is expected to be very efficient in the condensed phase. However, no triplet formylcarbene (**5**), which has an adiabatic ionization energy of 6.35 – 6.36 eV, was detected. To rationalize the experimental detection of oxirene (**1**), here we revisited the oxirene (**1**) → ketene (**2**) isomerization pathway at the CCSD(T)/aug-cc-pVTZ level of theory including harmonic zero-point energy (ZPE) obtained by the same method, with anharmonic corrections evaluated at the PBE0/aug-cc-pVTZ level (Fig. 4).

One can see that at this theoretical level, singlet formylcarbene (**6**) still optimizes as a local minimum, but the inclusion of ZPE lifts its energy above the energy of transition state separating it from oxirene (**1**), which is computed to be 3.7 kJ mol-1 more stable than singlet formylcarbene (**6**). Thus, the latter is unlikely to represent a stationary structure on the PES thus making the ISC to triplet formylcarbene (**5**) unlikely. The electronic vertical singlet-triplet gap in oxirene is 270.8 kJ mol–1, which is also reflected in the T1 diagnostic value of 0.0142. However, the T1 diagnostics of singlet formylcarbene (**6**) along with the transition states TS1 and TS2 (Fig. 4, T1 > 0.02) indicate that the CCSD(T) energies of these structures may not be “gold-standard” accurate as multireference effects are significant *(40)*. Much to our surprise the multireference character of formylcarbene has not been considered in previous computational studies. Therefore, we recomputed single-point energies of all coupled cluster stationary structures on the PES using the multireference perturbation theory CASPT2 (*41, 42*) with an active space including 16 electrons and 14 orbitals, which is the full valence active space. As evident from Fig. 4, the singlet formylcarbene (**6**) is positioned 20.8 kJ mol–1 above oxirene (**1**) and 2.7 kJ mol–1 higher in energy than **TS1**. Moreover, the CASPT2(16,14) calculations in the complete basis set (CBS) limit show that triplet formylcarbene (**5**) lies 1.2 kJ mol–1 above oxirene. It is important to note that the inclusion of diffuse basis functions in the basis set is critical to resolve this issue, e.g., at the CASPT2(16,14)/cc-pVTZ level singlet formylcarbene (**6**) is only 0.6 kJ mol–1 higher in energy than oxirene (**1**). Also, the convergence of the CASPT2 energies to the CBS limit is enhanced for the aug-cc-pVxZ basis sets as compared to cc-pVxZ (Table S7). Essentially, the non-existence of singlet formylcarbene (**6**) *(43)* as a stationary structure and a lower stability triplet formylcarbene (**5**) than predicted by the earlier theoretical calculations allow for the experimental detection of oxirene in the present study.

**Discussion**

     To conclude, the present study presents the very first preparation of the hitherto elusive oxirene (*c*-C2H2O, **1**) molecule – the prototype of a highly strained, 4π Hückel (anti)­aromatic heterocyclic organic molecule in low-temperature methanol−acetaldehyde ices and detection in the gas phase exploiting isomer selective soft photoionization reflectron time-of-flight mass spectrometry (PI-ReTOF-MS). Upon formation through isomerization of ketene (H2CCO, **2**), oxirene (**1**) is stabilized in the matrix via resonances with the vibrational modes of the matrix molecule (methanol). Oxirene is photolabile as demonstrated in an UV photolysis experiment. Considering the velocity of 254 m s−1 for the oxirene (**1**) subliming at an average temperature of 129 K along with the distance between the ice surface and the photoionization region of 2.0 ± 0.5 mm, the lifetime of the neutral oxirene (**1**) in the gas phase has to exceed 8 ± 2 μs, while the corresponding molecular ions have to survive at least 29 ± 1 μs before they reach the detector of the ReTOF-MS. The principle of a *resonance stabilization* of energetic transients upon formation with the matrix opens up the door for the synthesis of highly strained organics such as 1H-phosphirene (*c*-C2H2PH) – the isovalent counterpart of oxirene (**1**) – organic chemists have attempted to prepare in the laboratory for decades and predicts that oxirene (**1**) is likely synthesized in methanol-rich interstellar ices in cold molecular clouds from acetaldehyde followed by sublimation in the hot core stage such as of Sagittarius B2 (Sgr(B2)) thus manifesting molecular clouds and star-forming regions as natural laboratories on a macroscopic scale harboring exotic organics in deep space (*44-46*).

**Materials and Methods**

**Experimental Methods**

All experiments were carried out in a stainless steel ultra-high vacuum chamber (UHV) evacuated to a few 10−11 Torr by magnetically levitated turbomolecular pumps (Osaka, TG1300MUCWB and TG420MCAB) backed by an oil-free scroll pump (Edwards, GVSP30). A polished silver substrate was interfaced to a cold head and was cooled down to 5 K by a closed-cycle helium compressor (Sumitomo Heavy Industries, RDK-415E). The silver substrate can be translated vertically and rotated in the horizontal plane. After the wafer reached 5 K, methanol (CH3OH, Sigma-Aldrich, HPLC grade) and acetaldehyde (CH3CHO, Sigma-Aldrich, anhydrous, ≥99.5% purity) were co-deposited onto the wafer by introducing each reactant to the chamber at a pressure of 2 × 10−8 Torr via separate glass capillary arrays. Isotopically labeled methanol (CD3OH, Sigma-Aldrich, ≥98 atom % D) and isotopically labeled acetaldehyde (CD3CHO, CDN isotopes, ≥98 atom % D; CD3CDO, Sigma-Aldrich, ≥99 atom % D; 13CH313CHO, Sigma-Aldrich, 99 atom % 13C) were utilized in duplicate experiments to observe mass shifts of products. Laser interferometry was used to determine the thickness of the ice based on the interference pattern produced by helium: neon laser (CVI Melles Griot; 25-LHP-230; 632.8 nm) reflected off the surface at an angle of 2° relative to the ice surface normal (*47*). By averaging the refractive indexes of pure amorphous methanol ice (n = 1.33 ± 0.04) (*48*) and that of acetaldehyde (n = 1.303) (*49*) the ice thickness was calculated to be 740 ± 40 nm; this is thicker than the penetration depth of the electrons (330 ± 20 nm) to exclude interactions of the electrons with the substrate. The ice composition of methanol to acetaldehyde was determined to be 1.0 ± 0.3:1 by integrating the infrared features of methanol at 2825 cm−1 (ν3) and 3261 cm−1 (ν1) and acetaldehyde at 1122 cm−1 (ν8) and 1345 cm−1 (ν7)which have absorption coefficients of 5.3 × 10−18, 1.01 × 10−16, 6.6 × 10−19 and 1.1 × 10−18 cm molecule−1, respectively (*36, 48, 50, 51*).

After deposition, the samples were irradiated with 5 keV electrons at a current of 20 nA for 15 min or 60 min, which corresponds to doses of up to 2.5 eV molecule−1 for methanol and 3.8 eV molecule−1 for acetaldehyde, respectively, for 20 nA (60 min) according to CASINO simulations (*52*). An additional photolysis experiment at 304 nm for 5 hours was performed after the electron irradiation to photolyze synthesized oxirene. Infrared spectra (6,000 to 500 cm−1) of the ices were collected before, during, and after the irradiation or photolysis to track changes in the chemical composition using a Fourier transform infrared (FTIR; Thermo Nicolet 6700) spectrometer with 4 cm−1 spectral resolution. During the irradiation, FTIR spectra of the ices were recorded every 2 minutes. The FTIR spectra of the pristine ice are shown in figs. S1 and S2. Detailed assignments of the peaks are compiled in tables S1 and S2.

After the irradiation, the ices were heated from 5 K to 320 K at a rate of 0.5 K min−1 (temperature programmed desorption (TPD)) or first photolyzed by 304 nm light at 20 mW for 5 hours and then heated to 320 K at 0.5 K min−1 by a programmable temperature controller (Lakeshore 336). The 304 nm laser light was generated from the frequency doubling of 608 nm light, which was produced by using the second harmonic (532 nm) of a pulsed neodymium-doped yttrium aluminum garnet laser (Nd: YAG, Spectra-Physics, PRO-270, 30 Hz) to pump the Rhodamine 610/640 dye mixture. During the TPD phase, the sublimed molecules from the sample were analyzed by vacuum ultraviolet (VUV) photoionization reflectron time-of-flight mass spectrometry (PI-ReToF-MS). The VUV photons at 9.70 eV, 9.20 eV, and 8.25 eV were generated by resonant four-wave mixing (FWM) of two synchronized pulsed laser beams from two dye lasers (Sirah, Cobra-Stretch) pumped by two Nd: YAG lasers (table S5). To produce 9.70 eV light, the Rhodamine 610/640 dye mixture was pumped by the second harmonic (532 nm) of an Nd: YAG laser to obtain 606.948 nm, which produces ω1 = 202.316 nm via third harmonic generation. The Coumarin 480 dye was pumped by a second Nd: YAG laser to obtain ω2 = 484.982 nm, which then generated 9.70 eV photons in krypton gas by combining with 2ω1. To generate 9.20 eV and 8.25 eV photons, xenon gas was used as a nonlinear medium. The 9.20 eV light was generated by the difference FWM in pulsed gas jets of xenon gas with ω1 = 222.566 nm and ω2 = 638.667 nm, which were obtained via a double frequency of 445.132 nm from Coumarin 450 dye pumped by the third harmonic (355 nm) of an Nd: YAG laser and via 4-(Dicyanomethylene)-2-methyl-6-(4-dimethylaminostyryl)-4H-pyran (DCM) dye in dimethyl sulfoxide (DMSO) pumped by the second harmonic of an Nd: YAG laser, respectively. For 8.25 eV, ω1 = 249.628 nm and ω2 = 736.448 nm are used. The ω1 = 249.628 nm was obtained via a double frequency of 499.256 nm from Coumarin 503 dye pumped by an Nd: YAG laser. While the ω2 = 736.448 nm was generated from LDS (laser dye) 722 dye. The VUV light was spatially separated from other wavelengths (ω1; ω2;2ω1 + ω2; 3ω1; 3ω2) using a biconvex lithium fluoride (LiF) lens (ISP Optics) in an off-axis geometry and then directed at 2.0 ± 0.5 mm above the substrate surface to ionize subliming molecules during the TPD phase. The resulting molecular ions were mass analyzed with a ReTOF mass spectrometer (Jordan TOF Products, Inc.). The signals were amplified by a preamplifier (Ortec 9305), shaped with a 100 MHz discriminator (F-100TD, Advanced Research Instruments Corporation), and recorded by a multichannel scaler (FAST ComTec, MCS6A) with accumulation times of 2 minutes (3600 sweeps) at 30 Hz for each recorded mass spectra in 0.5 K min-1 during the TPD phase.

**Computational Methods**

All density functional theory (DFT) computations were carried out with Gaussian 16, Revision C.01. (*53*) For geometry optimizations and frequency computations, the DFT PBE0 functional (*54-56*) was employed utilizing the Dunning correlation consistent split valence basis set aug-cc‑pVTZ. (*57*) All coupled cluster computations were carried out with the CFOUR program package *(58)*. In general, the coupled cluster level of theory within the frozen core approximation including single, double, and perturbatively included triple excitations [CCSD(T)] *(59-61)* utilizing the Dunning correlation consistent split valence basis set aug-cc-pVTZ *(62)* was used for geometry optimizations and frequency calculations. To disclose the nature of all stationary points we computed the corresponding frequencies (0 imaginary frequencies for minima and exactly 1 imaginary frequency for transition states). Based on these geometries, the corresponding frozen-core coupled cluster (*59-61, 63*) CCSD(T)/aug-cc‑pVTZ and CCSD(T)/aug-cc‑pVQZ single point energies were computed utilizing the built-in extrapolation routine in ORCA 5.0.3 (*64*) and extrapolated to complete basis set limit (*65*) CCSD(T)/CBS(aug-T,Q) with CCSD(T)/aug-cc-pVTZ zero-point vibrational energies (ZPVEs) and PBE0/aug-cc-pVTZ anharmonic zero-point vibrational energy corrections. The adiabatic ionization energies were computed by taking the ZPVE corrected energy difference between the neutral and ionic species that correspond to similar conformations. As in general the difference between heavier isotopologues and standard isotopologues in the zero-point vibrational energy is marginal, we used the ZPVEs of standard isotopologues for IE calculation and assume them as the same for our experiments with heavier isotopologues.The calculated accuracy is within 0.01 eV for IE in comparison with the measured experimental ionization energy of ketene (**2**) (table S4). Computed Cartesian coordinates and vibration frequencies are listed in table S6. The predicted ultraviolet-visible (UV-Vis) spectra for oxirene (**1**) and ketene (**2**) were calculated at the TD-PBE0/cc-pVTZ level of theory (fig. S3). CASPT2 calculations were carried out using the MOLPRO 2021 package (*66*).

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**Figures**

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**Fig. 1**. **Structures of C2H2O isomers**. Bond length (in picometer), angles (in degrees), point groups, and electronic ground states are shown. The atoms are color-coded in white (hydrogen), gray (carbon), and red (oxygen). Ranges of computed adiabatic ionization energies (IEs) corrected for the electric field effect, and relative energies (Δ*E*) calculated at the CCSD(T)/aug-cc-pVTZ level of theory.

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**Fig. 2. PI-ReTOF-MS data recorded during the TPD phase of mixtures ices.** TPD desorption profiles of ion signals of interest detected at 9.70 eV (**top**) and 9.20 eV (**bottom**) in four irradiated (20 nA, 15 minutes) ice mixtures: (**A**) C2H2O+ (*m/z* = 42) in CH3OH–CH3CHO ice; (**B**) C2D2O+ (*m/z* = 44) in CH3OH–CD3CHO ice; (**C**) C2D2O+ (*m/z* = 44) in CD3OH–CD3CDO ice; (**D**) 13C2H2O+ (*m/z* = 44) in CH3OH–13CH313CHO ice. The sublimation events from 145 K to 200 K in CD3OH–CD3CDO ice (**C**, **bottom**) and CH3OH–13CH313CHO ice (**D**, **bottom**) are due to C2H2DO+ (*m/z* = 44) and 13CCH3O+ (*m/z* = 44), respectively. TPD desorption profiles of ion signal for *m/z* = 42 in irradiated (20 nA, 60 minutes) CH3OH–CH3CHO ices recorded at photoionization energy of 9.20 eV and 8.25 eV (**E**, **top**). For the photolysis experiment at 9.20 eV, the ice was photolyzed at 304 nm after the synthesis via electron exposure (**E**, **bottom**). The dashed line indicates the sublimation peak of oxirene. The atoms are color-coded in white (hydrogen), light blue (deuterium), grey (carbon), and red (oxygen).

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**Fig. 3. Peak absorptions (solid vertical lines) and absorption ranges (solid horizontal lines) of methanol (CH3OH, red), acetaldehyde (CH3CHO, black), and oxirene (*c*-C2H2O, blue).** Dotted vertical lines indicate the overlap positions between the absorption peaks of oxirene and reactants (CH3OH, red; CH3CHO, black). Detailed absorption positions and assignments for experimentally measured pure methanol and acetaldehyde icesas well as calculated oxirene are listed in table S3.

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**Fig. 4. Potential energy profile of the reaction coordinate from oxirene (1) to ketene (2).**

**Supplementary Materials**

Figs. S1 to S5

Tables S1 to S7