**An Unconventional Dark Radical Chemistry in Dense Molecular Cloud: Directed Gas-Phase Formation of Naphthyl Radicals**

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**ABSTRACT**

The synthetic pathways to aromatic molecules inside photon shielded dense molecular clouds remain a fundamental, unsolved enigma in astrochemistry and astrophysics with low-temperature molecular growth routes involving aromatic radicals, like prototype bicyclic naphthyl (C10H7•), implicated as key sources. Here, exploiting crossed molecular beam experiments augmented by electronic structure calculations, unexpected pathways are exposed leading to the gas phase formation of 1- and 2-naphthyl via barrierless bimolecular reactions of atomic carbon (C) with indene (C9H8) and of dicarbon (C2) with styrene (C8H8) accompanied by ring expansion and cyclization together with aromatization. These facile routes challenge conventional wisdom that aromatic radicals are formed in deep space solely via ‘bright’ gas phase photochemistry of their closed shell polycyclic aromatic hydrocarbon (PAH) precursors. A hitherto disregarded ‘dark’ aromatic radical chemistry with aromatic radicals synthesized via gas-phase reactions offers new concepts on the chemical evolution of the chemistry of dark molecular clouds eventually culminating in the rapid formation of aromatics, fullerenes, and carbonaceous nanostructures.

**1. INTRODUCTION**

     The ubiquitous presence of polycyclic aromatic hydrocarbons (PAHs) – organic molecules composed of fused benzene rings – along with their derivatives such as hydrogenated, ionized, and substituted counterparts in the interstellar medium and in carbonaceous chondrites like Orgueil, Allende, and Murchison has remained a key paradox in our Universe considering that interstellar PAHs are faster destroyed than synthesized.[1-6](#_ENREF_1) Accounting for up to 20% of the galactic carbon budget, rapid and barrierless bimolecular reactions commencing with prototype mono- (phenyl, C6H5•)[7](#_ENREF_7) and bicyclic (1-/2-naphthyl, C10H7•) aromatic radicals[8](#_ENREF_8) signify critical molecular mass growth processes to aromatic molecules in cold molecular clouds at temperatures as low as 10 K thus affording a sustainable solution to this puzzle. In particular, naphthalene (C10H8) and anthracene/phenanthrene (C14H10) can be prepared in the gas phase through the reactions of phenyl and 1-/2-naphthyl radicals, respectively, with vinylacetylene (C4H4) via the Hydrogen Abstraction−Vinylacetylene Addition (HAVA) mechanism.[7-9](#_ENREF_7) Except for the key role in forming acenes, helicenes, and phenacenes with fused benzene rings,[8](#_ENREF_8),[10](#_ENREF_10) naphthyl radicals (C10H7•) have also emerged as fundamental precursors to acenaphthylene (C12H8),[11](#_ENREF_11) fluorene (C13H10),[12](#_ENREF_12) and fluoranthene (C16H10)[13](#_ENREF_13) carrying five-membered rings, which represent critical molecular building blocks to non-planar PAHs like corannulene and even fullerenes (Figure S1).[12](#_ENREF_12),[13](#_ENREF_13) This classifies naphthyl radicals as central reaction intermediates in molecular mass growth processes to two- and three-dimensional carbonaceous nanostructures in deep space. Recently, mechanisms on distinct carbonaceous particle formation stages, especially the soot inception stage, were also explored.[14-17](#_ENREF_14)

     The recent detection of PAHs carrying up to four six-membered rings in the returned samples from the carbonaceous asteroids Ryugu[18](#_ENREF_18),[19](#_ENREF_19) and Bennu[20](#_ENREF_20),[21](#_ENREF_21) within the framework of the OSIRIS-REx mission suggest a hitherto unidentified low-temperature reservoir of aromatic molecules, which cannot be supported by the previously contemplated classical photolysis driven formation of reactive aromatic radicals like phenyl (C6H5•) and naphthyl (C10H7•) within cold molecular clouds. However, which mechanisms can be proposed to account for the formation of interstellar naphthyl radicals? While ultraviolet (UV) photolysis of naphthalene (C10H8) preferentially leads to atomic hydrogen loss forming naphthyl radicals (C10H7•),[2](#_ENREF_2) UV photons from the ambient interstellar radiation cannot reach the dense, inner regions of cold molecular clouds with the strength of photon field reduced at least by a factor of 103.[22-24](#_ENREF_22) A cosmic ray induced internal UV field present even deep inside cold molecular clouds could be too weak to drive an efficient photodissociation network and sufficient fractional abundances of polycyclic aromatic radicals such as naphthyl radicals (C10H7•) as the prototype representative.[25](#_ENREF_25),[26](#_ENREF_26) Therefore, ‘bright’ mechanisms leading to PAH radicals via photolysis of closed-shell PAHs have not been able to fully account for a plethora of aromatics detected on Ryugu[18](#_ENREF_18),[19](#_ENREF_19) and Bennu.[20](#_ENREF_20),[21](#_ENREF_21) Low-temperature bimolecular gas-phase routes to naphthyl radicals (C10H7•) and to other PAH radicals from astrochemically relevant reactants in general may afford an elegant and versatile ‘dark’ reservoir of PAH radicals such as naphthyl (C10H7•) *without* the need of photon-initiated molecular mass growth processes even in the darkest and coldest molecular clouds.[27](#_ENREF_27)

     The molecular beam technique represents a unique tool in understanding the chemical reactions and the underlying mechanisms on the molecular level. In particular, the crossed molecular beam (CMB) method coupled with universal mass spectrometric detection of the products allows us to unravel the formation routes to complex molecules including transient species in combustion flames and low-temperature environments of cold molecular clouds. Here, utilizing a universal CMB machine, we report the very first directed gas-phase synthesis of 1- and 2-naphthyl radicals (C10H7•, X2A') via the bimolecular reactions of ground state atomic carbon (C, 3P) with indene (C9H8, X1A') (reaction (1)) and of dicarbon molecule (C2, X1Σg+/a3Πu) with styrene (C6H5C2H3, X1A') (reaction (2)) under single collision conditions. Carbon and dicarbon beams are produced via laser ablation of a rotating graphite rod and seeding the ablated species in noble gases, whereas Krypton-seeded styrene and indene molecular beams are generated in the secondary source (Figure S2-3, Supporting Information). As demonstrated by the electronic structure calculations, both reactions appeared to be barrierless and exoergic thus providing facile routes to interstellar naphthyl radicals via two distinct pathways involving ring expansion (reaction (1)) and cyclization along with aromatization (reaction (2)) even in the darkest regions of molecular clouds from readily available organic reactants in these C1+C9 (carbon – indene) and C2+C8 (dicarbon – styrene) systems at 10 K. These findings challenge conventional wisdom that molecular mass growth processes in molecular clouds must be initiated via photolysis of closed shell gas phase PAHs through a ‘bright’ chemistry thus affording a viable alternative of a ‘dark’ chemistry initiated through bimolecular reactions leading to aryl radicals, fundamentally enhancing our understanding of the chemical evolution of the aromatic Universe we live in.

(1) C (12 amu) + C9H8 (116 amu) → C10H7• (127 amu) + H (1 amu)

(2) C2 (24 amu) + C8H8 (104 amu) → C10H7• (127 amu) + H (1 amu)

**2. RESULTS**

**2.1 Carbon-Indene and Dicarbon-Styrene Systems − Laboratory Frame**

      The reactive scattering experiments (1) and (2) were carried out in the gas phase under single collision conditions (Methods) (Figure 1). In the atomic carbon (C, 3P) **-** indene (C9H8, X1A') system, reactive scattering signal was observed at mass-to-charge (*m/z*) ratios of 127 (C10H7+/13CC9H6+) and 126 (C10H6+/13CC9H5+). Compared to *m/z* = 127, the relative intensity of the TOF spectra recorded at *m/z* = 126 was only 0.38 ± 0.03 with TOF spectra at both *m/z* values indistinguishable after scaling. These findings hint that signal at *m/z* = 127 originated from the formation of C10H7• radicals (127 amu) along with atomic hydrogen (1 amu) with ion counts at *m/z* = 126 associated with dissociative ionization of the parent species at *m/z* = 127. Considering the enhanced signal-to-noise (S/N) ratio at m/z = 127 compared to *m/z* = 126, TOF spectra were recorded at *m/z* = 127 from 30° to 65° in 5° intervals. These TOF spectra were then integrated to obtain the laboratory angular distribution (LAD) (Figure 1A and B). The LAD is essentially forward-backward symmetric and displays a maximum around the CM angle; this finding implies indirect scattering dynamics[28](#_ENREF_28) and the formation of long-lived C10H8 complex(es)[29](#_ENREF_29) ultimately decomposing via hydrogen elimination to C10H7• product(s). Therefore, our experimental data alone reveal the atomic carbon versus atomic hydrogen exchange pathway resulting in the gas phase preparation of C10H7• isomer(s) in the carbon **-** indene system.

     In case of the dicarbon (C2, X1Σg+/a3Πu) **-** styrene (C8H8, X1A') system (reaction (2)), reactive scattering signal was also detected at *m/z* = 127 and 126. In analogy to reaction (1), TOF data at *m/z* = 127 and 126 overlapped after scaling. Considering the better signal-to-noise (S/N) ratio at *m/z* = 127 as compared to *m/z* = 126, with relative intensities of 1.00 : 0.62 ± 0.03, TOF data were collected at *m/z* = 127 over a range of 50° in 5° intervals. Once again, the forward-backward symmetric LAD along with the detected mass-to-charge ratios alone provide evidence on the gas

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**Figure 1**. LAD (A, B), and TOFs (C, D) for the reactions of atomic carbon with indene (A, C) and dicarbon molecule with styrene (B, D). The solid circles with their error bars represent the normalized experimental distribution; the open circles indicate the experimental data. The red lines represent the best fits obtained. Atoms are color-coded in gray (carbon) and white (hydrogen).

phase formation of C10H7• (127 amu) isomer(s) along with atomic hydrogen (1 amu) via reaction (2) involving long-lived C10H8 intermediates (indirect scattering dynamics). Additional information on the position where the hydrogen loss originates from, i.e., the aromatic ring and/or the vinyl moiety, can be obtained by conducting experiments of dicarbon with D3-(C6H5C2D3) and D5-styrene (C6D5C2H3). For the C2**-**C6H5C2D3 system, signal was probed at *m/z* = 130 (C10H4D3+) and 129 (C10H3D3+/C10H5D2+) with the determined ratio of 1 : 1.22 ± 0.03 taking into account the contribution of natural 13C isotopes and differences of the spread recoil circles of the atomic hydrogen versus deuterium loss. For the C2**-**C6D5C2H3 system, a ratio of 1 (*m/z* = 132) : 0.78 ± 0.02 (*m/z* = 131) is obtained. These results expose that the atomic hydrogen loss can originate from both the vinyl moiety *and* the phenyl group (Supplementary Materials).

**2.2 Carbon-Indene and Dicarbon-Styrene Systems − Center-of-Mass Frame**

     The compelling identification of C10H7• isomer(s) in both the carbon**-**indene and dicarbon**-**styrene systems afford an elucidation of the nature of the isomer(s) along with the underlying mechanisms. This is achieved by converting the laboratory data (TOFs, LAD) into the center-of-mass reference frame yielding the translational energy (*P(E*T*)*) and angular flux (*T(θ)*) distributions (Figure 2).[30](#_ENREF_30) *First*, for both systems, best fits of the laboratory data are derived via a single atomic hydrogen loss channel involving reaction (1) and reaction (2). For the carbon**-**indene system (Figure 2A and B), the *P(E*T*)* function is characterized by a distribution maximum of 101 ± 9 kJ mol-1 suggesting a tight exit transition state upon the unimolecular decomposition of the C10H8 complex(es). Further, a high-energy cutoff of Emax = 328 ± 33 kJ mol-1, which represents the sum of the reaction exoergicity plus the collision energy (35.8 ± 1.2 kJ mol-1), is extracted for those products formed without internal excitation. This results in a reaction exoergicity of 293 ± 34 kJ mol-1. For the dicarbon**-**styrene reaction (Figure 2D and E), a distribution maximum of the *P(E*T*)* was also observed here at 66 ± 4 kJ mol-1, which also indicates a tight exit transition state a involving significant reorganization of the electron density upon decomposition of the C10H8 intermediates to the separated products. A *E*max of 384 ± 23 kJ mol-1 is revealed; considering the collision energy of 20.6 ± 0.5 kJ mol-1, reaction energies of -363 ± 23 kJ mol-1 and -371 ± 23 kJ mol-1 for dicarbon reactants in their X1Σg+ ground state and a3Πu excited state, respectively, are extracted. Fractions of 39 ± 7 % and of 34 ± 4 % of the available energy channeling into the translation degrees of freedom of the products are observed for reactions (1) and (2), respectively,



**Figure 2**. CM translational energy (a), angular flux distributions (b) and the flux contour map (c). The solid red lines and shaded areas define the best fit and the error limits, respectively.

also supporting indirect scattering dynamics.[29](#_ENREF_29) *Second*, the *T(θ)* functions exhibit a forward-backward symmetric pattern and peak intensities at 90° (sideway scattering). These findings reveal indirect dynamics involving the formation of the C10H8 intermediates holding a lifetime that is comparable to or exceeding their rotational period.[31](#_ENREF_31) The sideways scattering of both reactions exposes geometrical constraints inferring a hydrogen atom emission nearly parallel to the total angular momentum vector, i.e. perpendicularly to the rotational plane of the decomposing complex. These findings are also reflected in the flux contour maps, which contain the overall information on the sideway scattering process (Figure 2C and F).

**3. DISCUSSION**

     With the detection of C10H7• isomer(s) under single collision conditions in both the carbon**-**indene and dicarbon**-**styrene systems, we are now adjoining these results with electronic structure calculations carried out at an accuracy of ±10 kJ mol-1 to unravel the nature of the isomers formed and to elucidate the underlying reaction mechanisms (Figures 3-6; Supporting Information). Statistical calculations exploiting Rice−Ramsperger−Kassel−Marcus (RRKM) theory to predict unimolecular reaction rate constants and product branching ratios were also carried out (Supporting Information).

**3.1 Carbon-Indene System**

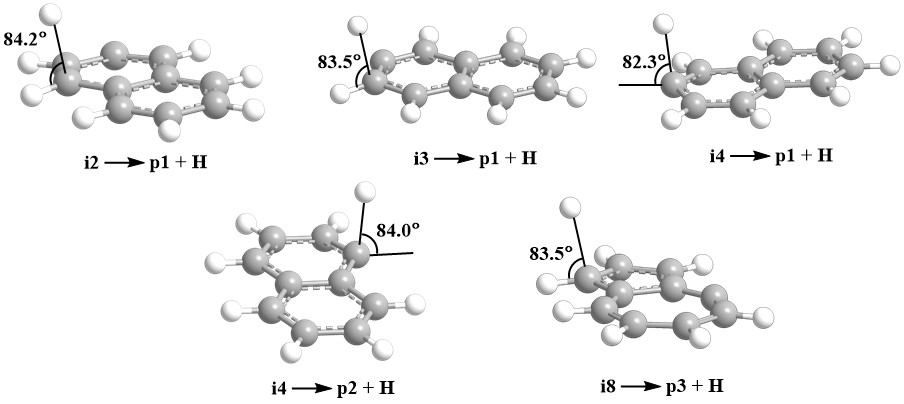
     For the carbon – indene system, the calculations were conducted on the triplet surface revealing three product channels (**p1**-**p3**) (Figure 3). A comparison of the experimentally determined reaction energy of -293 ± 34 kJmol-1 with the calculated reaction exoergicities suggests that at least the thermodynamically most stable 2-/1-naphthyl radicals (**p1/p2**; X2A'; ΔrG = -252/251 ± 10 kJmol-1) are formed. Formation of the 4-azulenyl isomer (**p3**; ΔrG = -123 ± 10 kJmol-1) carrying a fused five- and seven-membered ring might be masked in the low-energy section of the *P*(*E*T) and cannot be excluded at the present stage. Note that the computed reaction energies leading to 1- and 2-naphthyl of -251 ± 10 and -252 ± 10 kJmol-1 agree well with the predicted experimental enthalpies of formation from Active Thermochemical Tables (ATcT)[32](#_ENREF_32) of -261 ± 2 and -262 ± 2 kJmol-1, respectively. These results combined with the reaction energies obtained from experimental value of -293 ± 34 kJmol-1 support the identification of the 1-/2-naphthyl in the carbon-indene system.



**Figure 3**. Triplet potential energy surface (PES) for the reaction of atomic carbon (C, 3P) with indene (C9H8). The pathway highlighted in blue represent the dominant routes to naphthyl radicals.

     Which are the dominating reaction mechanisms for the gas phase preparation of the 1-/2-naphthyl radicals plus atomic hydrogen? The bimolecular reaction of ground state triplet atomic carbon with indene is initiated though barrierless additions to the π electronic system of the five-membered ring or the benzene moiety leading to intermediates **i1**, **i5**, or **i7**, which are stabilized by 215, 67, or 66 kJ mol-1 with respect to the separated reactants, respectively. Intermediates **i5** and **i7** are connected via **i6** – a tricyclic structure – through low lying transitions states ranging well below the energy of the separated reactants. Collision complex **i1** can undergoring opening of the annulated three-membered circle to triplet **i2** thus forming a naphthalene carbon skeleton. The collision complexes **i5** and **i1** interconnect through a barrier of only 8 kJ mol-1; this pathway should dominate compared to the **i5** → **i6** isomerization, which requires a barrier of 24 kJ mol-1 to be overcome. Once accessed, **i2** can undergo a facile unimolecular decomposition via atomic hydrogen elimination from the CH2 moiety to **p1** or isomerizes via hydrogen shift from the CH2 moiety to the neighboring CH group to **i3** followed by consecutive isomerization to triplet **i4** (naphthalene). Intermediates **i3** and **i4** can decompose to **p1** and/or **p2** plus atomic hydrogen via three exit channels: **i3** → **p1** + H, **i4** → **p1** + H, and **i4** → **p2** + H with exit transition states located 19, 28, and 24 kJ mol-1 above the energy of the separated products. It should be noted that addition pathways of atomic carbon across distinct chemically inequivalent carbon-carbon bonds of the benzene moiety were also located (Supporting Information). These processes lead via ring opening and atomic hydrogen loss to **p4** and **p5** carrying an adjacent five- and seven-membered ring with the radical center located at the seven-membered ring structure (Figure S4). The energetics of these reaction sequences (**i9** → **i10** → **p4** + H, **i11** → **i12** → **p5** + H) exhibit similar behaviors comparing to **i7** → **i8** → **p3** + H. Overall, the electronic structure calculations revealed barrierless pathways to the 1- and 2-naphthyl radicals (X2A') under single collision conditions in the atomic carbon – indene system via indirect scattering dynamics with **p1** (2-naphthyl) accessible via decomposition of intermediates **i2, i3**, and **i4** and **p2** (1-naphthyl) exclusively from **i4**.

     Which of these isomers dominates? Statistical RRKM (Rice–Ramsperger–Kassel–Marcus) theory is utilized to predict the branching ratios of C10H7• isomer (**p1**-**p3**). These calculations are conducted for three different scenarios commencing exclusively with initial intermediates of **i1**, **i5**, or **i7** within the limit of a complete energy randomization. The ultimate branching ratios depend strongly on the initial collision complex. At the experimental collision energy of 36 kJ mol-1, **i7**



**Figure 4.** Exit transition states lead to **p1**-**p3**. The angle for each emitting hydrogen atom is given with respect to the rotational plane of the decomposing complex.

decomposes predominantly to **p3** with a small fraction of 0.16 % dissociating to **p1** (Table S2). Those trajectories in which the bimolecular collision between atomic carbon and indene lead to **i1** and **i5**, **p1** will bethe highly prevailing product (99.99 and 99.84 %, respectively) formed under our experimental condition. Furthermore, the computed geometries of the exit transition states leading to **p1-p3** plus H (**i2** → **p1** + H, **i3** → **p1** + H, **i4** → **p1** + H, **i4** → **p2** + H, and **i8** → **p3** + H) from **i2**, **i3**, **i4** and **i8** reveal that the atomic hydrogen is emitted nearly perpendicular to the plane of the rotating complex at angles from 82.3° to 84.2° (Figure 4), which is consistent with the sideway scattering fitting of the *T*(*θ*) distributions.

    In conclusion, the formation of the 2-naphthyl radical (**p1**) proceeds via initial collision complexes **i1**, **i5**, and **i6** with the latter two complexes eventually isomerizing to **i1.** This intermediate ring opens to **i2**, which contains the naphthalene carbon skeleton followed by atomic hydrogen loss and aromatization from the methylene moiety to the 2-naphthyl radical (Figure 3).

**3.2 Dicarbon-Styrene System**

    For the dicarbon – styrene system, both the singlet and triplet C10H8 surfaces were explored (Figure 5A and B). It should be noted that the reaction energies of -363 ± 23 kJ mol-1 and -371 ± 23 kJ mol-1 for dicarbon reactants in their X1Σg+ ground state and a3Πu excited state, respectively, extracted from experiment closely match the computed values for the formation of 1-/2-naphthyl, -349/-350 ± 10 kJ mol-1 for singlet and -359/-361 ± 10 kJ mol-1 for triplet, and the reaction energies derived from the enthalpies of formation from ATcT (-356/-357 and -363/-364 ± 2 kJ mol-1), respectively.[32](#_ENREF_32) On the singlet surface, the reaction is initiated via barrierless additions of the dicarbon molecule to the π electron density of the benzene or vinyl moiety accessing collision complexes **i3'** and/or **i1'** (benzene ring) and/or **i2'** (vinyl group). All collision complexes can isomerize via ring closure to the bicyclic intermediate **i4'**. An exotic transition state involving a hydrogen migration and ring closure connects **i1'** to a tricyclic intermediate **i7'** through a significant barrier of 197 kJ mol-1; ring opening restores the benzene moiety and a hydrogenated four-carbon side chain (**i9'**). A hydrogen migration from the benzene moiety to the side chain accompanied by hydrogen migration accesses the naphthalene intermediate (**i15'**) which represents the global minimum of the C10H8 potential energy surface. Considering the collision complex **i2'**, a hydrogen shift from the CH2 group to the terminal carbon of the side chain leads to phenylvinylacetylene (**i8'**), which can undergo ring closure to **i12'**. Alternatively, a hydrogen migration in **i3'** connects to ortho-vinylethynyl benzene (**i6'**), which can subsequently ring closure to a bicyclic carbene intermediate (**i13'**). Inspecting **i4'**, its isomerization via ring contraction provides the three-membered ring intermediate **i11'**, which can undergo ring opening to **i15'**. Second, **i4'** is connected to **i5'** via ring expansion. This opens three routes: **i**) an energetically unfavorable rearrangement via a significant barrier of 240 kJ mol-1 accessing the azulene intermediate **i14'** followed by H elimination from the five- or seven-membered ring to **p6** or **p5** through loose transition states; **ii**) a ring contraction to the tricyclic intermediate **i10'**, which can subsequently isomerize to **i15'** via ring opening of the annulated three-membered cycle; **iii**) molecular hydrogen atom loss from the CH2 group and the bridging carbon to **p7**. These intermediates can undergo multiple unimolecular decomposition pathways via atomic hydrogen loss. The 2-naphthyl radical (**p1**) can be accessed from **i15'** and **i12'** via a loose and tight exit transition states, respectively; the 1-naphthyl radical (**p2**) could be generated from **i15'** and **i13'** also via loose and tight exit transition states, respectively.

     On the triplet surface, the dicarbon molecule can add barrierlessly to the terminal carbon atom of the side chain and the benzene moiety of the styrene forming initial intermediates **i16'** and **i17'** lying 198 and 125 kJ mol-1 below the separated reactants, respectively. Subsequent isomerization of the initial intermediates involves ring closure leads to **i18'** which opens three new routes: **i**) unimolecular decomposition via H elimination from the bridging carbon to **p8**; **ii**) H migration from the bridging carbon to the neighboring position forming **i20'** which can lose an atomic hydrogen from the CH group via a loose transition state to **p8** or from the CH2 group to **p1** through a tight transition state located 20 kJ mol-1 above the separated products; **iii**) H migration from the

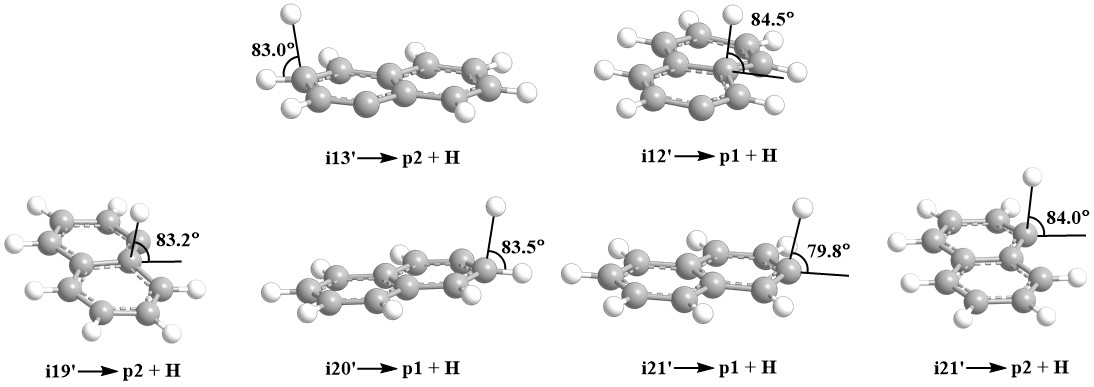




**Figure 5.** Singlet (A) and triplet (B) PES for the reaction of dicarbon (C2, X1Σg+/a3Πu) with styrene (C8H8, X1A'). The pathways highlighted in blue represent the dominant routes to naphthyl radicals.

CH2 group to the neighboring position leading to **i19'** which can undergo further H migration to **i21'** (naphthalene), the most stable intermediate on the triplet PES, or directly access **p2** with atomic hydrogen elimination originating from the bridging carbon. Once formed a 2,3-hydrogen migration in **i21'** connects it to **i20'** via a high barrier lying 164 kJ mol-1 above **i20'**, and the decomposition of **i21'** forms **p1** and **p2** via tight exit transition states lying 29 and 24 kJ mol-1 above the separated products. Note that the reaction of dicarbon with benzene was previously studied[33](#_ENREF_33) revealing the formation of the phenylethynyl radical (C6H5CC, X2A') on both singlet and triplet surfaces. According to the calculations, on the singlet surface, dicarbon (C2, X1Σg+) added to a carbon-carbon double bond leading to the methylidynecyclopropyl benzene followed by non-RRKM unimolecular decomposition to phenylethynyl (C6H5CC) plus atomic hydrogen. Recall that in the dicarbon-styrene reaction on the singlet surface, the similar addition mechanism leads to **i1'**. However, the existence of the vinyl moiety in **i1'** results in the distinct reaction dynamics, i.e., **i1'** is identified to undergo ring closure to bicyclic **i4'** or tricyclic **i7'** instead of the formation of ethynylstyryl (C2H3C6H4CC). Similarly, on the triplet surface, the addition of dicarbon (C2, a3Πu) to one carbon of the benzene molecule (C2**-**C6H6) or the benzene moiety of styrene (C2**-**C8H8) forms ethynylbenzene (C6H6CC) or ethynylstyrene (C8H8CC, **i17'**), respectively. Here, the influence of the vinyl moiety in styrene is also depicted in the subsequent ring closure to a bicyclic **i18'** isomerrather than the simple H elimination to ethynylstyryl (C2H3C6H4CC).

     It is interesting to compare the RRKM branching ratios for the C2**-**C8H8 system on the singlet surface with those on the triplet surface (Tables S3-4). On the singlet surface, the branching ratios depend strongly on the initial collision complex(es). RRKM calculations reveal that **p1** will be the dominant product if **i1'** and **i2'** are formed (67.3% for **i1'** and 94.0% for **i2'**), 32.3% of **i1'** and5.9% of **i2'** also decompose to **p2**; and **p2** is almost the only product from **i3'** (99.6%). On the triplet surface, since both **i16'** and **i17'** will isomerize to **i18'**, the formation of **i16'** and **i17'** lead to the same ratios of the final products: 97.7% of them decomposes to **p1** with the remaining 2.3% connecting to **p2**. Thus, on both singlet and triplet surfaces, our calculations predict the almost exclusive synthesis of naphthyl radicals (**p1** and/or **p2**) which is in line with the experiments.These findings arefully supported by the geometry calculations of the exit transition states leading to naphthyl radicals on both singlet (**i13'** and **i12'**) and triplet (**i19', i20'** and **i21'**) surfaces which reveal that the hydrogen atoms are emitted at angles from 80° to 85° relative to the rotation plane of the complexes (Figure 6).

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**Figure 6.** Exit transition states lead to 2-naphthyl (**p1**) and 1-naphthyl (**p2**) radicals. The angle for each emitting hydrogen atom is given with respect to the rotational plane of the decomposing complex.

     Last,the experimental findings of the isotopic substitution studies can be addressed (Figure S2). For the reaction of singlet dicarbon with D3-styrene (C6H5C2D3, hydrogen and deuterium atoms are color-coded in gray and light blue, respectively, in Figure S2), the formation of **i13'** will only lead to the atomic deuterium loss exit channel (from vinyl) to **p2**, whereas the formation of **i12'** leads to **p1** + H. Moreover, **i15'** can eliminate atomic hydrogen (benzene ring) and atomic deuterium (vinyl moiety) to **p1** or **p2** with a ratio of 1:1 or 3:1, respectively. For the reaction of singlet dicarbon with D5-styrene (C6D5C2H3, hydrogen and deuterium atoms are color coded in light blue and gray, respectively, in Figure S2), the formation of **i13'** will result in atomic hydrogen elimination from vinyl group to **p2**, **i12'** will undergo deuterium loss to **p1**;and the formation of **i15'** will lead to both atomic hydrogen and deuterium loss to **p1** or **p2** with H:D ratios of 1:1 and 1:3, respectively. In the triplet dicarbon **-** D3/D5-styrene (C6H5C2D3/C6D5C2H3) reactions (Figure S2B), it is clear that the formation of **i19'** will lead to **p2**+H in the C2**-**D3-styrene system and **p2**+D in the C2**-**D5-styrene system, whereas **i20'** or **i21'** can lose a deuterium atom and/or a hydrogen atom to naphthyl radical (**i20'→p1**+H/D, **i21'→p1/p2**+H/D). We can then conclude that atomic hydrogen can be emitted from the vinyl and/or phenyl groups on both singlet and triplet surfaces of the C2**-**C8H8 system, in excellent agreement with our experimental predictions.

     Overall, on the singlet surface of the C2**-**C8H8 reaction, four pathways of **i8'→i12'→p1**+H, **i4'→i11'→i15'→p1**+H, **i4'→i11'→i15'→p2**+H, and **i6'→i13'→p2**+H can be important in forming 1/2-naphthyl radicals (Figure 5A); their formation involves distinct collision complexes **i1'**-**i3'**. The triplet surface is simpler, and the formation of **i18'** is centralfor accessing the dominate reaction pathway **i18'→i20'→p1**+H with **i18'** connected to both initial intermediates of **i16'** and **i17'** (Figure 5B).

**3.3 Astrochemical modeling**

     The identification of the barrierless formation of 1/2-naphthyl radicals (C10H7) through two neutral-neutral reactions provides an opportunity of exploring the aromatic chemistrydeep inside the photon shielded dense molecular clouds. To investigate the implications of our laboratory results, we explore an enhanced neutral-neutral chemical reaction network, which incorporates the newly identified reactions of ground state carbon (C) with indene (C9H8) and of dicarbon (C2) with styrene (C8H8) and by excluding photodissociation. The details of the astrochemical modeling based on the UMIST (University of Manchester Institute for Science and Technology) database has been present before.[34-36](#_ENREF_34)



**Figure 7**. The fractional abundances of the styrene (C8H8, orange), indene (C9H8, red), naphthyl (C10H7, blue), and anthracene/phenanthrene (C14H10, magenta) are plotted as a function of time. Astronomically observed fractional abundances along with the uncertainties of indene are visualized by the horizontal bars.

     As depicted in the Figure 7, the modeling studies reveal that even in the darkest cold molecular cloud, the naphthyl radicals (C10H7) can be efficiently formed with predicted peak fractional abundances of 1.8 × 10-10 with respect to molecular hydrogen (H2) at 6.3 × 105 years. Reactions 1 (C + C9H8 → C10H7 + H) and 2 (C2 + C8H8 → C10H7 + H) contribute 96 and 4%, respectively. Furthermore, larger PAHs of anthracene/phenanthrene (C14H10) are suggested to be formed through reaction with vinylacetylene (C4H4)[8](#_ENREF_8) with peak fractional abundance of at least 4.0 × 10-11 with respect to molecular hydrogen. Considering that o-C6H4 (benzyne) has been detected recently with fractional abundances of (5.0 ± 1.0) × 10-11 in the cold molecular cloud TMC-1 via the Yebes 40m radio telescope,[37](#_ENREF_37) the predicted fractional abundances here suggest that the naphthyl radicals formed in photon shielded dense molecular clouds are a promising candidate for future astronomical observations.

**4. CONCLUSIONS.**

     Our combined crossed molecular beam experiments and electronic structure calculations exposed a novel concept on the low temperature gas phase synthesis of naphthyl radicals (C10H7•) – the simplest prototype of polycyclic aromatic hydrocarbon radical – inside photon shielded cold molecular clouds such as the Taurus Molecular Cloud (TMC-1). The formation of naphthyl radicals is tentatively studied before, In the laboratory, naphthyl radical can been prepared in single-crystal naphthalene matrix undergoing radioactive decay,[38](#_ENREF_38),[39](#_ENREF_39) or via photodissociation of di-1-naphthoylperoxide (DNPO) or thermolysis of 1,2-diethynylbenzene in solution.[40](#_ENREF_40),[41](#_ENREF_41) In flow reactor, naphthyl radicals might be formed as minor products from pyrolysis of ethynylbenzene in helium.[42](#_ENREF_42) In combustion conditions, HACA mechanism predicts that 1-naphthyl radical might be produced with a barrier of around 37 kJ mol-1, and 1- and 2-naphthyl radicals are mainly formed via the H-atom abstraction of naphthalene.[9](#_ENREF_9),[43](#_ENREF_43),[44](#_ENREF_44) Furthermore, the calculation also predict that naphthyl can be formed from phenylacetylene via the ethynyl addition mechanism (EAM).[45](#_ENREF_45) However, the barrierless gas phase pathways to naphthyl have been still elusive. Here, the barrierless bimolecular reactions of atomic carbon (C) with indene (C9H8) and of dicarbon molecule (C2) with styrene (C8H8) are driven by ring expansion and cyclization, respectively, together with aromatization to eventually form 1- and 2-naphthyl radicals in single collision events of bimolecular reactions involving readily available reactants. The reactants include indene identified as the most abundant aromatic and organic ring molecule to date in TMC-1 with fractional abundance of (0.8-1.4) × 10-9 relative to H2 and styrene predicted with the upper limit of fractional abundance of 10-8 in TMC-1.[46](#_ENREF_46),[47](#_ENREF_47) Considering that the hydrogen atom of the aromatic ring of the styrene and indene reactants can be replaced with any organic group, these bimolecular gas phase reactions represent a universal template toward the formation of the substituted naphthyl radicals. These facile routes challenge conventional wisdom that aromatic radicals are formed in deep space solely via ‘bright’ gas phase photochemistry of their closed shell polycyclic aromatic hydrocarbon (PAH) precursors. A hitherto disregarded ‘dark’ aromatic radical chemistry with aromatic radicals synthesized in the gas phase via chemical reactions rather than a ‘bright’ photochemistry offers a new concept on the chemical evolution of the chemistry of dark molecular clouds thus facilitating a better understanding of the origin and chemical evolution of carbon in our aromatic Universe.

**5. EXPERIMENTAL AND COMPUTATIONAL**

**5.1. EXPERIMENTAL METHODS − CARBON-INDENE SYSTEM:** The gas-phase bimolecular reaction of ground state carbon (C, 3P) with indene (C9H8, X1A') was investigated under single-collision conditions employing the crossed molecular beam (CMB) scattering technique with mass spectrometric detection in the time-of-flight (TOF) mode.[48](#_ENREF_48),[49](#_ENREF_49) Briefly, the supersonic beam of ground state carbon atoms was prepared using a laser ablation source by ablating a rotating graphite rod with a Nd:YAG laser (266 nm, 10-12 mJ pulse-1, 30 Hz) and seeding the ablated carbon atoms in helium carrier (4 atm) released by a piezoelectric valve at 60 Hz. The atomic carbon beam was skimmed and velocity selected by a chopper wheel operating at 120 Hz. Peak velocity and speed ratio of the chopped section of the primary beam were 2538 ± 46 m s-1 and 2.8 ± 0.3, respectively. In this manner carbon atoms are produced only in the ground 3P state. The indene (TCI, >99%) reactant was held in a bubbler and seeded in krypton (550 Torr, Praxair, 99.999%) and the supersonic beam was released by the second piezoelectric valve with the peak velocity and speed ratio of 367 ± 6 m s-1 and 15.2 ± 1.4, respectively. Two beams of the reactants are crossed at 90° in the interaction region under single-collision condtions leading to a collision energy of 35.8 ± 1.2 kJ mol-1 and a center-of-mass angle of 54.4 ± 0.5°. Experimental conditions were optimized to make sure that the primary beam contains only a small amount of dicarbon (<5% compared to atomic carbon).

**5.2. EXPERIMENTAL METHODS − DICARBON-STYRENE SYSTEM:** Dicarbon beam [C2 (X1Σg+/a3Πu)] was generated by the same ablation source described above. The 266 nm laser output at energy of 9-11 mJ pulse-1 and 4 atm Neon carrier (Ne, 99.9999%) was used. Experimental conditions and time sequences were optimized to maximize dicarbon concentrations in the supersonic beam. The ro-vibrational distribution of both singlet ground state (X1Σg+) and the lowest lying triplet state (a3Πu) of dicarbon beam was characterized by laser-induced fluorescence (LIF) [50](#_ENREF_50). In detail, the X1Σg+ state was probed via the Mulliken excitation (D1Σu+-X1Σg+) and the bimodal rotational distributions of both ν = 0, 1 were confirmed at fractions of 0.83 ± 0.10 (200 K: 0.44 ± 0.05; 1000 K: 0.39 ± 0.05) and 0.17 ± 0.04 (200 K: 0.06 ± 0.02; 1000 K: 0.11 ± 0.02). The rotational temperature (Trot) for the first and second vibrational levels of the a3Πu state were detected to be 240 ± 30 and 190 ± 30 K via the Swan band transition (d3Πg-a3Πu) with fractions of 0.67 ± 0.05 and 0.33 ± 0.05, respectively.[50](#_ENREF_50) A stainless-steel bubbler was used to hold the styrene reactant and krypton carrier introduced at 550 Torr during the reaction. The corresponding peak velocity of 372 ± 6 m s-1 and speed ratio of 12.8 ± 0.6 of the styrene beam were determined, resulting in a collision energy of 20.6 ± 0.5 kJ mol-1 and a center-of-mass angle of 48.9 ± 0.4° in the C2-C8H8 reaction (Table S1). In addition, the D3- and D5-styrene (CDN Isotopes, 99% D), i.e., C6H5C2D3 and C6D5C2H3, were employed to confirm the position of the hydrogen atom elimination.

     For both reactions, the neutral scattered products entering the triply differentially pumped, ultrahigh vacuum (6.6×10-12 Torr) detector were ionized via an electron impact ionizer and the laboratory data were mass-selected via a quadrupole mass spectrometer operating in the TOF mode. The details of our detector have been depicted elsewhere.[51](#_ENREF_51) In brief, the selected ions initiate a cascade-of-electron pulse by using a target biased at -22.5 kV and a photon pulse from an organic scintillator is generated. Finally, a Burle photomultiplier tube (-1.35 kV), a discriminator operating at 1.6 mV and a multichannel scaler were utilized. The detector unit is rotatable within the plane of the two beams and the laboratory data were collected at distinct angles. To obtain the LAD of selected m/z, the integrated TOF spectra signal collected are normalized with respect to the signal at the CM angle. The laboratory data including the LAD and TOF spectra were transformed into the CM frame using a forward-convolution routine for the underlying reaction dynamics. The CM translational energy (*P(E*T*)*) and angular (*T(θ)*) flux distributions are varied iteratively before a best fit of the laboratory data is derived. *I(u, θ) ~* *P(u) × T(θ)*,the reactive differential cross section with the product intensity (*I*) as a function of the velocity *u* and the CM angle *θ*, reveals an overall image of the reaction.[29](#_ENREF_29)

**5.3. COMPUTATIONAL:** Electronic structure calculations in the present work were carried out on the C10H8 potential energy surfaces accessed by the carbon - indene (triplet) and dicarbon – styrene (singlet and triplet) reactions. In particular, geometries of the reactants, products, intermediates, and transition states involved in these reactions were optimized at the density functional theory ωB97X-D/6-311G(d,p) level (Table S4),[52](#_ENREF_52),[53](#_ENREF_53) with vibrational frequencies and zero-point vibrational energies (ZPE) computed using the same theoretical approach. Single-point energies were further refined within the explicitly-correlated CCSD(T)-F12/cc-pVTZ-F12 method[54](#_ENREF_54),[55](#_ENREF_55) with ZPE(ωB97X-D/6-311G(d,p)) included, which normally provides the energetic parameters of local minima and transition states within ‘chemical accuracy’ of ~5-10 kJ mol-1 for hydrocarbons in terms of average absolute deviations.[56](#_ENREF_56) The GAUSSIAN 16[57](#_ENREF_57) and MOLPRO 2021[58](#_ENREF_58)program packages were used for the ab initio calculations. Rice-Ramsperger-Kassel-Marcus (RRKM) theory[59](#_ENREF_59),[60](#_ENREF_60) implemented in our Unimol code,[61](#_ENREF_61) were applied to compute energy-dependent rate constants of all unimolecular reaction steps on the C10H8 singlet and triplet surfaces. For H loss steps on the singlet C10H8 surface accessed by the styrene + C2(X1Σg+) reaction, which occur without distinct exit transition states, energy-dependent rate constants were computed using the microcanonical variational transition state theory µ-VTST. Here, the H loss potential energy surfaces (PES) were scanned at the ωB97X-D/6-311G(d,p) level of theory along the reaction coordinate represented by the length of the breaking C-H bonds. Vibrational frequencies for each optimized structure along the PES scan were computed at the same level of theory with projecting the reaction coordinate out. The ωB97X-D/6-311G(d,p) relative energies of these structures were scaled by a factor chosen to match the ωB97X-D/6-311G(d,p) relative energy of the C10H7 +H products to its value obtained at the highest CCSD(T)-F12/cc-pVTZ-F12 level. The optimized structures along the PES profile were considered as variational transition state (VTS) candidates. The unimolecular rate constants for the barrierless H loss were evaluated for each VTS candidate using the RRKM formalism and the minimal rate constant was then selected. Next, product branching ratios at the zero-pressure limit under single-collision conditions were assessed within the steady-state approximation.[59](#_ENREF_59),[60](#_ENREF_60) Assuming single-collision conditions, master equations for unimolecular reactions can be expressed as follows:

where [C]i and [C]j are concentrations of various intermediates or products and *k*n and *k*m are microcanonical rate constants for the formation and consumption of each intermediate *i*, respectively, computed using the RRKM theory. Only a single total-energy level was considered throughout, as for single-collision crossed-beam conditions. In such a case, rate constants of different unimolecular reaction steps are independent of each other, but relative product yields can be determined from the solution of the system of coupled master equations, which is reduced to a system of linear equations within the steady-state approximation, where the concentration of a chosen initial intermediate formed in the bimolecular reaction is taken as a constant and concentrations of all other unimolecular intermediates remain steady, so that their .

**SUPPORTING INFORMATION**

Explanation of the laboratory and center-of-mass frame, supplementary experimental results, experimental parameters (Table S1), statistical branching ratios for the triplet PES for the C+C9H8 reaction (Table S2), the singlet PES for the C2+C8H8 reaction (Table S3), the triplet PES for the C2+C8H8 reaction (Table S4), the optimized coordinates and calculated vibrational frequencies for stationary structures involved in the C-C9H8 reaction (Table S5), C2 (X1Σg+)-C8H8 reaction (Table S6) and C2 (a3Πu)-C8H8 reaction (Table S7); structures that might be formed from naphthyl radicals (Figure S1), Newton diagram of a reactive scattering event (Figure S2), cross sections of our universal CMB machine (Figure S3), additional pathways of the C-C9H8 reaction (Figure S4), singlet and triplet PESs of the reactions of C2 with D3/D5-styrene(Figure S5).

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**Conflicts of Interest**

There are no conflicts to declare.

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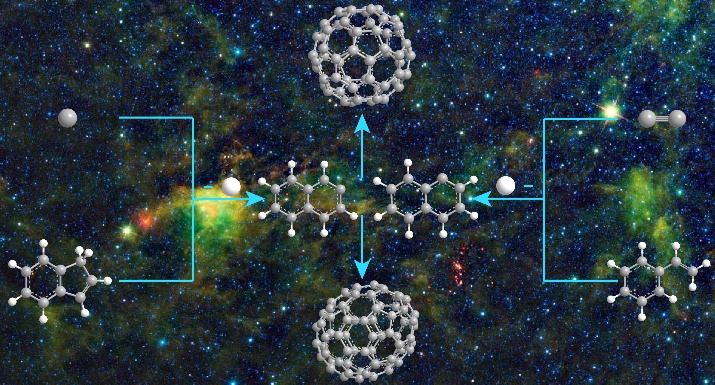
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