PCCP



View Article Online **PAPER**

medium†



Cite this: Phys. Chem. Chem. Phys., 2025, 27, 11413

Mario Largo, Miguel Sanz-Novob and Pilar Redondo ** **

Formation of hydroxy, cyano and ethynyl

derivatives of C4H4 isomers in the interstellar

The study of cyclic hydrocarbons is of utmost relevance in current astrochemical research, as they are considered to be among the most significant reservoirs of carbon in the interstellar medium. However, while unsaturated cyclic hydrocarbons with three, five, and six carbon atoms have been widely investigated, the highly strained antiaromatic cyclobutadiene (c-C₄H₄) still remains uncharted. Here, we employed high-level CCSD(T)-F12/cc-pVTZ-F12//B2PLYPD3/aug-cc-pVTZ theoretical calculations to analyze whether the cyano (CN), ethynyl (CCH), and hydroxy (OH) derivatives of c-C₄H₄ and its structural isomers butatriene (H2CCCCH2) and vinylacetylene (H2CCHCCH) can readily form via the gasphase reaction: $C_4H_4 + X \rightarrow C_4H_3X + H$ (where X = CN, CCH, and OH). For each system, we thoroughly explored the corresponding potential energy surfaces, identifying their critical points to enable a detailed analysis of the thermochemistry. Hence, we found various exothermic pathways for the formation of CN and CCH derivatives of butatriene and vinylacetylene, with no net activation barriers, while the formation of the OH derivatives is in general less favorable. Prior to the mechanistic study, we also analyzed the complete conformational panorama and stability of all the derivatives at the CCSD(T)-F12/cc-pVTZ-F12 level. Overall, $c-C_4H_3CN$ and $c-C_4H_3CH$ emerge as particularly promising candidates for interstellar detection, provided that the parental c-C₄H₄ is present in the gas phase. These findings highlight the potential for detecting polar derivatives of c-C₄H₄ as indirect evidence of its presence in the ISM, as it appears to be "invisible" to radioastronomical observations. Also, this study underscores the need for future laboratory and theoretical efforts to characterize the spectroscopic properties of the proposed derivatives, paving the way for their eventual identification in space.

Received 27th February 2025, Accepted 1st May 2025

DOI: 10.1039/d5cp00781j

rsc.li/pccp

1 Introduction

In recent times, among the wide variety of molecules identified to date in the interstellar medium (ISM; see McGuire, Iimenez-Serra et al. for a recent census), cyclic hydrocarbons have drawn considerable attention from the astrophysical community. In particular, the presence of benzene, the simplest aromatic unit, and diverse polycyclic aromatic hydrocarbons (PAHs) has been confirmed through the detection of infrared (IR) emission features in a variety of interstellar environments.³⁻⁸

However, pure cyclic hydrocarbons often exhibit a very low or even zero permanent dipole moment, which makes them

"invisible" to rotational spectroscopy, further hampering their radioastronomical identification. An exception to this trend is cyclopropenylidene (c-C₃H₂; μ_b = 3.27 D), the first cyclic molecule identified in the ISM, 9-11 as well as cyclopentadiene (c-C₅H₆; μ_b = 0.416 D) and indene (c-C₉H₈; μ_a = 0.50 D and $\mu_b = 0.37 \text{ D}$, ¹² which have been recently detected toward the dark cold pre-stellar core Taurus molecular cloud 1 (TMC-1). This source accounts for the majority of new interstellar detections of cyclic systems, achieved during the course of two sensitive molecular line surveys: the Q-band ultrasensitive inspection journey to the obscure TMC-1 environment (QUIJOTE)¹² and the GBT observations of TMC-1: hunting aromatic molecules (GOTHAM).13 The complete list of interstellar cyclic molecules includes various cyano (-CN) and ethynyl (-CCH) derivatives, where one of the hydrogen atoms is replaced by a CN or CCH radical, highlighting benzonitrile (c-C₆H₅CN), ¹³ 1- and 2-cyanonaphthalene (c-C₁₀H₇CN), ¹⁴ the fivemembered ring 1-cyanocyclopentadiene (c-C₅H₅CN), ¹⁵ ethynylcyclopropenylidene (c-C₃H-CCH), 12 two isomers of ethynylcyclopentadiene (c-C₅H₅CCH), and its structural isomer

^a Computational Chemistry Group, Departamento de Química Física y Química Inorgánica, Universidad de Valladolid, E-47011 Valladolid, Spain. E-mail: pilar.redondo@uva.es

^b Centro de Astrobiología (CAB), CSIC-INTA, Carretera de Ajalvir km 4, Torrejón de Ardoz, 28850 Madrid, Spain

[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d5cp00781j

fulvenallene (c-C₅H₄CCH₂).¹⁷ Very recently, several isomers of the cyano-substituted derivatives of the three- and four-ring acenaphthylene (c-C₁₂H₇)¹⁸ and pyrene (c-C₁₆H₉)^{19,20} have also been identified toward TMC-1, ranking among the largest molecules ever detected in space.

In this context, while unsaturated hydrocarbons with three, five, and six carbon atoms have been successfully detected in the ISM, the more simple but also highly strained cyclobutadiene (also known as [4]annulene, c-C₄H₄) still remains uncharted. To date only its structural isomer vinylacetylene (CH₂CHCCH, global minimum in energy)²¹ has been detected in the ISM, along with the related cyano derivative isomers H₂CCHCCCN and trans-HCNCCHCCH. ^{22,23} This species is followed in energy by a cumulene form, butatriene (H₂CCCCH₂) located at 7.7 kcal mol⁻¹ above CH₂CHCCH at the CCSD(T)/ccpVTZ level, while c-C₄H₄ is found as the fourth most stable isomer, located at 33.4 kcal mol⁻¹.²¹ Its inherent instability can be rationalized in terms of the presence of 4n delocalized π electrons in it, which result in a so-called antiaromatic species. Nevertheless, in a recent study, Wang et al.24 have presented the first bottom-up formation of c-C₄H₄ in low-temperature acetylene (C2H2) ices exposed to energetic electrons, which simulates secondary electrons produced by the passage of cosmic rays. Once formed, c-C₄H₄ could perhaps be released into the gas phase through various thermal and non-thermal processes, serving as a promising candidate for astronomical searches, including James Webb Space Telescope (JWST) observations. Alternatively, its derivatization will infer a sizable dipole moment to the molecule and, therefore, its detection via rotational spectroscopy - both in the laboratory and in space - would be feasible. This, in turn, could serve as indirect evidence of the presence of c-C₄H₄ in the ISM.

Overall, the formation of CN and CCH derivatives of unsaturated hydrocarbons is suggested to proceed through the reaction of the hydrocarbon with the corresponding radicals, which occur rapidly at low temperatures.²⁵ This type of process has been proposed as pathways for the formation of c-C₆H₅CN, ¹³ c-C₁₀H₇CN, ¹⁴ and c-C₅H₅CN, ¹⁵ and are also thought to drive the relative abundance of the cyanopyrene isomers found in TMC-1.20 In addition, a theoretical study explored the possibility of forming derivatives of cyclopropenylidene c-C₃HX. Theoretically, the reaction of c-C₃H₂ with 16 radicals was analyzed, and four cyclopropenylidene derivatives (c-C₃HX, X = CN, OH, F, NH₂) were proposed as potential species for detection.26 In a following study, these authors investigated the possible formation of cyclopropenylidene disubstituted with CN and CCH radicals in space.²⁷

In this work, we have analyzed the formation processes of cyano, ethynyl and hydroxy derivatives of the structural isomers cyclobutadiene (c-C₄H₄), butatriene (H₂CCCCH₂), and vinylacetylene (H₂CCHCCH) through the following reaction: C₄H₄ + X \rightarrow C₄H₃X + H, where X = CN, CCH, and OH. We highlight that all of these radicals have already been detected in the interstellar medium, further motivating the exploration of the aforementioned routes. We have also studied the thermochemistry of the different processes and analyzed the critical points

of each potential energy surface (PES) to determine possible activation barriers.

2 Computational methods

Ab initio and density functional theory (DFT) methodologies have been employed to study the hydroxy, cyano, and ethynyl derivatives of cyclobutadiene (c-C₄H₄), butatriene (H₂CCCCH₂), and vinylacetylene (H2CCHCCH), along with all the intermediates and transition states identified on the respective potential energy surfaces (PES) associated with their formation reactions. We note that the selection of computational levels prioritizes maximum accuracy while maintaining reasonable computational efficiency.

At the DFT level, we have chosen the hybrid functionals M08HX²⁸ and MPWB1K,²⁹ which generally provide excellent results in both thermochemistry and kinetics. Additionally, the double-hybrid B2PLYPD3 functional30 was employed. This functional includes Hartree-Fock exchange and a perturbative second-order correlation part, together with a Grimmes D3BJ empirical dispersion term.³¹ For these three functionals, Dunning's correlation consistent triple-zeta basis set, aug-ccpVTZ, 32 was used, which includes both polarization and diffuse functions on all elements.

For ab initio calculations, we have selected the explicitly correlated coupled cluster theory with single and double excitations, including triplet excitations through a perturbative treatment, known as CCSD(T)-F12,33 in conjunction with the ccpVTZ-F12 basis set.³⁴ Recent studies³⁵ have shown that CCSD(T)-F12/cc-pVTZ-F12 optimized structures and energies are in excellent agreement with that obtained using a more complete "composite" scheme, but with a much lower computational cost. The calculated T1 diagnostic at the CCSD level³⁶ was found to be below 0.02 for all characterized structures. This value is within the accepted threshold for systems that are well described by a single-reference wavefunction, supporting the reliability of our single-reference calculations.

Harmonic vibrational frequency calculations were carried out for each optimized geometry at every of employed level of theory. This analysis allowed us to characterize the structures as either minima (all real frequencies) or transition states (with one imaginary frequency) and to determine the zero point vibrational energy (ZPVE). To verify that transition states connect the correct minima, intrinsic reaction coordinate (IRC) calculations³⁷ were performed for each transition state. Additionally, singlepoint energy calculations at the CCSD(T)-F12/cc-pVTZ-F12 level were performed on the DFT-optimized structures.

All calculations were performed using the GAUSSIAN 16,38 MOLPRO³⁹ program packages, which already implement the required methods, basis sets, and geometry optimization procedures.

3 Results and discussion

In this section, we will first analyze the geometrical parameters and the stabilities of the different isomers obtained by

substituting a hydrogen atom with the hydroxyl (OH), cyano (CN), and ethynyl (CCH) radicals in cyclobutadiene, butatriene and vinylacetylene hydrocarbons. Afterward, we will examine the reactions between these hydrocarbons and the three radicals, which allows us to assess the feasibility of forming such substituted species.

To streamline the description of the computational levels employed here, we will adopt the following notation throughout the discussion: M08HX, MPWB1K, and B2PLYPD3 refer to calculations performed at these levels in conjunction with the aug-cc-pVTZ basis set; CCSD(T)-F12 denotes calculations performed at the CCSD(T)-F12/cc-pVTZ-F12 level; and CC-F12// M08HX, CC-F12//MPWB and CC-F12//B2PLD refer to energy calculations at the CCSD(T)-F12/cc-pVTZ-F12 level using geometries optimized at the M08HX/aug-cc-pVTZ, MPWB1K/augcc-pVTZ, and B2PLYPD3/aug-cc-pVTZ levels, respectively.

3.1 Structure and stability of the OH, CN, and CCH derivatives of C₄H₄ isomers

The geometries of the cyclobutadiene derivatives, 2-hydroxycyclobuta-1,3-diene (c-C₄H₃OH), 2-cyanocyclobuta-1,3-diene (c-C₄H₃CN), and 2-ethynylcyclobuta-1,3-diene (c-C₄H₃CCH) are shown in Fig. 1, while the structures of the 1-hydroxybuta-1,2,3triene (H₂C₄HOH), 1-cyanobuta-1,2,3-triene (H₂C₄HCN), and 1ethynylbuta-1,2,3-triene (H₂C₄HCCH) are given in Fig. 2. Since the four hydrogen atoms of vinylacetylene are non-equivalent, the substitution of one of them will lead to four different structures. For instance, the cyano-substituted compounds identified are 1-vinylcyano-acetylene (H2CCHCCCN), 1-vinyl-1cyano-acetylene (H2CCCNCCH), trans-1-vinyl-2-cyano-acetylene

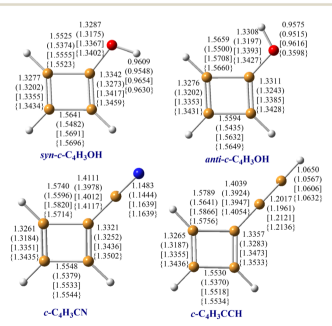


Fig. 1 Geometrical parameters of the cyclobutadiene derivates, $c-C_4H_3X$ with X = OH, CN, and CCH, calculated at the M08HX/aug-cc-pVTZ, MPWB1K/aug-cc-pVTZ (in parentheses), B2PLYPD3/aug-cc-pVTZ (in brackets) and CCSD(T)-F12/cc-pVTZ-F12 (in curly bracket) levels. Distances are given in Angstroms.

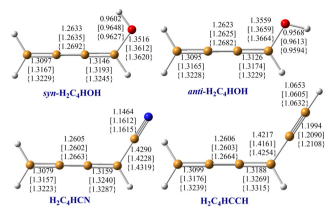


Fig. 2 Geometrical parameters of the butatriene derivates, H2CCCCHX with X = OH, CN, and CCH, calculated at the M08HX/aug-cc-pVTZ, B2PLYPD3/aug-cc-pVTZ (in brackets) and CCSD(T)-F12/cc-pVTZ-F12 (in curly bracket) levels. Distances are given in Angstroms.

(trans-HCNCCHCCH) and cis-1-vinyl-2-cyano-acetylene (cis-HCNCCHCCH). The optimized structures of these vinylacetylene derivatives are shown in Fig. 3. For all the hydroxyl isomers, we have identified two conformers, distinguished by the orientation of the hydrogen atom in the hydroxyl group. These conformers are denoted with the prefixes syn or anti, indicating whether the hydrogen is oriented towards the double bond or in the opposite direction, respectively. The relative energies for the monosubstituted hydrocarbons obtained at the different computational levels are provided in Table 1.

Additionally, as a reference, we report the optimized geometries of cyclobutadiene, butatriene, and vinylacetylene, as well as those of the OH, CN, and CCH radicals, at the levels of calculation employed in this study. These geometries are reported as ESI† in Fig. S1, and the relative energies for the hydrocarbons are summarized in Table S1 (ESI†). The relative stability data given in Table S1 (ESI†) confirm that the most stable isomer is vinylacetylene, followed by butatriene and cyclobutadiene. The relative energy values calculated at the CC-F12//M08HX and CC-F12//B2PLD levels are nearly identical (7.64, 7.74 kcal mol⁻¹ for butatriene, and 33.22 and 33.06 kcal mol⁻¹ for cyclobutadiene, relative to vinylacetylene) and differ by less than 0.3 kcal mol⁻¹ from previously reported values at the CCSD(T) level using B3LYP-optimized geometries. 21,24 Our results obtained at the CCSD(T)-F12 level place butatriene at 9.03 kcal mol⁻¹ and cyclobutadiene at 32.97 kcal mol⁻¹ above vinylacetylene. This indicates that optimizing the geometries at the CCSD(T)-F12 level has a higher impact on the energy than changing the type of functional. This fact is consistent with the differences observed in the geometries at the various levels (Fig. S1, ESI†), where it can be seen that the geometries optimized at the CCSD(T)-F12 level are the closest to the experimental data. 40-42 The variations between CCSD(T)-F12 and experimental values for the C-C bond lengths in the C_4H_4 isomers are on the order of hundredths of an angstrom.

We find that for monosubstituted hydrocarbons with OH, CN and CCH radicals, there are only slight variations in the structural parameters compared to the parent hydrocarbons.

Paper PCCP

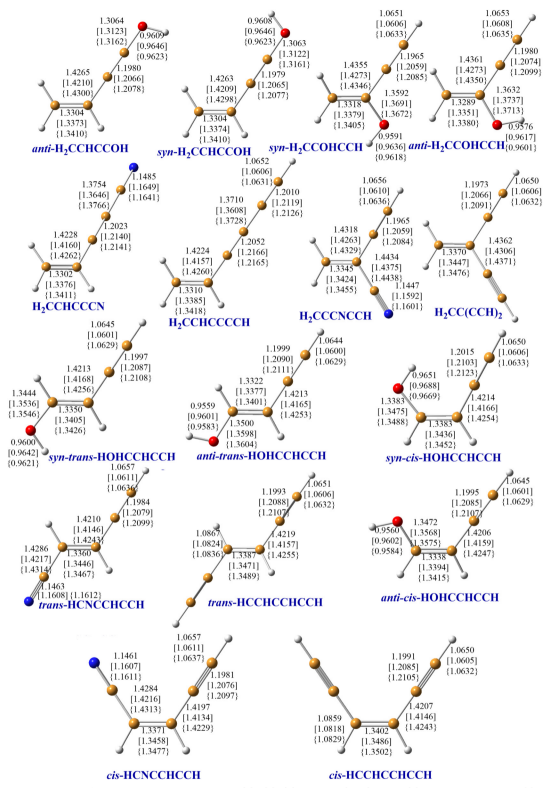


Fig. 3 Geometrical parameters of the vinylacetylene derivates, H_2 CCHCCH(X) with X = OH, CN, and CCH, calculated at the M08HX/aug-cc-pVTZ, B2PLYPD3/aug-cc-pVTZ (in brackets) and CCSD(T)-F12/cc-pVTZ-F12 (in curly bracket) levels. Distances are given in Angstroms.

These changes are primarily observed in the carbon bond where the substitution occurs. Specifically, the C-C distance (or distances) adjacent to the substituted hydrogen increases, as shown in Fig. 1–3. Regarding the different calculation levels

employed, it can be seen that the geometries optimized using the double hybrid functional B2PLYPD3 are the closest to those calculated at the CCSD(T)-F12 level, while the geometries obtained using the hybrid functional MPWB1K are the furthest.

PCCP

Table 1 Relative energies (in kcal mol⁻¹) of the hydroxy, cyano and ethynyl derivatives of cyclobutadiene, butatriene and vinylacetylene computed at different levels; ZPV energies are included

Molecule	M08HX	CC-F12//M08HX	B2PLYPD3	CC-F12//B2PLD	CCSD(T)-F12
OH derivatives from the c-C ₄ H ₄ /H ₂	CCCCH ₂ /H ₂ CCHC	CH + OH reaction			
anti-c-C ₄ H ₃ OH (¹ A')	37.28	34.62	39.41	34.52	35.66
syn-c-C ₄ H ₃ OH (¹ A')	36.46	33.81	38.55	33.70	34.55
syn-H ₂ C ₄ HOH (¹ A')	10.89	12.29	10.50	12.40	13.41
anti-H ₂ C ₄ HOH (¹ A')	13.19	14.54	12.88	14.68	16.18
anti-H ₂ CCHCCOH (¹ A')	12.68	13.52	13.39	13.67	14.67
syn-H ₂ CCHCCOH (¹ A')	12.67	13.52	13.37	13.66	14.70
syn-H ₂ CCOHCCH (¹ A')	5.86	4.81	5.26	4.73	6.02
anti-H ₂ CCOHCCH (¹ A')	4.89	3.95	4.45	3.97	5.39
syn-trans-HOHCCHCCH (¹ A')	3.33	3.10	3.09	3.12	3.73
anti-trans-HOHCCHCCH (1A')	3.97	3.75	3.78	3.77	4.77
syn-cis-HOHCCHCCH (¹ A')	0.00	0.00	0.00	0.00	0.00
anti-cis-HOHCCHCCH (1A')	3.89	3.52	4.12	4.13	4.68
CN derivatives from the c-C ₄ H ₄ /H ₂ !	CCCCH ₂ /H ₂ CCHC	CH + CN reaction			
$c-C_4H_3CN$ ($^1A'$)	33.56	30.36	35.21	30.20	31.14
$H_2C_4HCN(^1A')$	7.08	9.03	7.65	9.16	9.05
H ₂ CCHCCCN (¹ A')	-0.23	0.05	-1.48	0.13	0.07
H ₂ CCCNCCH (¹ A')	3.81	3.00	3.58	2.91	2.92
trans-HCNCCHCCH (¹ A _g)	0.00	0.00	0.00	0.00	0.00
cis-HCNCCHCCH (¹Ag)	0.45	0.34	0.28	0.34	0.32
CCH derivatives from the c-C ₄ H ₄ /H	I ₂ CCCCH ₂ /H ₂ CCH	CCH + CCH reaction			
c-C ₄ H ₃ CCH (¹ A')	36.11	32.36	38.67	32.12	32.52
$H_2C_4HCCH(^1A')$	10.13	11.46	11.64	11.56	11.51
H ₂ CCHCCCCH (¹ A')	0.00	0.00	0.00	0.00	0.00
$H_2CC(CCH)_2$ ($^1A'$)	6.67	5.16	7.47	5.07	5.19
trans-HCCHCCHCCCH (¹ A _g)	2.89	2.33	3.94	2.28	2.34
cis-HCCHCCHCCCH (¹ A _g)	3.22	2.58	4.17	2.61	2.63

Therefore, the latter has only been used for cyclobutadiene derivatives.

The relative stability of the isomers c-C₄H₃X, H₂C₄HX, and $H_2CCHCC(X)$, with X = OH, CN and CCH (Table 1), follows a similar trend for all the radicals: the monosubstituted cyclobutadiene isomers are the most unstable species in all cases, while the monosubstituted vinylacetylene isomers are the most stable. This trend mirrors that observed for C₄H₄ hydrocarbons.

Afterward, we will focus on the stability of the four vinyl acetylene derivatives as a function of the substitution position. The monosubstituted hydrocarbons, in one of the two hydrogen atoms attached to the terminal carbon of the vinyl group, cis-HXCCHCCH and trans-HXCCHCCH exhibit similar stabilities, with energy differences of less than 1 kcal mol⁻¹ for the CCH radical. In contrast, substitution at the non-terminal hydrogen of the vinyl group leads to the most unstable compounds, H2CCXCCH. For the hydroxy radical, the most favorable structures are obtained by substitution of the terminal hydrogens of the vinyl group, with the most stable isomer being syn-cis-HOHCCHCCH. When the ethynyl radical is involved, the substitution of the acetylenic hydrogen gives rise to the most stable structure, H2CCHCCCCH. Whereas for the cyano radical, substitutions at the terminal hydrogens of the vinyl and acetylene groups results in quasi-isoenergetic isomers (the energy differences between H2CCHCCCN, cis-HCNCCHCCH and trans-HCNCCHCCH isomers are less than $0.5 \text{ kcal mol}^{-1}$). These relative stabilities arise from various factors. The formation of H_2 CCHCCX isomers where the radicals have π -type electrons favors delocalization along the carbon chain. Conversely, more electrophilic substituents prefer to attack positions where the carbon has a larger negative partial charge (i.e., the terminal carbon of the vinyl group). Another factor to take into consideration is the steric effect, which may make substitution at less crowded positions more favorable when replacing a hydrogen atom with radicals.

An analysis of the relative energies at the different computational levels presented in Table 1 shows that the CC-F12// M08HX and CC-F12//B2PLD results are closer to those obtained at the CCSD(T)-F12 level than those derived at the DFT levels (M08HX or B2PLYPD3). The results that most closely match the CCSD(T)-F12 data are those calculated using the B2PLYPD3 geometries (CC-F12//B2PLD), with energy differences of less than 0.5 kcal mol⁻¹ for the CN and CCH derivatives, and approximately 1 kcal mol⁻¹ for the OH derivatives.

3.2 Formation of the OH, CN, and CCH derivatives of C₄H₄ isomers

As a potential route for the formation of the hydroxyl, cyano and ethynyl derivatives of cyclobutadiene, butatriene and vinylacetylene, we have considered the following reaction:

$$C_4H_4 + X \rightarrow C_4H_3X + H \tag{1}$$

where C₄H₄ represents the three isomers obeying the general formula C₄H₄ and X denotes the OH, CN or CCH radicals.

The intermediates (I) and transition states (TS) located on the different potential energy surfaces (PES) are labeled using a subscript indicating the related hydrocarbon (cyc, but, or vin, for reactions with cyclobutadiene, butatriene, or vinylacetylene) and the radical (OH, CN, or CCH) involved in the reaction under study. The reactions studied all proceed on the doublet PES. Geometric optimizations have been performed at the CCSD(T)-F12 level exclusively for reactants and products. The relative energies calculated at this level on the DFT-optimized geometries (CC-F12//DFT) are in close agreement with those obtained directly at the CCSD(T)-F12 level. In particular, the reaction energies computed at the CC-F12//B2PLD level generally differ by about 0.5 kcal mol⁻¹ from the CCSD(T)-F12 values.

3.2.1 Reaction of cyclobutadiene with OH, CN, and CCH. The relative energies of the characterized stationary points for the reactions between cyclobutadiene and the OH, CN, and CCH radicals are shown in Table 2 for the different calculation levels. The corresponding reaction profiles are schematically depicted in Fig. 4.

The formation of hydroxy, cyano, and ethynyl derivatives of cyclobutadiene are exothermic processes. The most favorable one is the production of c-C₄H₃CCH followed by that of c-C₄H₃CN, whose reaction energies are -30.27 and -22.62 kcal mol⁻¹ at the CCSD(T)-F12 level, respectively. The formation of the two hydroxyl-substituted conformers is slightly exothermic (with energies of -0.58 and -1.51 kcal mol⁻¹ at the CCSD(T)-F12 level for *anti*-c-C₄H₃OH and *syn*-c-C₄H₃OH,

respectively). Thus, from an energetic point of view, the formation would be possible at the low temperature conditions of the ISM, provided that barrierless reaction pathways are available.

The reaction mechanism for the substitution process is analogous for all three radicals, as illustrated in Fig. 4, which depicts the characterized stationary points in their respective potential energy surfaces (PESs). For simplicity, in the reaction involving the OH radical, only the reaction pathways corresponding to the *syn* conformers have been represented, while the energy values for the *anti* conformers are provided in Table 2.

As a representative example, we consider the reaction mechanism of cyclobutadiene with the CN radical. The process begins with the insertion of the CN radical into one of the carbons giving rise to the intermediate denoted as $I1_{\rm cyc-CN}$ located at -95.75 kcal $\rm mol^{-1}$ below the reactants at the CC-F12//B2PLD level. This step does not involve any transition state. Once the intermediate $I1_{\rm cyc-CN}$ is formed, the elimination of the hydrogen atom through the transition state $\rm TS1_{\rm cyc-CN}$ (located about -23.05 kcal $\rm mol^{-1}$ at the CC-F12//B2PLD level) leads to the product c-C₄H₃CN. Alternatively, the intermediate $\rm I1_{\rm cyc-CN}$ can undergo isomerization to form the more stable intermediate $\rm I2_{\rm cyc-CN}$ (-101.79 kcal $\rm mol^{-1}$ at the CC-F12//B2PLD level) through hydrogen migration from the carbon where the CN insertion occurred to the adjacent carbon. This

Table 2 Relative energies (in kcal mol⁻¹) for the stationary points located along the gas-phase reaction paths of cyclobutadiene with hydroxyl, cyano, and ethynyl radicals computed at different levels; ZPV energies are included

Molecule	M08HX	CC-F12//M08HX	MPWB1K	CC-F12//MPWB	B2PLYPD3	CC-F12//B2PLD	CCSD(T)-F12
Reaction c-C ₄ H ₄ + OF	· I						
$c-C_4H_4 + OH$	0.00	0.00	0.00	0.00	0.00	0.00	0.00
anti-c-C ₄ H ₃ OH + H	-2.86	-0.66	-2.44	-0.57	-1.17	-0.80	-0.58
syn-c-C ₄ H ₃ OH + H	-3.68	-1.48	-3.37	-1.42	-2.03	-1.62	-1.51
anti-I1 _{cyc-OH}	-75.96	-73.02	-78.08	-72.93	-74.05	-72.51	
syn-I1 _{cvc-OH}	-74.32	-71.26	-76.35	-71.14	-72.30	-70.72	
anti-I2 _{cyc-OH}	-80.75	-75.78	-83.09	-75.78	-77.88	-75.42	
syn-I2 _{cvc-OH}	-80.51	-75.61	-82.92	-75.64	-77.71	-75.30	
anti-TS1 _{cvc-OH}	-1.78	0.02	-2.01	-0.23	1.94	1.29	
syn -TS1 $_{cvc-OH}$	-2.27	-0.50	-3.13	-0.61	0.79	0.57	
anti- $TS2_{cyc-OH}$	-28.51	-22.95	-30.10	-22.99	-25.81	-22.95	
syn -TS2 $_{cyc-OH}$	-29.73	-24.20	-31.29	-24.13	-26.99	-24.07	
anti-TS3 _{cyc-OH}	-3.36	-0.56	-2.48	-1.01	-1.24	-0.83	
syn-TS3 _{cyc-OH}	-4.03	-1.33	-3.31	-1.75	-2.07	-1.69	
TS4-I1 _{cyc-OH}	-73.76	-70.58	-75.79	-70.56	-71.68	-70.13	
$TS5-I2_{cyc-OH}$	-77.63	-72.98	-79.93	-73.01	-74.69	-72.60	
Reaction c-C ₄ H ₄ + CN	1						
$c-C_4H_4 + CN$	0.00	0.00	0.00	0.00	0.00	0.00	0.00
$c-C_4H_3CN + H$	-30.88	-23.65	-31.40	-23.64	-28.07	-23.51	-22.62
$I1_{cvc-CN}$	-106.45	-96.48	-109.45	-96.47	-101.40	-95.75	
I2 _{cyc-CN}	-113.45	-102.76	-117.48	-102.75	-108.48	-101.79	
$TS1_{cyc-CN}$	-30.26	-22.74	-34.06	-22.43	-23.82	-23.05	
$TS2_{cyc-CN}$	-61.07	-49.36	-64.01	-49.34	-55.94	-48.78	
$TS3_{cyc-CN}$	-31.20	-23.58	-31.06	-23.55	-23.64	-22.16	
Reaction c-C ₄ H ₄ + CC	ЭН						
$c-C_4H_4 + CCH$	0.00	0.00	0.00	0.00	0.00	0.00	0.00
$c-C_4H_3CCH + H$	-34.39	-30.90	-35.66	-31.01	-35.60	-30.86	-30.27
I1 _{cvc-CCH}	-107.08	-101.08	-110.74	-101.16	-106.00	-100.40	
I2 _{cyc-CCH}	-117.35	-110.08	-122.10	-110.52	-116.28	-109.49	
$TS1_{cyc-CCH}$	-33.73	-30.07	-39.12	-57.26	-32.77	-30.86	
TS2 _{cyc-CCH}	-64.67	-56.66	-68.25	-56.75	-63.38	-56.14	
TS3 _{cyc-CCH}	-34.85	-30.94	-35.56	-31.16	-35.86	-31.02	

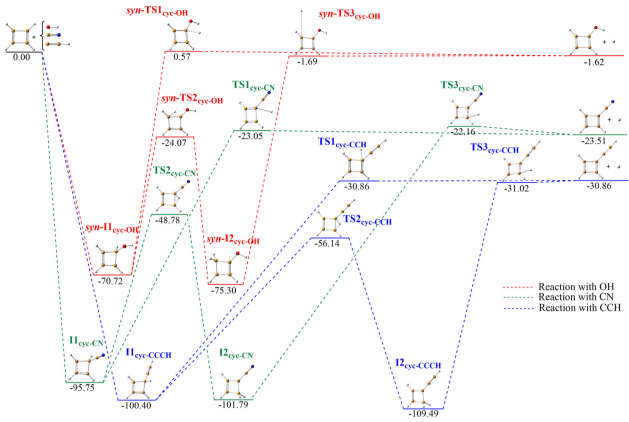


Fig. 4 Relative energies (in kcal mol⁻¹) for the stationary points located along the gas phase reaction of cyclobutadiene with hydroxyl, cyano, and ethynyl radicals computed at the CCSD(T)-F12/cc-pVTZ-F12//B2PLYPD3/aug-cc-pVTZ level. ZPV energies are included.

step involves the transition state TS2_{cvc-CN} located -48.78 kcal mol⁻¹ below the reactants at the CC-F12//B2PLD level. Subsequently, the elimination of one of the hydrogen atoms bonded to the carbon in I2_{cvc-CN} also yields the formation of c-C₄H₃CN. This step involves the transition state TS3_{cvc-CN}, which is clearly below the reactants $(-22.16 \text{ kcal mol}^{-1} \text{ at the CC-F12//B2PLD level})$. Therefore, the formation of c-C₄H₃CN via both pathways proceeds without a net activation barrier, indicating that the reaction is feasible under ISM conditions. The reaction mechanisms can be summarized as follows:

$$c-C_4H_4 + CN \rightarrow I1_{cyc-CN} \rightarrow TS1_{cyc-CN} \rightarrow c-C_4H_3CN + H$$
(2)

$$\begin{array}{c} \text{c-C}_4\text{H}_4 + \text{CN} \rightarrow \text{I1}_{\text{cyc-CN}} \rightarrow \text{TS2}_{\text{cyc-CN}} \rightarrow \text{I2}_{\text{cyc-CN}} \rightarrow \text{TS3}_{\text{cyc-CN}} \rightarrow \\ \text{c-C}_4\text{H}_3\text{CN} + \text{H} \end{array} \tag{3}$$

Regarding the reaction between cyclobutadiene and the ethynyl radical, it shows a reaction profile analogous to that observed for the corresponding reaction with the cyano radical. However, in the case of the CCH radical, the identified stationary points and the final product possess greater relative stability compared to those found in the CN system, with energy differences ranging from 10 to 20 kcal mol⁻¹. As illustrated in Fig. 4, the transition states TS1_{cyc-CCH} and TS3_{cvc-CCH}, when zero-point vibrational energy (ZPVE)

corrections are included, show energies equal to or slightly lower than that of the reaction product, c-C₄H₃CCH.

As previously indicated, the formation of the two conformers of the hydroxy derivatives, syn- and anti-c-C₄H₃OH, is only barely exothermic. As shown in Table 2 and Fig. 4, both synand anti-TS1_{cvc-OH} transition states lie slightly above the reactants $(0.57 \text{ kcal mol}^{-1} \text{ and } 1.29 \text{ kcal mol}^{-1} \text{ at the CC-F12}//$ B2PLD, respectively), suggesting that the formation of the hydroxy derivative could proceed in the ISM via the syn- and anti-12_{cyc-OH} intermediates. This path involves the transition states syn- and anti-TS3cvc-OH lying below the reactants $(-0.83 \text{ kcal mol}^{-1} \text{ and } -1.69 \text{ kcal mol}^{-1} \text{ at the CC-F12}//$ B2PLD, respectively). In Table 2, for the reaction of $c-C_4H_4$ with radical OH we include transition states TS4-I1_{evc-OH} and TS4-I2_{cvc-OH}, which correspond to the interconversion between the syn and anti conformers of intermediates I1_{cvc-OH} and I2_{cvc-OH}, respectively. As shown, the activation barriers for interconversion are significantly lower than the energy of the reactants (-70.13 and -72.60 kcal mol⁻¹ at the CC-F12//B2PLD level of theory, respectively). This suggests that interconversion to the more stable conformer is energetically favorable under ISM conditions.

In summary, the results for the reaction of cyclobutadiene with OH, CN, and CCH radicals indicate that the formation of cyano, ethynyl, and hydroxy derivatives of cyclobutadiene is thermodynamically feasible and involves no net activation barriers. Among these, the most favorable processes involve

Paper

Table 3 Relative energies (in kcal mol⁻¹) for the stationary points located along the gas-phase reaction paths of butatriene with hydroxyl, cyano, and ethynyl radicals computed at different levels; ZPV energies are included

Molecule	M08HX	CC-F12//M08HX	B2PLYPD3	CC-F12//B2PLD	CCSD(T)-F12
Reaction H ₂ CCCCH ₂ + OH					
$H2C_4H_4 + OH$	0.00	0.00	0.00	0.00	0.00
syn-H ₂ C ₄ HOH + H	1.60	2.59	2.41	2.40	2.49
anti-H ₂ C ₄ HOH + H	3.90	4.83	4.80	4.68	4.75
syn-H ₂ CCOHCCH + H	-3.43	-4.90	-2.82	-5.27	-5.06
anti-H ₂ CCOHCCH + H	-4.39	-5.76	-3.63	-6.03	-5.86
syn-I1 _{but-OH}	-47.05	-47.25	-44.96	-47.03	
anti-I1 _{but-OH}	-48.19	-48.33	-46.15	-48.04	
syn-TS1 _{but-OH}	-0.26	-1.97	0.05	-1.89	
anti-TS1 _{but-OH}	-1.11	-2.77	-0.71	-2.64	
Reaction H ₂ CCCCH ₂ + CN					
$H_2C_4H_2 + CN$	0.00	0.00	0.00	0.00	0.00
$H_2C_4HCN + H$	-26.50	-19.40	-23.13	-19.23	-19.27
H ₂ CCCNCCH + H	-29.78	-25.44	-27.20	-25.49	-25.40
I1 _{but-CN}	-77.08	-70.21	-71.98	-69.72	
I2 _{but-CN}	-75.77	-68.43	-70.50	-68.06	
$TS1_{but-CN}$	-26.54	-22.33	-23.73	-21.78	
TS2 _{but-CN}	-23.07	-16.37	-18.61	-15.67	
$TS3_{but-CN}$	-38.48	-28.90	-30.84	-28.06	
Reaction H ₂ CCCCH ₂ + CCH					
$H_2C_4H_2 + CCH$	0.00	0.00	0.00	0.00	0.00
$H_2C_4HCCH + H$	-29.52	-26.23	-30.12	-26.10	-25.85
$H_2CC(CCH)_2 + H$	-32.98	-32.53	-34.30	-32.59	-32.18
I1 _{but-CCH}	-79.13	-76.41	-78.04	-75.92	
I2 _{but-CCH}	-77.06	-73.96	-75.88	-72.26	
TS1 _{but-CCH}	-29.92	-29.70	-31.02	-29.10	
$TS2_{but-CCH}$	-26.17	-23.40	-25.55	-22.68	
$TS3_{but-CCH}$	-46.74	-41.19	-43.23	-40.39	

the formation of c-C₄H₃CN and c-C₄H₃CCH derivatives, which appears as thrilling candidates for interstellar detection. Moreover, this derivatization infers a significant dipole moment to the parental and apolar c-C₄H₄, enabling its detection by means of rotational spectroscopy in the laboratory but also in space. In this context, the main challenge for such measurements lies in generating sufficient quantities of these highly unstable antiaromatic species in the gas phase. However, emerging laboratory techniques, such as direct-absorption millimeter and submillimeter spectroscopy of desorbed species from ice samples, 43 may succeed where conventional spectroscopic methods are likely to fail. Furthermore, the eventual detection of CN and CCH derivatives could serve as indirect evidence for the presence of $c-C_4H_4$ in the interstellar medium (ISM).

3.2.2 Reaction of butatriene with OH, CN, and CCH. The stationary points on the doublet potential energy surfaces (PESs) corresponding to the reactions between butatriene and the hydroxy, cyano, and ethynyl radicals at various computational levels are summarized in Table 3. The reaction profiles are depicted in Fig. 5. As in the case of cyclobutadiene reacting with OH, only the pathway associated with the anti conformer is presented, as it is the most thermodynamically favorable.

As shown in Table 3, the formation of 1-cyanobuta-1,2,3triene (H₂C₄HCN) and 1-ethynylbuta-1,2,3-triene (H₂C₄HCCH) are significantly exothermic, with reaction energies of -19.27kcal mol⁻¹ and -25.85 kcal mol⁻¹, respectively, at the CCSD(T)-F12 level. On the other hand, the formation of 1-hydroxybuta-1,2,3-triene (H₂C₄HOH) is slightly endothermic for both the syn

and anti conformers, with computed reaction energies of 2.49 kcal mol⁻¹ and 4.75 kcal mol⁻¹, respectively, at the same level of theory. Consequently, the formation of these hydroxycontaining species is not feasible under the low-temperature conditions characteristic of the ISM. Additionally, we consider the vinyl acetylene derivatives 1-vinyl-1-cyano-acetylene (H2CCCNCCH), and 1-vinyl-1-ethynyl-acetylene (H₂CC(CCH)₂), and the syn- and anti-1-vinyl-1-hydroxy-acetylene (H2CCOHCCH) as potential reaction products, as they can also form through the interaction of butatriene with the aforementioned radicals. These reaction pathways are exothermic, with reaction energies of -25.40, -32.18, -5.06, and -5.86 kcal mol⁻¹, respectively, at the CCSD(T)-F12 level. The corresponding reaction pathways for their formation are also illustrated in Fig. 5.

We then analyze the reaction profile starting with the interaction between butatriene and the CN radical as a reference. The CN radical can either bond to one of the central carbon atoms of butatriene, leading to the intermediate I1but-CN, or to one of the terminal carbon atoms, yielding intermediate $I2_{but-CN}$. An analysis of the reaction coordinate in both cases indicates that these intermediates form without passing through a transition state. Both are significantly more stable than the reactants, with relative energies of -69.72 kcal mol⁻¹ and -68.06 kcal mol⁻¹, respectively, at the CC-F12//B2PLD level. From I1_{but-CN}, the elimination of a hydrogen atom from the terminal carbon farthest from the CN group leads to the formation of the most exothermic product, the vinylacetylene derivative H2CCCNCCH. This transformation occurs through

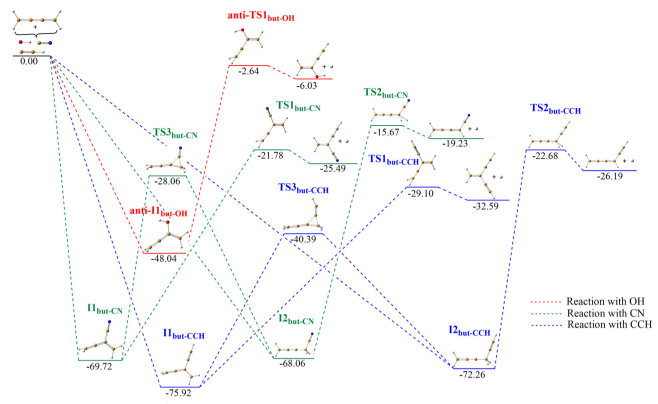


Fig. 5 Relative energies (in kcal mol⁻¹) for the stationary points located along the gas phase reaction of butatriene with cyano, and ethynyl radicals computed at the CCSD(T)-F12/cc-pVTZ-F12//B2PLYPD3/aug-cc-pVTZ level. ZPV energies are included.

the TS1_{but-CN} transition state, which is located below the energy of the reactants $(-21.78 \text{ kcal}^{-1} \text{ at the CC-F12//B2PLD level})$. The overall reaction process can be summarized as follows:

$$\mathrm{H_{2}C_{4}H_{2}} + \mathrm{CN} \rightarrow \mathrm{I1_{but-CN}} \rightarrow \mathrm{TS1_{but-CN}} \rightarrow \mathrm{H_{2}CCCNCCH} + \mathrm{H}$$
(4)

Alternatively, if one of the hydrogens in I1_{but-CN} is removed from the carbon closest to the CN group, a cyclic product is formed. This process involves an activation barrier of more than 30 kcal mol⁻¹. For this reason, it has not been included, as it is, in principle, an unfeasible process under ISM conditions.

Starting from the intermediate I2_{but-CN}, the elimination of one hydrogen from the carbon to which the CN radical is attached leads to the formation of the butatriene derivative, H₂C₄HCN. This process takes place through the TS2_{but-CN} transition state, located -15.67 kcal mol⁻¹ below the reactants. The overall process can be summarized as:

$$H_2C_4H_2 + CN \rightarrow I2_{but-CN} \rightarrow TS2_{but-CN} \rightarrow H_2C_4HCN + H$$
 (5)

As can be seen in Fig. 5, the intermediate I2_{but-CN} can also isomerize to the more stable one, I1_{but-CN}, through the TS3_{but-} _{CN} transition state, which is located 28.06 kcal mol⁻¹ below the

Concerning the reaction of butatriene with the CCH radical, its profile closely parallels that of the reaction with the CN radical and, similarly to the reactions with cyclobutadiene, the various PES stationary points along the reaction paths are relatively more stable. Overall, the results derived for the reaction of butatriene with CN and CCH radicals indicate that the butatriene derivatives, H2C4HCN and H2C4HCCH, can be readily produced, but the formation of vinylacetylene derivatives, H₂CCCNCCH and H₂CC(CCH)₂, is even more favorable from the energetic (more exothermic) point of view and exhibits lower barriers.

A similar result is obtained for the reaction of H₂C₄H₂ with the OH radical, for which, from a thermodynamic point of view, only the formation of the vinylacetylene derivatives, syn- and anti-H₂CCOHCCH, is feasible (see Table 3). As shown in Fig. 5, these pathways proceed via the syn- and anti-syn-I1_{but-OH} intermediates by elimination of a hydrogen from the terminal carbon farthest from the OH group. The transition states involved, syn- and anti-TS1_{but-OH}, are located 1.86 and $2.64 \text{ kcal mol}^{-1}$ below the reactants. Therefore, the formation of syn- and anti-H₂CCOHCCH is also feasible.

3.2.3 Reaction of vinylacetylene with OH, CN, and CCH. Finally, we analyze the reactions of OH, CN, and CCH radicals with vinylacetylene. The relative energies of reactants, products, intermediates, and transition states characterized on their respective PESs - at the different calculation levels - are summarized in Table 4. The corresponding reaction profiles for OH, CN, and CCH are depicted in Fig. 6, 7, and 8, respectively.

We first consider the reaction of H₂CCHCCH with the OH radical. As can be seen in Table 4, the reaction is only exothermic for the formation of the vinylacetylene

Table 4 Relative energies (in kcal mol⁻¹) for the stationary points located along the gas-phase reaction paths of vinylacethylene with hydroxyl, cyano, and ethynyl radicals computed at different levels; ZPV energies are included

Molecule	M08HX	CC-F12//M08HX	B2PLYPD3	CC-F12//B2PLD	CCSD(T)-F12
Reaction H ₂ CCHCCH + OH					
H_2 CCHCCH + OH	0.00	0.00	0.00	0.00	0.00
syn-H ₂ CCHCCOH + H	8.72	11.44	11.47	11.40	11.47
anti-H ₂ CCHCCOH + H	8.73	11.45	11.48	11.41	11.50
syn-H ₂ CCOHCCH + H	1.91	2.74	3.36	2.47	2.59
anti-H ₂ CCOHCCH + H	0.95	1.87	2.55	1.71	1.80
syn-trans-HOHCCHCCH + H	-0.61	1.03	1.19	0.86	0.99
anti-trans-HOHCCHCCH + H	0.03	1.68	1.88	1.51	1.60
syn-cis-HOHCCHCCH + H	-3.94	-2.07	-1.90	-2.26	-2.12
anti-cis-HOHCCHCCH + H	-0.06	1.44	2.21	1.87	1.96
syn-I1 _{vin-OH}	-37.94	-35.56	-34.78	-35.46	
syn-I2 _{vin-OH}	-37.35	-35.61	-34.86	-35.37	
syn -TS1 $_{ m vin-OH}$	3.81	4.95	5.11	5.10	
syn-TS2 _{vin-OH}	0.39	1.79	1.93	1.90	
syn-TS3 _{vin-OH}	-36.60	-35.00	-34.11	-34.77	
Reaction H ₂ CCHCCH + CN					
H ₂ CCHCCH + CN	0.00	0.00	0.00	0.00	0.00
H ₂ CCHCCCN + H	-28.47	-20.75	-26.09	-20.52	-20.60
H ₂ CCCNCCH + H	-24.43	-17.80	-21.02	-17.75	-17.75
trans-HCNCCHCCH + H	-28.24	-20.80	-24.60	-20.65	-20.67
cis-HCNCCHCCH + H	-27.79	-20.45	-24.33	-20.31	-20.35
$I1_{vin-CN}$	-72.78	-63.54	-66.98	-62.89	
I2 _{vin-CN}	-52.53	-44.53	-46.87	-44.37	
I3 _{vin-CN}	-69.92	-61.09	-63.93	-60.54	
I4 _{vin-CN}	-69.45	-60.67	-63.60	-60.18	
TS1 _{vin-CN}	-23.14	-15.88	-19.80	-15.00	
TS2 _{vin-CN}	-16.20	-9.99	-12.49	-9.66	
TS3 _{vin-CN}	-24.18	-17.03	-20.09	-16.43	
TS4 _{vin-CN}	-23.72	-16.67	-19.81	-16.05	
TS5 _{vin-CN}	-9.76	-4.51	-9.10	-4.34	
TS6 _{vin-CN}	-68.70	-60.16	-63.10	-59.72	
Reaction H ₂ CCHCCH + CCH					
H ₂ CCHCCH + CCH	0.00	0.00	0.00	0.00	0.00
H ₂ CCHCCCCH + H	-34.30	-30.05	-35.59	-29.92	-29.71
$H_2CC(CCH)_2 + H$	-27.64	-24.89	-28.12	-24.85	-24.52
trans-HCCHCCHCCH + H	-31.42	-27.73	-31.65	-27.64	-27.37
cis-HCCHCCHCCH + H	-31.09	-27.48	-31.42	-27.31	-27.07
I1 _{vin-CCH}	-74.76	-69 . 47	-72.88	-68.83	
I2 _{vin-CCH}	-54.23	-50.25	-52.54	-50.13	
I3 _{vin-CCH}	-70 . 88	-65 . 97	-68.82	-65.51	
I4 _{vin-CCH}	-70.23	-65.39	-68.33	-64.97	
TS1 _{vin-CCH}	-28.73	-25.03	-29.03	-24.17	
TS2 _{vin-CCH}	-19.30	-17.09	-19.53	-16.79	
TS3 _{vin-CCH}	-27.35	-24.06	-27.09	-23.46	
TS4 _{vin-CCH}	-26.96	-23.74	-26.87	-23.10	
TS5 _{vin-CCH}	-4.56	-3.38	-7.49	-2.98	
TS6 _{vin-CCH}	-69.46	-64.86	-67.75	-64.45	

derivative, syn-cis-HOHCCHCCH with a reaction energy of -2.12 kcal mol⁻¹ at the CCSD(T)-F12 level. The reaction energy for obtaining the syn-trans-HOHCCHCCH isomer is close to zero at all the different levels employed. Therefore, we will consider the formation of these two products whose reaction profiles are shown in Fig. 6. When the OH radical interacts with the terminal carbon of the acetylene group, two intermediates, $\textit{syn-}\text{I1}_{\text{vin-OH}}$ and $\textit{syn-}\text{I2}_{\text{vin-OH}}$, are obtained, depending on the dihedral angle formed between the OH radical and the CCH group of vinylacetylene. The formation of these intermediates occurs directly, without the involvement of transition states and they are located below the reactants (-35.46, and -35.37 kcal mol⁻¹ at the CC-F12//B2PLD level, respectively). Subsequent hydrogen elimination from the OH-bonded carbon

in syn-I1 $_{vin-OH}$ and syn-I2 $_{vin-OH}$ leads to the formation of the products syn-cis-HOHCCHCCH and syn-trans-HOHCCHCCH. These processes proceed through the transition states syn- $TS1_{vin-OH}$ and syn- $TS2_{vin-OH}$, which lie 1.90 and 5.10 kcal mol⁻¹ above the reactants at the CC-F12//B2PLD level, respectively. Given these energy barriers, the formation of hydroxy derivatives of vinylacetylene through these reaction pathways are unlikely under the typical temperature and density conditions of the ISM.

On the other hand, the formation pathways of the four cyano-substituted derivatives of vinylacetylene, H2CCHCCCN, H₂CCCNCCH, trans-HCNCCHCCH, and cis-HCNCCHCCH, through the substitution of a hydrogen atom in vinylacetylene for the cyano (CN) radical, are exothermic, with reaction

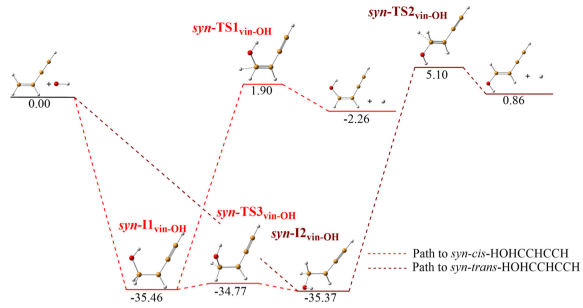
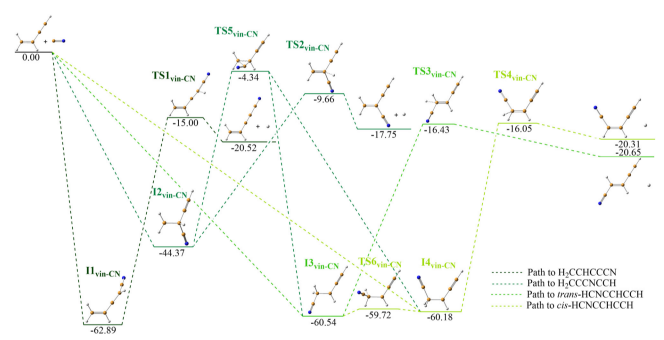


Fig. 6 Relative energies (in kcal mol⁻¹) for the stationary points located along the gas phase reaction of vinylacetylene with the hydroxyl radical computed at the CCSD(T)-F12/cc-pVTZ-F12//B2PLYPD3/aug-cc-pVTZ level. ZPV energies are included.



 $\textbf{Fig. 7} \quad \text{Relative energies (in kcal mol}^{-1}\text{) for the stationary points located along the gas phase reaction of vinylacetylene with cyano radical computed at the property of the stationary points located along the gas phase reaction of vinylacetylene with cyano radical computed at the property of the stationary points located along the gas phase reaction of vinylacetylene with cyano radical computed at the property of the stationary points located along the gas phase reaction of vinylacetylene with cyano radical computed at the property of the stationary points located along the gas phase reaction of vinylacetylene with cyano radical computed at the property of the stationary points located along the gas phase reaction of vinylacetylene with cyano radical computed at the property of the stationary points located along the gas phase reaction of vinylacetylene with cyano radical computed at the property of the stationary points located along the gas phase reaction of vinylacetylene with cyano radical computed at the property of th$ the CCSD(T)-F12/cc-pVTZ-F12//B2PLYPD3/aug-cc-pVTZ level. ZPV energies are included.

energies ranging from -17.75 to -20.67 kcal mol⁻¹, as summarized in Table 4. The corresponding reaction profiles for the formation of these cyano derivatives are depicted in Fig. 7.

The reaction initiates with the addition of the CN radical to one of the carbon atoms in vinylacetylene. When CN attaches to the carbon of the ethynyl moiety, intermediate I1_{vin-CN} is formed. Alternatively, if the CN radical binds to the carbon of the vinyl group that is bonded to just one hydrogen,

intermediate $I2_{vin-CN}$ is obtained. Furthermore, intermediates I3_{vin-CN} and I4_{vin-CN} arise from the attachment of CN to the terminal carbon of the vinyl group in different orientations. All intermediates lie significantly below the reactants, with relative energies of -62.89, -44.37, -60.54, and -60.18 kcal mol⁻¹ at the CC-F12//B2PLD level, respectively. From these intermediates, the elimination of a hydrogen atom bonded to the carbon where the CN radical was initially added results in the

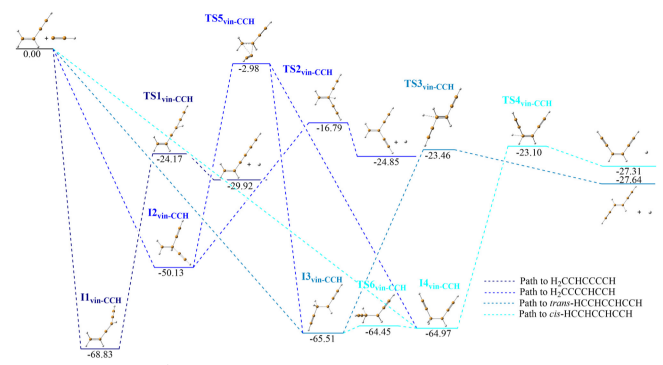


Fig. 8 Relative energies (in kcal mol⁻¹) for the stationary points located along the gas phase reaction of vinylacetylene with ethynyl radical computed at the CCSD(T)-F12/cc-pVTZ-F12//B2PLYPD3/aug-cc-pVTZ level. ZPV energies are included.

formation of the final cyano-substituted vinylacetylene products H2CCHCCCN, H2CCCNCCH, trans-HCNCCHCCH, and cis-HCNCCHCCH. These steps proceed through transition states TS1vin-CN, TS2vin-CN, TS3vin-CN, and TS4vin-CN, which are located -15.00, -9.66, and -16.43 and -16.05 kcal mol⁻¹ below the reactants at the CC-F12//B2PLD level, respectively. Additionally, we include in the reaction profile transition state $TS5_{vin-CN}$, located at -4.34 kcal mol^{-1} relative to the reactants at the CC-F12//B2PLD level, which corresponds to the isomerization of intermediate I2_{vin-CN} into the more stable ones I3_{vin-CN} and I4_{vin-CN}. We also characterized the transition state TS6_{vin-CN}, which is associated with the rotation of the terminal vinyl carbon, facilitating the interconversion of ${\rm I3_{vin-CN}}$ to ${\rm I4_{vin-}}$ CN. This transition state is positioned slightly above the intermediates, with a relative energy of -59.72 kcal mol⁻¹ at the CC-F12//B2PLD level. As shown in the Fig. 7, the formation pathways of the four cyano-substituted vinylacetylene isomers follow analogous mechanisms, which can be exemplified by the formation of H₂CCHCCCN:

$$\label{eq:H2CCHCCH+CN} \text{H}_2\text{CCHCCH} + \text{CN} \rightarrow \text{I1}_{\text{vin-CN}} \rightarrow \text{TS1}_{\text{vin-CN}} \rightarrow \text{H}_2\text{CCHCCCN} + \text{H}$$
 (6)

These results indicate that the formation of the four cyanosubstituted derivatives of vinylacetylene is thermodynamically feasible in the ISM from the proposed CN addition. However, determining the relative abundances of each isomer requires a detailed kinetic study of the processes, which falls beyond the scope of this work and will be undertaken in the near future. As noted in the introduction, two cyanovinylacetylene isomers, H2CCHCCCN and

trans-HCNCCHCCH, have been detected in the TMC-1 molecular cloud. 22,23 The reaction described here represents a potential synthetic pathway contributing to their formation.

As shown in Table 4, the formation of the four ethynylsubstituted derivatives of vinylacetylene, H₂CCHCCCCH, H₂CC(CCH)₂, trans-HCCHCCHCCH, and cis-HCCHCCHCCH, from the reaction between vinylacetylene and the CCH radical is exothermic, with reaction energies of -29.71, -24.52, -27.37, and -27.07 kcal mol⁻¹ at the CCSD(T)-F12 level, respectively. The corresponding reaction pathways leading to these products are depicted in Fig. 8. It is worth noting that these pathways closely resemble those described in Fig. 7 for the analogous reaction with the CN radical. The reaction between the CCH radical and vinylacetylene leads to the formation of four intermediates, I1_{vin-CCH}, I2_{vin-CCH}, I3_{vin-CCH}, and I4vin-CCH. These intermediates undergo hydrogen atom elimination from the carbon where the CCH radical has bonded, resulting in the final products. The transition states involved in these processes, TS1_{vin-CCH}, TS2_{vin-CCH}, TS3_{vin-CCH}, and $TS4_{vin-CCH}$, are located well below the reactants (-24.17, -16.79, -23.46, and -23.10 kcal mol⁻¹ at the CC-F12//B2PLD level). Thus, the formation of isomers H₂CCHCCCCH, H₂CC(CCH)₂, trans-HCCHCCHCCH, and cis-HCCHCCHCCH, from the CCH addition to vinylacethylene, is feasible under ISM conditions, as all processes proceed without a net activation barrier. In contrast, the higher TSs reported above for the reactions with the OH radicals can be rationalized in terms of a reduced delocalization of π -electrons in the double bonds, compared to CN- and CCH-bearing species, which benefit from greater stabilization due to a more extended conjugation.

3.3 Astrophysical implications

Our theoretical findings demonstrate that the cyano, ethynyl, and hydroxy derivatives of the highly strained antiaromatic cyclic c-C₄H₄, can readily form under ISM conditions, provided that the parental species c-C₄H₄ is available in the gas phase, as supported by its recent laboratory detection in low-temperature ice analogs.24 Subsequent desorption could occur through different thermal and non-thermal processes (e.g., in the hotcore stage or due to the action of large-scale shocks), depending on the physical properties of the astronomical environment. Furthermore, we emphasize that while c-C₄H₄ is "invisible" to radioastronomy due to its lack of a permanent dipole moment, the detection of its OH, CN and CCH derivatives could serve as indirect evidence of the presence of c-C₄H₄ in the ISM. Therefore, the present results highlight the need for future theoretical and laboratory studies on the spectroscopic characterization of these polar derivatives that facilitate their astronomical identification, which will help shed light on the role of antiaromatic systems in astrochemistry.

The results obtained for the reactions of butatriene with CN and CCH radicals suggest that the formation of butatriene derivatives, specifically H₂C₄HCN and H₂C₄HCCH, is energetically feasible. However, the formation of vinylacetylene derivatives, such as H₂CCCNCCH and H₂CC(CCH)₂, is even more favorable, exhibiting higher exothermicity and lower activation barriers. In the case of the OH radical, only the formation of vinylacetylene derivatives, syn- and anti-H2CCOHCCH, is found to be viable. On the basis of these results, kinetic analysis is necessary to fully assess the viability of butatriene derivative formation under interstellar conditions.

Four distinct isomers can be obtained via the CN and CCH addition reaction to vinylacetylene: H2CCHCCCN, H2CCCNCCH, trans-CNHCCHCCH, and cis-CNHCCHCCH, H2CCHCCCCH, H₂CC(CCH)₂, trans-HCCHCCHCCH, and cis-HCCHCCHCCH. Bearing in mind that H2CCHCCCN and trans-HCNCCHCCH, have been already detected in the cold dark cloud TMC-1,22,23 our results suggest that the CN and CCH addition reactions to vinylacetylene represent a plausible formation pathway leading to these species. In this context, future theoretical efforts will focus on analyzing the kinetics behind the proposed routes, which will be essential for deriving and subsequently rationalizing the observed isomeric ratio.

Overall, our results indicate that cyano and ethynyl derivatives can be readily formed from C₄H₄ hydrocarbons. In contrast, the formation of hydroxy derivatives appears significantly less favorable, despite the high abundance of OH radicals in the interstellar medium (ISM). These results are consistent with the fact that, to date, only CN and CCH derivatives of pure cyclic hydrocarbons have been definitely identified.

4 Conclusions

In this study, we have performed a theoretical investigation of the stability and formation processes for the cyano (CN), ethynyl (CCH) and hydroxy (OH) derivatives of the C4H4 structural isomers: cyclobutadiene (c-C₄H₄), butatriene (H₂CCCCH₂), and vinylacetylene (H2CCHCCH). In particular, we proposed the gasphase reaction: $C_4H_4 + X \rightarrow C_4H_3X + H$, where X = CN, CCH, and OH. For each system, we have analyzed in detail the thermochemistry and identified the critical points involved in each PES to determine possible activation barriers. The main conclusions of this work are the following:

- The results for the reaction starting with the cyclic system, cyclobutadiene (c-C₄H₄) have shown that the formation of cyano, ethynyl, and hydroxy derivatives are thermodynamically feasible and involve no net activation barriers. Among these, the most favorable pathways involve the formation of c-C4H3CN and c-C₄H₃CCH derivatives, which emerge as promising candidates for interstellar detection via JWST (IR) observations or through radioastronomical observations, once rotational laboratory spectroscopic data become accessible.
- Regarding the formation of the butatriene (H₂CCCCH₂) derivatives, we found that endothermic reaction pathways are unfeasible to the formation of the hydroxy derivatives under ISM conditions. However, the suggested PESs also allow the formation of various vinylacetylene derivatives syn- and anti-H₂CCOHCCH, which proceeds with no net activation barrier. In contrast, the CN and CCH addition reactions are exothermic and lead to the formation of H₂C₄HCN and H₂C₄HCCH without net activation barriers. Interestingly, we also found that the formation of the vinylacetylene derivatives H2CCCNCCH and H₂CC(CCH)₂ is even more favorable from the energetic (more exothermic) point of view and shows lower barriers.
- As for vinylacetylene (H2CCHCCH), while the formation of the OH derivatives is again unlikely under the typical conditions of the ISM due to the net activation barrier, we have identified feasible routes for the formation of all CN and CCH substituted derivatives. In this context, future theoretical efforts will focus on analyzing the kinetics behind the proposed routes.
- Finally, concerning the different computational levels employed in this study, the reaction energies calculated at the CCSD(T)-F12 level deviate by less than 1.0 kcal mol⁻¹ from those obtained at the CC-F12//B2PLD level, where single-point energy calculations were performed on geometries optimized at the B2PLYPD3 level. The results obtained with this approach are highly consistent. Therefore, the influence of molecular geometry is comparatively less significant than the incorporation of electron correlation energy.

Author contributions

PR and MSN conceived the project. PR and ML carried out the quantum-chemical computations. PR and MSN drafted the manuscript. All authors discussed the results and reviewed the manuscript.

Data availability

The data supporting this article have been included as part of the ESI.† Additional data are available from the corresponding author upon reasonable request.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

Financial support from the Spanish Ministerio de Ciencia e Innovación (PID2020-117742GB-I00/AEI/10.13039/501100011033) is gratefully acknowledged. M. S.-N. acknowledges the Juan de la Cierva Postdoctoral Fellow project JDC2022-048934-I, funded by the Spanish Ministry of Science, Innovation and Universities/State Agency of Research MICIU/AEI/10.13039/501100011033 and by the European Union "NextGenerationEU/PRTR". M. S.-N. also acknowledges funding from grant No. PID2022-136814NB-I00 funded by MICIU/AEI/10.13039/501100011033 and by ERDF, UE.

References

- 1 B. A. McGuire, Astrophys. J., Suppl. Ser., 2022, 259, 30.
- 2 I. Jimenez-Serra, C. Codella and A. Belloche, arXiv, 2025, preprint, arXiv:2503.17104, DOI: 10.48550/arXiv.2503.17104.
- 3 A. G. G. M. Tielens, Annu. Rev. Astron. Astrophys., 2008, 46, 289-337.
- 4 I. Garca-Bernete, D. Rigopoulou, A. Alonso-Herrero, F. R. Donnan, P. F. Roche, M. Pereira-Santaella, A. Labiano, L. Peralta de Arriba, T. Izumi, C. Ramos Almeida, T. Shimizu, S. Hönig, S. Garca-Burillo, D. J. Rosario, M. J. Ward, E. Bellocchi, E. K. S. Hicks, L. Fuller and C. Packham, Astron. Astrophys., 2022, 666, L5.
- 5 R. Chown, A. Sidhu, E. Peeters, A. G. G. M. Tielens, J. Cami, O. Berné, E. Habart, F. Alarcón, A. Canin, I. Schroetter, B. Trahin, D. Van De Putte, A. Abergel, E. A. Bergin, J. Bernard-Salas, C. Boersma, E. Bron, S. Cuadrado, E. Dartois, D. Dicken, M. El-Yajouri, A. Fuente, J. R. Goicoechea, K. D. Gordon, L. Issa, C. Joblin, O. Kannavou, B. Khan, O. Lacinbala, D. Languignon, R. Le Gal, A. Maragkoudakis, R. Meshaka, Y. Okada, T. Onaka, S. Pasquini, M. W. Pound, M. Robberto, M. Röllig, B. Schefter, T. Schirmer, S. Vicente, M. G. Wolfire, M. Zannese, I. Aleman, L. Allamandola, R. Auchettl, G. A. Baratta, S. Bejaoui, P. P. Bera, J. H. Black, F. Boulanger, J. Bouwman, B. Brandl, P. Brechignac, S. Brünken, M. Buragohain, A. Burkhardt, A. Candian, S. Cazaux, J. Cernicharo, M. Chabot, S. Chakraborty, J. Champion, S. W. J. Colgan, I. R. Cooke, A. Coutens, N. L. J. Cox, K. Demyk, J. D. Meyer, S. Foschino, P. Garca-Lario, L. Gavilan, M. Gerin, C. A. Gottlieb, P. Guillard, A. Gusdorf, P. Hartigan, J. He, E. Herbst, L. Hornekaer, C. Jäger, E. Janot-Pacheco, M. Kaufman, F. Kemper, S. Kendrew, M. S. Kirsanova, P. Klaassen, S. Kwok, A. Labiano, T. S. Y. Lai, T. J. Lee, B. Lefloch, F. Le Petit, A. Li, H. Linz, C. J. Mackie, S. C. Madden, J. Mascetti, B. A. McGuire, P. Merino, E. R. Micelotta, K. Misselt, J. A. Morse, G. Mulas, N. Neelamkodan, R. Ohsawa, A. Omont, R. Paladini, M. E. Palumbo, A. Pathak, Y. J. Pendleton, A. Petrignani, T. Pino, E. Puga, N. Rangwala, M. Rapacioli, A. Ricca, J. Roman-Duval, J. Roser, E. Roueff, G. Rouillé, F. Salama, D. A. Sales, K. Sandstrom, P. Sarre, E. Sciamma-O'Brien, K. Sellgren, S. S. Shenoy, D. Teyssier,

- R. D. Thomas, A. Togi, L. Verstraete, A. N. Witt, A. Wootten, H. Zettergren, Y. Zhang, Z. E. Zhang and J. Zhen, Astron. Astrophys., 2024, 685, A75.
- 6 J. Cernicharo, A. M. Heras, A. G. G. M. Tielens, J. R. Pardo, F. Herpin, M. Guélin and L. B. F. M. Waters, Astrophys. J., Lett., 2001, 546, L123-L126.
- 7 B. Tabone, G. Bettoni, E. F. van Dishoeck, A. M. Arabhavi, S. Grant, D. Gasman, T. Henning, I. Kamp, M. Güdel, P. O. Lagage, T. Ray, B. Vandenbussche, A. Abergel, O. Absil, I. Argyriou, D. Barrado, A. Boccaletti, J. Bouwman, A. Caratti o Garatti, V. Geers, A. M. Glauser, K. Justannont, F. Lahuis, M. Mueller, C. Nehmé, G. Olofsson, E. Pantin, S. Scheithauer, C. Waelkens, L. B. F. M. Waters, J. H. Black, V. Christiaens, R. Guadarrama, M. Morales-Calderón, H. Jang, J. Kanwar, N. Pawellek, G. Perotti, A. Perrin, D. Rodgers-Lee, M. Samland, J. Schreiber, K. Schwarz, L. Colina, G. Östlin and G. Wright, Nat. Astron., 2023, 7, 805-814.
- 8 A. M. Arabhavi, I. Kamp, T. Henning, E. F. van Dishoeck, V. Christiaens, D. Gasman, A. Perrin, M. Güdel, B. Tabone, J. Kanwar, L. B. F. M. Waters, I. Pascucci, M. Samland, G. Perotti, G. Bettoni, S. L. Grant, P. O. Lagage, T. P. Ray, B. Vandenbussche, O. Absil, I. Argyriou, D. Barrado, A. Boccaletti, J. Bouwman, A. Caratti o Garatti, A. M. Glauser, F. Lahuis, M. Mueller, G. Olofsson, E. Pantin, S. Scheithauer, M. Morales-Calderón, R. Franceschi, H. Jang, N. Pawellek, D. Rodgers-Lee, J. Schreiber, K. Schwarz, M. Temmink, M. Vlasblom, G. Wright, L. Colina and G. Östlin, Science, 2024, 384, 1086-1090.
- 9 P. Thaddeus, J. M. Vrtilek and C. A. Gottlieb, Astrophys. J., 1985, 299, L63.
- 10 S. C. Madden, W. M. Irvine, H. E. Matthews, P. Friberg and D. A. Swade, Astron. J., 1989, 97, 1403.
- 11 M. Agúndez and V. Wakelam, Chem. Rev., 2013, 113, 8710–8737.
- 12 J. Cernicharo, M. Agúndez, C. Cabezas, B. Tercero, N. Marcelino, J. R. Pardo and P. de Vicente, Astron. Astrophys., 2021, 649, L15.
- 13 B. A. McGuire, A. M. Burkhardt, S. Kalenskii, C. N. Shingledecker, A. J. Remijan, E. Herbst and M. C. McCarthy, Science, 2018, 359, 202-205.
- 14 B. A. McGuire, R. A. Loomis, A. M. Burkhardt, K. L. K. Lee, C. N. Shingledecker, S. B. Charnley, I. R. Cooke, M. A. Cordiner, E. Herbst, S. Kalenskii, M. A. Siebert, E. R. Willis, C. Xue, A. J. Remijan and M. C. McCarthy, Science, 2021, 371, 1265-1269.
- 15 M. C. McCarthy, K. L. K. Lee, R. A. Loomis, A. M. Burkhardt, C. N. Shingledecker, S. B. Charnley, M. A. Cordiner, E. Herbst, S. Kalenskii, E. R. Willis, C. Xue, A. J. Remijan and B. A. McGuire, Nat. Astron., 2021, 5, 176-180.
- 16 J. Cernicharo, M. Agúndez, R. I. Kaiser, C. Cabezas, B. Tercero, N. Marcelino, J. R. Pardo and P. de Vicente, Astron. Astrophys., 2021, 655, L1.
- 17 J. Cernicharo, R. Fuentetaja, M. Agúndez, R. I. Kaiser, C. Cabezas, N. Marcelino, B. Tercero, J. R. Pardo and P. de Vicente, Astron. Astrophys., 2022, 663, L9.
- 18 J. Cernicharo, C. Cabezas, R. Fuentetaja, M. Agúndez, B. Tercero, J. Janeiro, M. Juanes, R. I. Kaiser, Y. Endo,

- A. L. Steber, D. Pérez, C. Pérez, A. Lesarri, N. Marcelino and P. de Vicente, Astron. Astrophys., 2024, 690, L13.
- 19 G. Wenzel, T. H. Speak, P. B. Changala, R. H. J. Willis, A. M. Burkhardt, S. Zhang, E. A. Bergin, A. N. Byrne, S. B. Charnley, Z. T. P. Fried, H. Gupta, E. Herbst, M. S. Holdren, A. Lipnicky, R. A. Loomis, C. N. Shingledecker, C. Xue, A. J. Remijan, A. E. Wendlandt, M. C. McCarthy, I. R. Cooke and B. A. McGuire, Nat. Astron., 2025, 9, 262-270.
- 20 G. Wenzel, I. R. Cooke, P. B. Changala, E. A. Bergin, S. Zhang, A. M. Burkhardt, A. N. Byrne, S. B. Charnley, M. A. Cordiner, M. Duffy, Z. T. P. Fried, H. Gupta, M. S. Holdren, A. Lipnicky, R. A. Loomis, H. T. Shay, C. N. Shingledecker, M. A. Siebert, D. A. Stewart, R. H. J. Willis, C. Xue, A. J. Remijan, A. E. Wendlandt, M. C. McCarthy and B. A. McGuire, Science, 2024, 386, 810-813.
- 21 D. Cremer, E. Kraka, H. Joo, J. A. Stearns and T. S. Zwier, Phys. Chem. Chem. Phys., 2006, 8, 5304-5316.
- 22 J. Cernicharo, M. Agúndez, C. Cabezas, N. Marcelino, B. Tercero, J. R. Pardo, J. D. Gallego, F. Tercero, J. A. López-Pérez and P. de Vicente, Astron. Astrophys., 2021, 647, L2.
- 23 K. L. Kelvin Lee, R. A. Loomis, A. M. Burkhardt, I. R. Cooke, C. Xue, M. A. Siebert, C. N. Shingledecker, A. Remijan, S. B. Charnley, M. C. McCarthy and B. A. McGuire, Astrophys. J., Lett., 2021, 908, L11.
- 24 J. Wang, J. H. Marks, A. K. Eckhardt and R. I. Kaiser, J. Phys. Chem. Lett., 2024, 15, 1211-1217.
- 25 A. B. Vakhtin, D. E. Heard, I. W. Smith and S. R. Leone, Chem. Phys. Lett., 2001, 348, 21-26.
- 26 A. R. Flint and R. C. Fortenberry, Astrophys. J., 2022, 938, 15.
- 27 A. R. Flint, A. G. Watrous, B. R. Westbrook, D. J. Patel and R. C. Fortenberry, Astron. Astrophys., 2023, 671, A95.
- 28 Y. Zhao and D. G. Truhlar, J. Chem. Theory Comput., 2008, 4, 1849-1868.
- 29 Y. Zhao and D. G. Truhlar, J. Phys. Chem. A, 2004, 108, 6908-6918.
- 30 S. Grimme, J. Chem. Phys., 2006, 124, 034108.
- 31 S. Grimme, S. Ehrlich and L. Goerigk, J. Comput. Chem., 2011, 32, 1456-1465.
- 32 T. H. Dunning, K. A. Peterson and A. K. Wilson, J. Chem. Phys., 2001, 114, 9244-11.

- 33 G. Knizia, T. B. Adler and H.-J. Werner, J. Chem. Phys., 2009, 130, 054104,
- 34 K. A. Peterson, T. B. Adler and H.-J. Werner, J. Chem. Phys., 2008, 128, 084102.
- 35 P. Redondo, M. Sanz-Novo and C. Barrientos, Mon. Not. R. Astron. Soc., 2023, 527, 8659-8670.
- 36 T. J. Lee and P. R. Taylor, Int. J. Quantum Chem., 1989, 36, 199-207.
- 37 K. Fukui, Acc. Chem. Res., 1981, 14, 363-368.
- 38 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, Gaussian ~ 16 Revision C.01, Gaussian Inc., Wallingford, CT, 2016.
- 39 H. Werner, P. Knowles, G. Knizia, F. Manby, M. Schütz, P. Celani, W. Györffy, D. Kats, T. Korona, R. Lindh et al., Cardiff, UK, 2019.
- 40 K. H. Hellwege and A. M. Hellwege, Landolt-Bornstein: Group II: Atomic and Molecular Physics Volume 7: Structure Data of Free Polyatomic Molecules, Springer-Verlag, Berlin, 1976.
- 41 K. P. Huber and G. Herzberg, Molecular Spectra and Molecular Structure. IV. Constants of Diatomic Molecules, Van Nostrand Reinhold Co., 1979.
- 42 K. Kuchitsu, Structure of Free Polyatomic Molecules Basic Data, Springer, Berlin, 1998.
- 43 S. N. Milam, K. Yocum, L. Mora, E. Todd, P. Gerakines and S. Widicus Weaver, American Astronomical Society Meeting Abstracts #235, 2020, p. 418.01.