

Cite this: *Chem. Sci.*, 2020, 11, 5089

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Ultraviolet photochemistry of ethane: implications for the atmospheric chemistry of the gas giants†

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Chemical processing in the stratospheres of the gas giants is driven by incident vacuum ultraviolet (VUV) light. Ethane is an important constituent in the atmospheres of the gas giants in our solar system. The present work describes translational spectroscopy studies of the VUV photochemistry of ethane using tuneable radiation in the wavelength range $112 \leq \lambda \leq 126$ nm from a free electron laser and event-triggered, fast-framing, multi-mass imaging detection methods. Contributions from at least five primary photofragmentation pathways yielding CH₂, CH₃ and/or H atom products are demonstrated and interpreted in terms of unimolecular decay following rapid non-adiabatic coupling to the ground state potential energy surface. These data serve to highlight parallels with methane photochemistry and limitations in contemporary models of the photoinduced stratospheric chemistry of the gas giants. The work identifies additional photochemical reactions that require incorporation into next generation extraterrestrial atmospheric chemistry models which should help rationalise hitherto unexplained aspects of the atmospheric ethane/acetylene ratios revealed by the Cassini–Huygens fly-by of Jupiter.

Received 26th March 2020

Accepted 29th April 2020

DOI: 10.1039/d0sc01746a

rsc.li/chemical-science

Introduction

Understanding, and perhaps one day exploiting, the environment of extraterrestrial bodies is a central objective of planetary science. The gas giants in our solar system (Jupiter, Saturn, Uranus and Neptune) are rich in molecular chemistry and remain targets of intense scientific study. Like Earth, each of

these planets orbits the sun with its own eccentricity and obliquity leading to seasonal variations in incident solar radiation and thus a cycling chemical composition with latitudinal and altitudinal variations in the abundances of the various molecular constituents.¹ Absorption of near-infrared solar radiation by methane (CH₄) makes important contributions to heating the upper atmospheres (stratospheres) of these planets.^{1–3} Methane contributes less to stratospheric cooling, however, which is more reliant on emission from ethane (C₂H₆) and acetylene (C₂H₂).¹ Understanding the balance and interplay between CH₄ and C₂H₆/C₂H₂ is central to understanding the atmospheric dynamics of the gas giants.

Chemical processing in the stratospheres of the gas giants is driven by incident vacuum ultraviolet (VUV) light,⁴ even in the distant, gas-poor giants Uranus and Neptune.⁵ Numerous possible reactions merit consideration, but common photochemical models for these planetary atmospheres necessarily employ a reduced set pruned from a much larger library of reactions, along with their corresponding rates/branching fractions. These models describe many aspects of the atmospheres of Saturn and Jupiter reasonably well^{1–3} but have recognised shortcomings. For example, the dominant C₂H₆ and C₂H₂ generation mechanisms are assumed to involve secondary reactions following photolysis of CH₄.^{6–8} But both the Cassini–Huygens fly-by of Jupiter and terrestrial measurements reveal very different meridional and latitudinal distributions for C₂H₆ and C₂H₂. Such would be surprising if both species are tightly

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† Electronic supplementary information (ESI) available: Details of the experimental and computational methods used in this study; ionisation energies and dissociative ionisation thresholds for various C_xH_y (x = 1, 2, y ≤ 6) species of present interest; mass spectra showing CH₂ (m/z 14) and CH₃ (m/z 15) products from photolysis of C₂H₆ at λ = 125.6, 121.6, 118.2 and 112.0 nm using two different FEL pulse energies; CH₂⁺ and CH₃⁺ ion images analysed to prepare the P(TKER) and β(TKER) distributions shown in Fig. 4 and a representative C₂H₃⁺ ion image acquired following λ = 121.6 nm photolysis of a jet-cooled C₂H₆ sample. See DOI: 10.1039/d0sc01746a



coupled to methane photolysis.^{3,9,10} Neglect of ion-molecule chemistry has been suggested as one possible explanation for this discrepancy,^{3,11,12} but it is also appropriate to question the inputs to the commonly used photochemical schemes. These draw on data⁸ from a range of (often indirect) sources, including predictions, wherein chemical pathways have been included or removed on the basis of how well the model fits the measurements. Ethane is an important participant in these models and, whilst VUV photolysis is accepted as its main destruction route,^{13,14} the dominant fragmentation pathways and photo-products are not well determined.

Early laboratory studies of C₂H₆ photolysis at the resonance wavelengths emitted by a xenon lamp ($\lambda = 147.0$ and 129.5 nm) deduced the involvement of (at least) three fragmentation pathways. Two involve loss of H₂ or two H atoms, the other yields CH₄ + CH₂ products.¹⁵ Subsequent studies using Kr and Ar resonance lamps ($\lambda = 123.6$ and 106.7/104.8 nm, respectively) suggested additional primary fragmentation channels, to CH₃ + CH₃ and, particularly, H + C₂H₅ products.^{16–18} These studies all involved careful end-product analysis but could not distinguish primary photofragmentation processes from secondary reactions following photolysis, nor yield any dynamical information. More recent imaging studies showed formation of H atoms following C₂H₆ photolysis at the Lyman- α wavelength ($\lambda = 121.6$ nm, the most intense VUV wavelength in the solar spectrum), with an isotropic velocity distribution peaking at low kinetic energies and a weak tail extending to higher energies. The form of this distribution was attributed to initial C–H bond fission, yielding a fast H atom and an electronically excited C₂H₅^{*} fragment, followed by a second (slow) H atom from unimolecular decay of the latter.¹⁹

The present study employs two cutting-edge technologies – the intense, pulsed VUV free electron laser (FEL) at the Dalian Coherent Light Source (DCLS)²⁰ and an event-triggered, fast framing, Pixel Imaging Mass Spectrometry (PIMMS2) sensor²¹ – to advance understanding of C₂H₆ photochemistry and to identify similarities and differences with the photochemistry of both lighter (*i.e.* CH₄) and heavier (*e.g.* propane (C₃H₈)) alkanes. The reported data derive from two sets of collision-free experiments: (i) multi-mass velocity-map ion imaging (PIMMS2 detected)²² studies following one-color VUV photolysis of ethane and ‘universal’ (*i.e.* not quantum state selected) photoionisation of CH₂ and CH₃ photoproducts at four (FEL-produced) wavelengths in the range $112.0 \leq \lambda \leq 125.6$ nm, and (ii) VUV photolysis at $\lambda = 121.6$ nm (using photons generated by four wave mixing outputs from a tabletop pulsed laser) and subsequent detection of H atom products using the high resolution H-atom Rydberg tagging technique.^{23,24} The experimental procedures have all been described previously and are thus confined to the ESL.[†]

Results and discussion

(a) Ethane absorption and the energetics of its various dissociation channels

Fig. 1 shows the chosen photolysis wavelengths superimposed on the electronic absorption spectrum of ethane.^{25,26} As with the

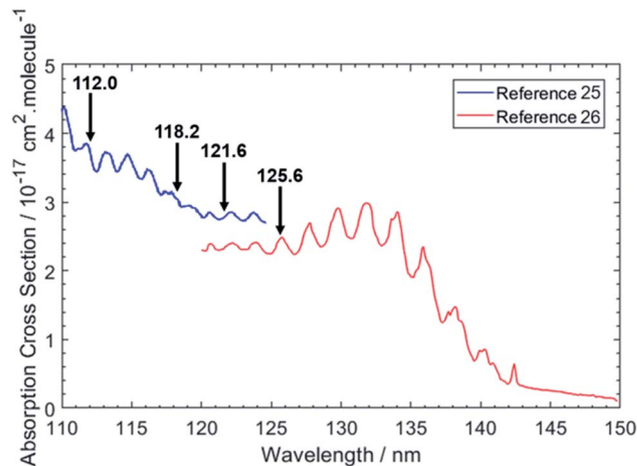


Fig. 1 VUV absorption spectrum of C₂H₆ from ref. 25 and 26. The data from ref. 26 were extracted manually, while the data from ref. 25 were retrieved from ref. 48. The black arrows indicate the photolysis wavelengths (in nm) used in the present study.

other alkanes, the absorption of C₂H₆ lies in the VUV region but, uniquely amongst the alkanes, its room temperature absorption spectrum displays resolved vibronic structure. This structure is attributed to transitions from the near degenerate highest occupied 3a_{1g} and 1e_g valence orbitals to orbitals with dominant 3p Rydberg character. One or more of these are suggested to have significant antibonding valence σ^* character also.²⁷ Excitations to the 3s Rydberg orbital in C₂H₆ are predicted at lower energies, but to be weak – as a result of the molecular center of symmetry – thus distinguishing the 3s Rydberg excitations of ethane from those in CH₄ and the heavier alkanes which all show large absorption cross-sections. This seemingly simple description hides a wealth of potential complexity, however. The degeneracy of the ground (\tilde{X}^2E_g) state of the C₂H₆⁺ cation is lifted by Jahn–Teller distortion, and the structure and dynamics of the resulting cation states are further complicated by the energetic proximity of the low lying \tilde{A}^2A_{1g} excited state – with the result that even a full interpretation of the threshold photoelectron spectrum of C₂H₆ remains elusive.²⁸ Such interactions must also affect the Rydberg states of current interest – since they share the same ion core(s) – and thus affect the absorption spectrum shown in Fig. 1.

Contemporary computational chemistry methods have enabled global investigations of the ground (S₀) state potential energy surfaces (PESs) for species involved in the early stages of the pyrolysis of ethane and other C₁–C₃ hydrocarbons²⁹ but have yet to be directed at the excited state photochemistry of any but the very simplest alkanes. Fig. 2 shows the lower-lying dissociation limits of C₂H₆. The predicted minimum energy conical intersections were located using the global reaction route mapping (GRRM) method and are discussed later. The S₀ state correlates adiabatically with the ground state products from either C–C or C–H bond fission (*i.e.* ground state CH₃ + CH₃ and H + C₂H₅ fragments). The former is the weaker bond, and the formation of ¹CH₂ + CH₄ products is attributed to an (essentially barrierless) H atom transfer between the incipient CH₃



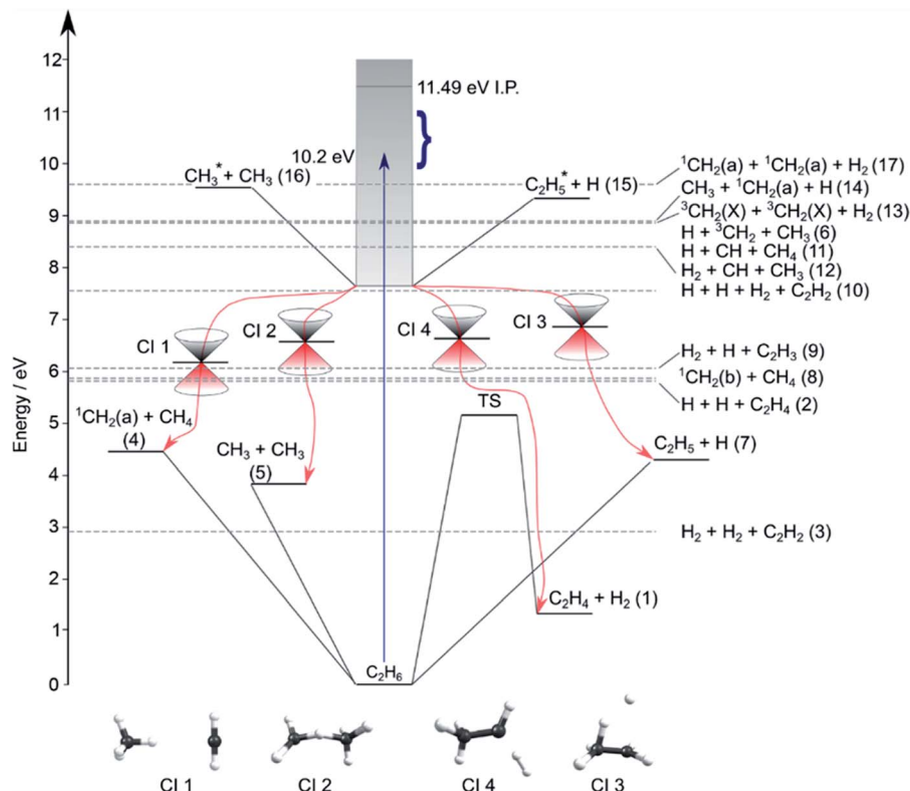


Fig. 2 Energy diagram depicting the excited states of ethane, the thresholds for forming various product combinations (as labelled in Table 1) and illustrative members of families of low-lying CIs (labelled CI1–CI4, identified using the GRRM method) that could facilitate non-adiabatic coupling of excited state population to the S_0 PES and thence to the various dissociation products. Representative structures of these CIs are shown at the foot of the figure. The shaded region indicates the energies spanned by excited electronic states of C_2H_6 , the density of which increases as the ionisation potential (IP) is approached. The vertical arrow shows the energy of a Lyman- α photon and the bracket indicates the range of photolysis photon energies explored in this study.

radicals.²⁹ The energetic thresholds for these three processes are all lower than the calculated barrier to H_2 elimination on the S_0 PES (~ 5.1 eV).²⁹ As Fig. 2 also shows, many more spin-allowed fragmentation channels are energetically accessible following electronic excitation of ethane. Table 1 lists no fewer than 17 chemically intuitive channels that require less than the 10.2 eV of energy provided by a Lyman- α photon. Of these, 8, 7, 6 and 5 of the channels yield, respectively, H atoms, H_2 molecules, CH_2 and CH_3 radicals amongst the dissociation products. Such commonalities provide a major challenge for quantitative studies of the primary photochemistry of ethane (and larger alkanes). Of particular relevance to the present study, the reduced models currently used to describe the atmospheric chemistry of Jupiter and Saturn^{1–3} recognise just reactions (1)–(5) in Table 1.

(b) Ion imaging studies

Fig. 3 shows a representative time-of-flight mass spectrum (TOF-MS) of the ions formed following FEL excitation (at $\lambda = 121.6$ nm) of a jet-cooled sample of C_2H_6 in helium. The spectrum is dominated by a peak associated with H^+ ions. This is unsurprising, given that this wavelength is resonant with the Lyman- α transition from the ground ($n = 1$) state of the H atom.

The remainder of the TOF-MS, displayed on a $5\times$ expanded vertical scale, reveals two clumps of partially-resolved peaks corresponding to CH_x^+ ($x = 2, 3$) and $C_2H_y^+$ ($y = 3–6$) ions. The most intense features in the latter are associated with $C_2H_3^+$ and $C_2H_5^+$ ions. Tables S1 and S2 in the ESI† list relevant adiabatic ionisation and dissociative ionisation thresholds, respectively, and show that four of the neutral products of particular interest (*i.e.* CH_2 , CH_3 , C_2H_3 and C_2H_5) are amenable to single photon ionisation at 10.2 eV, with dissociative ionisation only a (potential) concern if any of these species carry high levels of internal excitation. Note, however, that the observation of some parent $C_2H_6^+$ ion signal highlights the difficulty of completely excluding multiphoton processes even when operating at threshold FEL pulse intensities (<100 nJ).

The inset to Fig. 3 shows that the relative intensity of the CH_2^+ signal increases as the excitation wavelength is decreased. Note that the data shown in the inset were recorded with the detector sensitivity raised for just the relevant narrow range of mass/charge (m/z) ratios, thus allowing averaging over many more acquisitions and improved signal-to-noise ratios. The λ -dependent trend in the CH_2^+ signal is also recognisable in spectra recorded using higher FEL pulse energies but, as shown in Fig. S1,† the relative peak intensities are also pulse energy dependent. Such variations are not unexpected, given the



Table 1 Possible spin-allowed fragmentation channels for C_2H_6 following absorption of a photon with energy $E_{\text{phot}} < 10$ eV. Reactions (1)–(5) were used to describe the photoinduced loss of C_2H_6 in the recent modelling of the stratospheres of Saturn and Jupiter,^{1,3} reactions (6) and (7) are implicated in the present data interpretation and the remaining reactions are numbered in order of increasing reaction enthalpy (calculated from data in ref. 47)

Products	$\Delta_r H$ (0 K)/eV	Dissociation channel
$C_2H_4 + H_2$	1.34	1
$C_2H_4 + H + H$	5.82	2
$C_2H_2 + H_2 + H_2$	3.08	3
$CH_4 + {}^1CH_2(\tilde{a})$	4.46	4
$CH_3 + CH_3$	3.81	5
$CH_3 + {}^3CH_2(\tilde{X}) + H$	8.55	6
$C_2H_5 + H$	4.30	7
$CH_4 + {}^1CH_2(\tilde{b})$	5.89	8
$C_2H_3 + H_2 + H$	6.07	9
$C_2H_2 + H_2 + H + H$	7.56	10
$CH_4 + CH + H$	8.40	11
$CH_3 + CH + H_2$	8.40	12
${}^3CH_2(\tilde{X}) + {}^3CH_2(\tilde{X}) + H_2$	8.81	13
$CH_3 + {}^1CH_2(\tilde{a}) + H$	8.94	14
$C_2H_5^+ + H$	9.33	15
$CH_3^+ + CH_3$	9.54	16
${}^1CH_2(\tilde{a}) + {}^1CH_2(\tilde{a}) + H_2$	9.59	17

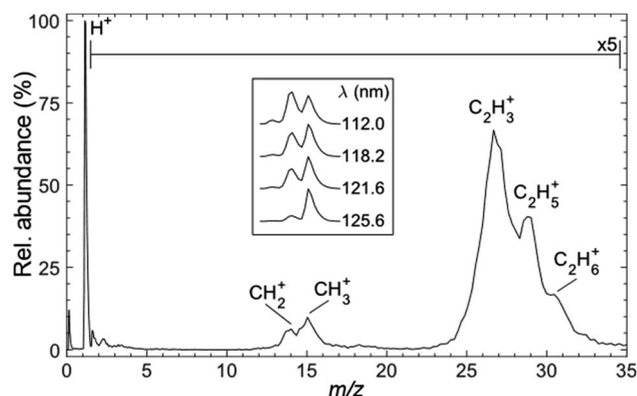


Fig. 3 Time-of-flight mass spectrum of ions formed following photoexcitation of a jet-cooled sample of C_2H_6 seeded in He at $\lambda = 121.6$ nm ($h\nu = 10.19$ eV). Inset: expanded spectra illustrating the λ -dependence of the m/z 14 (CH_2^+) and m/z 15 (CH_3^+) peak intensities.

inevitable differences in the wavelength and internal energy dependent photoionisation cross-sections for CH_3 , 1CH_2 and 3CH_2 radicals.

Use of the PIMMS2 sensor affords not just TOF mass spectra such as those presented in Fig. 3, but also an ion image for each mass channel, in a single acquisition. This provides velocity distributions for each ion peak in Fig. 3. Since C–C bond rupture processes are likely to be pivotal in the cycling of ethane and methane and thus to have a major effect on the atmospheric dynamics, we first present kinetic energy distributions of CH_2 and CH_3 fragments (monitored *via* the corresponding ions) from the photofragmentation of ethane.

Fig. 4 presents the total kinetic energy release $P(\text{TKER})$ distributions (calculated on the basis that the partner to the observed fragment carries all of the remaining mass) and TKER-dependent best-fit recoil anisotropy (β) parameters³⁰ obtained from analysing the ion images retrieved from the central time slice of the TOF-MS peaks corresponding to (A, B) CH_2^+ and (C, D) CH_3^+ ions recorded at FEL wavelengths $\lambda = 125.6$ nm (9.87 eV), 121.6 nm (10.19 eV), 118.2 nm (10.49 eV) and 112.0 nm (11.07 eV). Note that the signal intensities at $\text{TKER} > 35\,000\text{ cm}^{-1}$ are too low for recoil anisotropy parameters to be fitted satisfactorily. Fig. 4A and C also show the corresponding $[P(\text{TKER})]^{1/2}$ plots (dotted lines) to allow better visualisation of the high TKER data. The raw ion images are shown in Fig. S2 of the ESI.†

The distributions derived from the CH_2^+ ion images (Fig. 4A) assume that the co-fragments are CH_4 (*i.e.* that the CH_2 fragments derive from reaction (4) in Table 1). This assumption must be correct for the more translationally excited CH_2 products, which display an anisotropic velocity distribution characterised by a positive recoil anisotropy parameter, $\beta \sim +0.5$ – 0.7 (Fig. 4B), *i.e.* the CH_2 fragments recoil preferentially along the axis parallel to the polarisation vector ϵ of the photolysis laser photons. But the $P(\text{TKER})$ distributions extend to $\text{TKER} \sim 0$ – implying substantial internal excitation of some of the CH_2 and/or CH_4 fragments. Indeed, as Table 1 shows, the chosen photon energies exceed the thresholds for several three-body fragmentation processes that yield CH_2 products. Some or all of channels (6), (13), (14) and (17) in Table 1 could contribute to the increased low-TKER product yield observed at the two shortest excitation wavelengths – a point to which we return later. Thus the precise form of the $P(\text{TKER})$ distribution at low TKER is ill-defined, since the momentum conservation arguments used to convert a measured CH_2 fragment velocity (derived from the image radius) into a TKER value are likely not to apply in a three-body dissociation. But this does not negate the conclusions that (i) the relative yield of slow fragments in the CH_2^+ images increases with increasing photon energy and (ii) the slower fragments, which display minimal recoil anisotropy ($\beta \sim 0$), likely arise *via* one or more of the three-body fragmentation processes.

The distributions derived from the CH_3^+ ion images (Fig. 4C) peak at $\text{TKER} \sim 0$ and show a tail extending to higher TKER that becomes more anisotropic (to positive β) and relatively more intense as the photolysis wavelength is reduced. As can be deduced from Table 1, the maximum possible TKER of CH_3 fragments formed *via* reaction (5) following excitation at $\lambda = 121.6$ nm (Fig. 4C) would be ~ 6.38 eV ($\sim 51\,500\text{ cm}^{-1}$); the high-TKER tails of the $P(\text{TKER})$ distributions shown in Fig. 4C (derived assuming C–C bond fission) extend to values for which the direct C–C bond fission channel (5) is the only possible one photon induced CH_3 fragment formation pathway. Most of the imaged CH_3 fragments appear with much lower TKER, however. Table 1 shows several potential sources of slow CH_3 radicals, including three-body dissociations (6), (12) and (14) and the production of an electronically excited CH_3 partner (channel (16)), the relative likelihoods of which are discussed below.



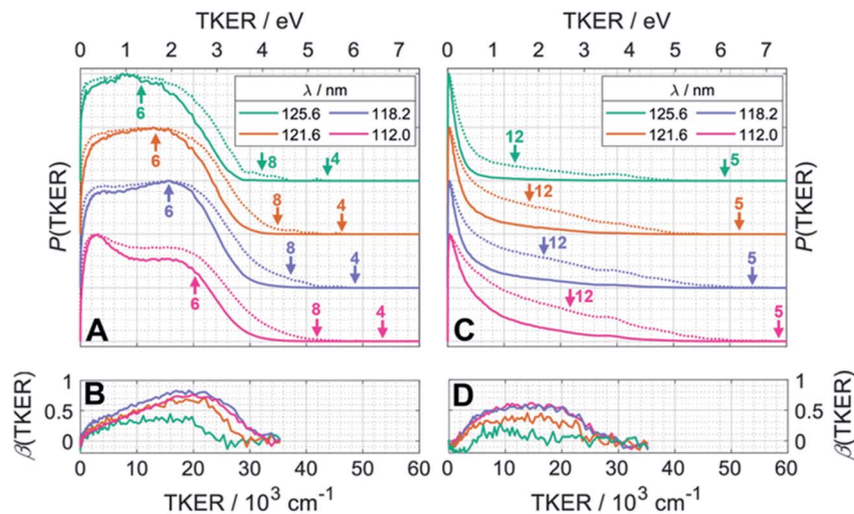


Fig. 4 $P(\text{TKER})$ and $\beta(\text{TKER})$ distributions derived from the analysis of ion images (Fig. S2†) for (A and B) CH_2 and (C,D) CH_3 fragments from photolysis of a jet-cooled C_2H_6 sample, with the TKER shown in both cm^{-1} and eV (bottom and top axes, respectively). The distributions for each photolysis wavelength are offset vertically for display purposes and the dotted lines in plots (A) and (C) show the square root of $P(\text{TKER})$ – normalised to the same maximum value – in order to accentuate the high TKER tails. The TKER_{max} values associated with the two-body fragmentation channels (4), (5) and (8) as well as the most exoergic three body fragmentations yielding CH_2 and/or CH_3 fragments (channels (12) and (6)) are also indicated by vertical arrows in panels (A) and (C). TKER_{max} values for other many-body fragmentations can be derived from Table 1 but, as noted in the text, given the assumed TOF to TKER conversion scheme it is likely that the products from such many-body fragmentations would appear at TKER values well below TKER_{max} .

(c) H atom photofragment time-of-flight (TOF) spectra

H atom TOF spectra were recorded following photolysis of a jet-cooled ethane sample in He at $\lambda = 121.6$ nm with ϵ aligned at, respectively, $\phi = 0, 54.7$ and 90° to the detection axis and converted to the corresponding $P(\text{TKER})$ and $\beta(\text{TKER})$ distributions, shown in Fig. 5A and B, by assuming C_2H_5 as the co-fragment. The fastest products have $\text{TKER} \sim 35\,000\text{ cm}^{-1}$ (~ 4.3 eV). This TKER value is greater than that reported in the earlier imaging study at this wavelength¹⁹ but still well below the maximum allowed by energy conservation assuming single C–H bond fission in ethane (channel (7), for which $\text{TKER}_{\text{max}} \sim 6.5$ eV). In contrast to the case of CH_4 , however, the H atom recoil velocity distribution is essentially isotropic.

(d) Active photofragmentation channels

The present work identifies fragments formed by VUV photolysis of C_2H_6 , assures that these arise *via* collision-free unimolecular dissociation and affords insights into the fragmentation dynamics. The translational spectroscopy data for the CH_2 , CH_3 and H atom products hint at similarities in the fragmentation mechanisms following VUV photoexcitation of C_2H_6 and CH_4 and it is useful to summarise current knowledge of the photofragmentation dynamics of CH_4 to provide context for the discussion that follows.

Only the ground (S_0) state and a repulsive triplet excited state of CH_4 correlate with the lowest energy C–H bond fission limit (associated with $\text{H} + \text{CH}_3$ products). The first excited singlet (S_1) state of CH_4 correlates adiabatically with $\text{H} + \text{CH}_3^*$ products; the electronically excited CH_3^* fragments predissociate rapidly to $\text{H} + {}^1\text{CH}_2(\tilde{a})$ products.³¹ (Here and henceforth, we use superscript * and # symbols to indicate, respectively, electronically and

rovibrationally excited products). Nonetheless, experiments find a substantial quantum yield of ground state C–H bond fission products following VUV photoexcitation of CH_4 , and the

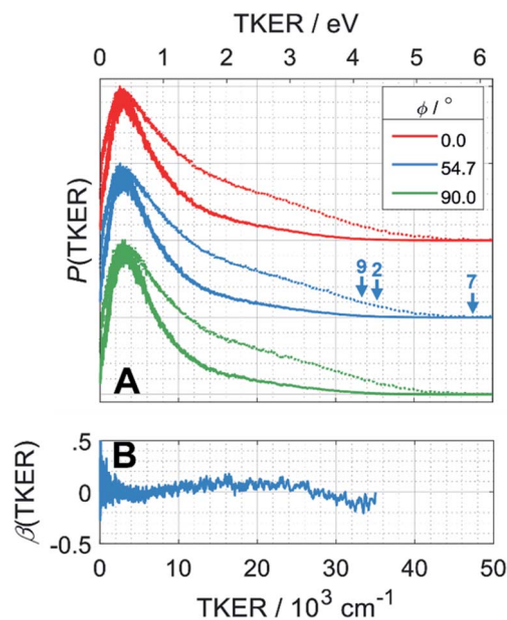


Fig. 5 (A) $P(\text{TKER})$ distribution distributions derived from H atom TOF spectra recorded following photolysis of a jet-cooled C_2H_6 sample at $\lambda = 121.6$ nm with the ϵ vector aligned at $\phi = 0, 54.7$ and 90° to the detection axis and (B) the $\beta(\text{TKER})$ distribution derived therefrom. As in Fig. 4, the individual data sets are offset vertically for display purposes and the high TKER part of the data are accentuated by also plotting $[P(\text{TKER})]^{1/2}$ distributions (dotted lines). The TKER_{max} values associated with primary C–H bond fission (channel (7)) and the two lowest energy three-body dissociation channels (2) and (9) from Table 1 are indicated by vertical arrows.



H atom products display anisotropic recoil velocity distributions – implying that the photoexcited molecules dissociate on a time scale that is much shorter than the rotational period of the parent molecule (which is estimated to be a few picoseconds).^{30,32–34} These findings highlight the importance of non-adiabatic couplings *via* conical intersection (CIs) between the S_1 and S_0 PESs.^{35,36} Theory shows that, to form H + CH₃ products, one C–H bond in the photoexcited CH₄ must start stretching and sweep through the plane defined by the other atoms to access the S_0 PES and dissociate. Angular momentum conservation requires that the resulting CH₃ products are highly rotationally excited; indeed, some of these CH₃(\tilde{X})[#] fragments are formed with so much internal energy that they dissociate further – to H + CH₂ and/or H₂ + CH products. Rival distortions have also been identified, whereby photoexcited CH₄ molecules dissociate by eliminating H₂. Theory suggests that the partner CH₂ fragments in this case are formed in the \tilde{a}^1A_1 state (for dissociations occurring after non-adiabatic coupling to the parent S_0 PES) and the \tilde{b}^1B_1 state (if dissociation occurs adiabatically on the excited state PES).³⁶

Quantitative simulations of the early time nuclear motions following photoexcitation of C₂H₆ remain challenging but global reaction route mapping (GRRM)^{37,38} calculations (summarised in the ESI†) can offer important insights by predicting low-lying conical intersections (CIs) between the PESs for the S_0 and S_1 states. The present VUV photoexcitations will populate one or more S_n ($n > 1$) states of ethane, but we henceforth assume that molecules excited to these higher S_n states undergo efficient non-radiative coupling to the S_1 state. As Fig. 2 showed, the S_1 state of ethane correlates with H + C₂H₅[#] and CH₃ + CH₃[#] products (channels (15) and (16) in Table 1). The C₂H₅[#] and CH₃[#] species are both unstable and dissociate to give, respectively, H + C₂H₄ (ref. 39–41) and H + ¹CH₂(\tilde{a}) (ref. 31) products. As in CH₄, C₂H₆(S_1) molecules can also decompose after non-adiabatic coupling to the S_0 PES. The nuclear distortions required to access the predicted CIs between the S_1 and S_0 PESs (shown in Fig. 2) correlate well with ¹CH₂ elimination once an H atom has inserted between the two C atoms, with C–C or C–H bond fissions and with loss of H₂. The present study is sensitive to the first three fragmentation processes, which are considered in turn. Given the photon energies involved and the multi-dimensional nature of many of these distortions, we can anticipate that (as in the case of CH₄ (ref. 30, 33 and 34)) many of the polyatomic products will be formed with sufficient internal energy that they will fragment further.

CH₂ radical formation. The imaging studies reveal CH₂ fragments, with non-zero β values, implying that these are again formed on a timescale shorter than the parent rotational period. The $P(\text{TKER})$ distributions extend to values where the partner fragment can only be CH₄, but not to sufficiently high TKER values to allow unambiguous determination of the electronic state of the CH₂. Spin-conservation arguments suggest that CH₂ radicals formed in tandem with CH₄ will be in their \tilde{a}^1A_1 state (for dissociations that occur following non-adiabatic coupling at a CI with the S_0 PES) and/or \tilde{b}^1B_1 state (for dissociation on the S_1 PES). But the distributions also extend to TKER ~ 0 , showing that one or other or both fragments are formed with a broad

spread of internal energies. The photoexcitation energies are sufficient to induce three-body fragmentations and, simply on energetic grounds, any of channels (6), (13), (14) and (17) in Table 1 could contribute to signal in the CH₂⁺ ion images. Of these, unimolecular decay of any sufficiently internally excited CH₄[#] partner would be expected to contribute to the yield of (slow) H and CH₃ products, *i.e.* the net reaction (14) in Table 1.

Fig. 4A shows an additional feature at low TKER in the distributions derived from the CH₂⁺ images measured at the two shorter wavelengths. This might signify the opening of a new (three-body) route to ¹CH₂ products, but this feature more likely indicates the presence of ³CH₂ photoproducts: The ground states of the ³CH₂ radical and the CH₂⁺ cation have very similar geometries. Photoionisation of ³CH₂ thus tends to be vibrationally adiabatic (*i.e.* to favour $\Delta v = 0$ transitions)^{42–44} and, from Table S1,† should only be expected at $E_{\text{phot}} > 10.39$ eV (*i.e.* at $\lambda < 119.3$ nm). Note that the feature at low TKER in the $P(\text{TKER})$ distributions shown in Fig. 4A appears to ‘turn on’ and become more prominent as the photon energy is tuned above this threshold. Several possible routes to forming ³CH₂ products can be envisaged, including the unimolecular decay of highly internally excited CH₃[#] fragments (from initial C–C bond fission) or of C₂H₅[#] fragments (following primary C–H bond fission) after non-adiabatic coupling to the S_0 state – as discussed below. Both would contribute to net process (6) in Table 1, though not necessarily exhibit similar energy disposals.

C–C bond fission. The tails of the $P(\text{TKER})$ distributions derived from the CH₃⁺ images extend to TKER values that can only be accommodated by assuming C–C bond fission and formation of two CH₃ radicals (*i.e.* reaction (5) in Table 1). Most of the measured CH₃ fragment velocities imply TKER values far below the maximum allowed by energy conservation, however. Focussing first on the high TKER region in Fig. 4C, the CH₃ fragment yield is clearly rising with decreasing TKER, indicating a preference that one (or both) CH₃ fragments from reaction (5) are formed internally excited. Such energy disposal would likely be a consequence of the nuclear motions that enable non-adiabatic coupling to the S_0 PES. Again, the non-zero β parameter revealed by the CH₃ images imply that these nuclear motions and the ensuing C–C bond fission on the S_0 PES also occur on a timescale shorter than the parent rotational period.

In principle, the entire $P(\text{TKER})$ distribution derived from the CH₃ image could be attributed to channel (5) if the fragmentation dynamics were heavily biased towards forming very highly internally excited CH₃[#] products. The unimolecular decay of these CH₃[#] fragments would be a source of the ³CH₂ products inferred above (reaction (6)), and these ³CH₂ products would be expected to display a similar translational energy distribution to that of the CH₃[#] products (since the light H atom partner would take the bulk of any excess energy released in the secondary fragmentation). Such expectations are consistent with the experimental data and, as noted above, the non-observation of a peak attributable to ³CH₂ products at longer wavelengths (*e.g.* at $\lambda = 121.6$ nm) need not imply that CH₃[#] fragments are not formed but simply that the ³CH₂ products from their decay are not amenable to photoionisation at the longer wavelengths.



The dominance of translationally 'cold' (*i.e.* internally 'hot') CH₃ products in the *P*(TKER) distributions is striking, however. Table 1 suggests several other potential sources of slow CH₃ products. Adiabatic dissociation on the S₁ PES to CH₃ + CH₃^{*} products is an interesting contender. This process is exoergic at all wavelengths studied, though the adiabatic S₁ PES will likely exhibit a barrier at short R_{C-C} bond extensions as the Rydberg function acquires increasing σ* antibonding valence character.⁶ The CH₃^{*} radicals would be unstable with respect to H + ¹CH₂(\bar{a}) products.³¹ Again, the H atoms would carry most of any kinetic energy release, so the translational energy distributions of any ¹CH₂(\bar{a}) fragments formed in this way should broadly mirror that of their CH₃^{*} precursor. ¹CH₂(\bar{a}) fragments are amenable to photoionisation at all wavelengths investigated in the present work, but the TKER distributions derived from the CH₂⁺ images measured at the longer excitation wavelengths show no 'spike' at low TKER – suggesting that any contribution to the ¹CH₂(\bar{a}) yield from adiabatic dissociation to CH₃ + CH₃^{*} products on the S₁ PES must be small compared to that from reaction (4).

C–H bond fission

The *P*(TKER) distribution derived from the H atom TOF measurements (Fig. 5A) extends to TKER values that can only be attributed to prompt C–H bond fission following VUV photoexcitation of C₂H₆, *i.e.* to reaction (7) in Table 1. The C₂H₅ co-fragments are formed with a very broad spread of internal energies. Analogy with CH₄ suggests that this energy disposal is a consequence of the nuclear motions that promote C–H bond fission by non-adiabatic coupling to the S₀ PES.^{6,35} Most of the 'C₂H₅' products assumed in deriving the *P*(TKER) distribution have sufficient internal energy to dissociate further – by loss of another H atom (net reaction (2)), or H₂ (net process (9)), or both (net channel (10)), or to two CH_x species (*e.g.* *via* net channels (6) or (11) in Table 1).⁴² However, the smoothly varying *P*(TKER) distributions shown in Fig. 5A suggest that such overall three- (or more-) body dissociations occur sequentially, *i.e. via* a prompt C–H bond fission and subsequent unimolecular decay of the resulting C₂H₅[#] radicals. The low-TKER peak in Fig. 5A could also indicate an adiabatic contribution to the overall dissociation, yielding H + C₂H₅^{*} as primary products – as suggested in the earlier imaging study at λ = 121.6 nm.¹⁹ Any C₂H₅^{*} fragments would dissociate, yielding H + C₂H₄ products with a spread of translational energies^{39,45} (*i.e.* net reaction (2)). Many of the C₂H₄ products formed by decay of C₂H₅[#] or C₂H₅^{*} species may well be formed with sufficient internal energy to decay yet further, to H + C₂H₃ (vinyl) radical products, or by eliminating H₂ to yield C₂H₂. The former products are observed in the present study, *via* the C₂H₃⁺ peak in the TOF-MS in Fig. 3 and corresponding small ion image shown in Fig. S3 of the ESI,[†] but the current work is blind to C₂H₂ products – which were identified by end-product analysis in the early VUV photolysis studies of C₂H₆ under collisional conditions.^{16–18} For completeness, we note that C₂H₂ products could also arise *via* sequential H₂ eliminations from, first, C₂H₆[#] (formed by non-adiabatic coupling to the S₀ state) and then from the resulting C₂H₄[#] intermediates (*i.e.* net reaction (3) in Table 1). C₂H₂

formation by loss of four H atoms from C₂H₆ is energetically forbidden at the VUV wavelengths of current interest.

The present study does not return quantum yields and, as noted above, is silent regarding some molecular elimination channels. But it certainly identifies several active fragmentation channels and provides new insights into the likely fragmentation dynamics. The present analysis finds no compelling evidence for adiabatic dissociation on an excited state PES – implying efficient non-adiabatic coupling between excited states of C₂H₆ and to the S₀ PES. Many of the present interpretations align with the results of recent quasi-classical trajectory surface hopping calculations for the next larger alkane, propane (C₃H₈), following excitation at λ = 157 nm, wherein it was concluded that most dissociations occur after internal conversion to the S₀ PES, that the energy disposal in the resulting fragments is governed by dynamical rather than statistical factors, and that the three-body fragmentation processes occur sequentially.⁴⁶

(e) Implications for modelling the atmospheres of the gas giants

This work provides detailed new insights into the VUV photochemistry of ethane. The results discussed in detail in the above subsections are summarised below in terms of their implications for modelling the atmospheres of the gas giants. These new results must influence future models:

(i) The HRA-PTS studies reveal kinetic energy distributions extending to values that, on energetic grounds, can only be attributed to prompt C–H bond fission, confirming primary C–H bond fission yielding H + C₂H₅ products (reaction (7) in Table 1). This reaction does not feature in current models used to describe the chemical processing in the stratospheres of the gas giants. Most of the C₂H₅ species are formed with enough internal energy to decay further, almost certainly yielding some H + C₂H₄ products. The present study thus supports inclusion of reaction (2) in the modelling and implies that the two H atoms in that case are lost sequentially.

(ii) The kinetic energy distributions derived from the CH₃⁺ ion images extend to TKER values that can only be attributed to C–C bond fission yielding two CH₃ radicals, confirming that the C–C bond fission channel (reaction (5)) is active and supporting its inclusion in the modelling. The finding that the *P*(TKER) distributions peak at TKER ~0 implies that one of the CH₃ fragments is generally formed with sufficient internal energy to decay further. If C–C bond fission completes after non-adiabatic coupling to the S₀ PES, the resulting CH₃[#] fragments most likely decay to H + ³CH₂(\bar{X}) products (*i.e.* net reaction (6)). This reaction is not included in the current model and, according to the present analysis, will have significantly higher quantum yield than reaction (5).

(iii) The imaging studies confirm formation of CH₂ fragments, with a smooth *P*(TKER) distribution that extends to TKER values such that the partner fragment can only be CH₄. Spin-conservation arguments and the deduced efficiency of non-adiabatic coupling to the S₀ PES suggest that these faster CH₂ fragments are formed in the \bar{a}^1A_1 state. The inclusion of reaction (4) in the photochemical modelling is vindicated.



(iv) The primary fragmentations and resulting product energy disposals following VUV photoexcitation of ethane are shown to be governed by dynamical rather than statistical factors; three-body dissociations are commonplace and occur sequentially. Clearly, quantitative branching ratios for the various active channels are still needed, but the present work offers several clear pointers. Reaction (7) and, particularly, the three-body fragmentation (6) are active and require incorporation in future modelling. The yield of (currently neglected) reaction (6) is deduced to be larger than that of reaction (5). The processes revealed in this study all involve relatively 'prompt' C–H or C–C bond fission, after non-adiabatic coupling to the S_0 PES. As Fig. 2 shows, the respective bond energies are lower than the energy barriers to C_2H_4 formation by H_2 elimination on the S_0 PES. Analogy with CH_4 suggests that any H_2 and C_2H_4 products formed *via* process (1) will be both translationally and vibrationally excited. The likelihood that the $C_2H_4^\#$ species would have sufficient internal energy to surmount the barrier to eliminating a further H_2 (to yield C_2H_2 or H_2CC) is unclear. We further note that the substantial (~60%) branching into C_2H_x species following VUV photoexcitation of C_2H_6 assumed in the current planetary atmospheric photochemistry models derives from indirect measurements made more than half a century ago, and is predicated on an assumption that the decomposition of the internally excited $C_2H_4^\#$ species formed *via* reaction (1) would mimic that deduced following VUV photoexcitation of strategically deuterated ethene (CH_2CD_2) molecules.^{16,17} Such an assumption must be questionable, given the differences in available energy and the recognised importance of dynamics (*i.e.* the topographies of, and non-adiabatic couplings between, the PESs sampled in the two cases) in determining the product branching and energy disposal. It seems likely that the current models overestimate the relative yield of C_2H_x (particularly C_2H_2) photoproducts.

Conclusions

Translational spectroscopy methods employing two cutting-edge technologies – the Dalian Coherent Light Source (DCLS) Free Electron Laser (FEL) and a fast-framing PImMS2 camera – have revealed many new insights into the rich (and hitherto largely impenetrable) VUV photochemistry of ethane. The present findings should serve to stimulate *ab initio* molecular dynamics simulations of this prototypical alkane following photoexcitation at VUV wavelengths and substantial refinements of the models currently used to describe the atmospheric photochemistry of the gas giants. This study (i) concludes that, as in CH_4 , the VUV photochemistry of ethane is driven by efficient non-adiabatic coupling to, and subsequent direct (and sequential) dissociations on the S_0 PES, (ii) highlights the need to revise current photochemical models of the stratospheric photochemistry of the gas giants – by incorporating the hitherto neglected C–H bond fission channel (7) and the three-body decomposition (6) to $CH_3 + {}^3CH_2 + H$ products and downgrading the relative yield of primary C_2H_x photoproducts – and (iii) emphasises the pressing need for quantitative product branching fractions. Stratospheric C_2H_6 production in the gas

giants is driven by VUV photodissociation of CH_4 , but the present analysis implies that the subsequent photochemical coupling between C_2H_6 and C_2H_2 is likely to be weaker than currently assumed.

Data and materials availability

The raw ion event lists, H-atom TOF spectra and calculation log files are available from the authors upon reasonable request.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (NSFC Center for Chemical Dynamics (Grant No. 21688102)), the National Natural Science Foundation of China (Grant No. 21673232, 21922306, and 21873099), the Strategic Priority Research Program of the Chinese Academy of Sciences (Grant No. XDB17000000), and the international partnership program of Chinese Academy of Sciences (No. 121421KYSB20170012). CSH is grateful for funding from the Australian Research Council (ARC, DE200100549), MNRA is grateful for funding from the Engineering and Physical Sciences Research Council (EPSRC, EP/L005913) and both CSH and MNRA are grateful to the NSFC Center for Chemical Dynamics for the award of Visiting Fellowships. The authors thank the FEL staff (Lei Shi, Hongli Ding, Kai Tao, Yuhuan Tian, Guanglei Wang, Dongxu Dai) for technical support, and Professor Satoshi Maeda and Dr Yu Harabuchi (Hokkaido University) for use of the GRRM software and computational resources.

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