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Low Temperature Synthesis and Phosphorescence of Methylcyanotriacetylene

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6
7 ABSTRACT
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10 This paper reports on UV-stimulated synthesis of methylecyanotriacetylene carried out in
11 cryogenic rare gas matrices via coupling of smaller precursors: propyne and cyanodiacetylene.
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13 The detection was possible due to the strong visible $\alpha^3A' \rightarrow X^1A_1$ phosphorescence of CH_3C_7N ,
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15 discovered in the course of this work. The ensuing measurements of electronic spectroscopy
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17 revealed the formally forbidden $B^1E - X^1A_1$ system, as well as the allowed one $E^1A_1 - X^1A_1$,
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19 with origins at approx. 3.32 eV and 5.4 eV, respectively. It was also possible to revisit the
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21 spectroscopic characterization of cyanotriacetylene, HC_7N , formed in parallel to the title
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23 photoproduct. Spectral assignments were assisted with a density functional theory study.
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3 INTRODUCTION
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6 The $\text{H}(\text{C}\equiv\text{C})_n\text{CN}$ molecules were observed with microwave spectroscopy, up to $n = 4$, in diverse
7 extra-terrestrial sources.¹⁻⁶ Methylated cyanopolyacetylenes $\text{CH}_3(\text{C}\equiv\text{C})_n\text{CN}$ are also of
8 astrochemical interest; methylcyanoacetylene, $\text{CH}_3\text{C}_3\text{N}$, was identified in the Taurus Molecular
9 Cloud (TMC)⁷ and more recently in the Sagittarius B2 cloud near the Galactic center.⁸ The
10 presence of methylcyanodiacetylene, $\text{CH}_3\text{C}_5\text{N}$, in TMC has also been confirmed.^{9,10} This latter
11 compound appears, along with $\text{CH}_3\text{C}_3\text{N}$ and $\text{CH}_3\text{C}_7\text{N}$, in astrochemical kinetic models.¹⁰⁻¹³ It
12 may originate, according to Quan and Herbst,¹⁰ from the dissociative recombination of
13 $\text{CH}_3\text{C}_5\text{NH}^+$ cations with electrons or from the encounter of neutral species CN and $\text{CH}_3\text{C}_4\text{H}$.
14 Markwick *et al.*¹² considered the reaction of HC_5N with CH_3OH_2^+ . $\text{CH}_3\text{C}_3\text{N}$ was also included in
15 modelling of Titan's atmosphere^{14,15}, and indeed the mass-spectrometric data from Cassini
16 spacecraft suggested the presence of both $\text{CH}_3\text{C}_3\text{N}$ and $\text{CH}_3\text{C}_5\text{N}$.¹⁴
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32 Addition of a methyl group to a cyanopolyacetylenic chain changes its physical properties. For
33 example, we could see partial thermal decomposition of HC_5N occurring even at -78°C , while the
34 methylated analogue, $\text{CH}_3\text{C}_5\text{N}$, remains stable at least its melting point (92°C ¹⁶). The same trend
35 was observed by us for HC_3N and $\text{CH}_3\text{C}_3\text{N}$: at room temperature, the condensed phase of the
36 latter is stable while the former decomposes and has to be stored below -20°C .
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45 The smallest of methylcyanopolyacetylenes, $\text{CH}_3\text{C}_3\text{N}$, has been the subject of numerous
46 spectroscopic investigations.¹⁷⁻²⁴ For $\text{CH}_3\text{C}_5\text{N}$, the next member of the series, gas-phase
47 rotational transitions^{25,26} and infrared absorption²⁷ were reported, as well as vibrational²⁸ and
48 electronic²⁹ spectra of pure solid and of rare gas matrix-isolated compound. It was reported that
49 photolysis of $\text{CH}_3\text{C}\equiv\text{CH}$ (propyne) mixed with cyanoacetylene leads to $\text{CH}_3\text{C}_5\text{N}$.¹⁶ Kerisit *et al.*³⁰
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3 suggested that $\text{CH}_3\text{C}_7\text{N}$ might analogously form in photochemical reactions involving gaseous
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5 $\text{CH}_3\text{C}\equiv\text{C}-\text{C}\equiv\text{CH}$ (1,3-pentadiyne) and HC_3N precursors (the identification was based on a single
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7 ^1H NMR peak).
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11 The first report to mention $\text{CH}_3\text{C}_7\text{N}$ (methylcyanotriacetylene, 6-methyl-1-cyanohepta-1,3,5-
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13 triyne, octa-2,4,6-triynenitrile) discussed its difficult synthesis, and provided infrared, as well as
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15 UV absorption spectra (for an ether solution).³¹ Microwave transitions were subsequently
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17 investigated.^{26,32,33} Theoretical results are available for rotational constants, molecular geometry,
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19 polarizability, and electric dipole moment.^{34,35}
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24 Our earlier studies^{36–38} demonstrated the appearance of mono- and dicyanopolyacetylenes in
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26 cryogenic UV-stimulated processes, out of shorter $\text{H}(\text{C}\equiv\text{C})_n\text{CN}$ species reacting with one another
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28 or with acetylenic-type molecules $\text{H}(\text{C}\equiv\text{C})_m\text{H}$. Such syntheses have already been shown to work
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30 for cyanodiacetylene (HC_5N),^{36,38} cyanotriacetylene (HC_7N),³⁷ and cyanotetraacetylene
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32 (HC_9N).³⁹ In particular, HC_7N was detected in UV-irradiated (H_2 discharge lamp or 193 nm
33
34 laser) solid Ar doped with either $\text{HC}_3\text{N} + \text{C}_4\text{H}_2$ or $\text{HC}_5\text{N} + \text{C}_2\text{H}_2$ mixture.³⁷ HC_7N was also
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36 produced by electric discharges in a $\text{HC}_3\text{N}/\text{Kr}$ mixture, prior to the trapping onto a cold
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38 substrate.⁴⁰ In all these cryogenic approaches, the generated cyanopolyacetylenes were found to
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40 emit strong phosphorescence, which proved to be the most sensitive spectroscopic indicator of
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42 their presence. However, no photochemically-stimulated cryogenic synthesis has as yet been
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44 reported for any methylated cyanopolyacetylene. Here we aimed at verifying the utility of such
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46 an approach to the formation of $\text{CH}_3\text{C}_7\text{N}$ by coupling the precursors $\text{CH}_3\text{C}_2\text{H}$ and HC_5N .
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48 Propyne, similarly to HC_5N , features an easily photodetachable acetylenic hydrogen atom.^{41,42}
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51 The targeted molecule has indeed appeared along with HC_7N , offering an opportunity to compare
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3 the electronic transitions of the two closely related products and to extend the prior knowledge on
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5 UV-Vis absorption^{43–46} and low temperature phosphorescence^{37,40} of HC₇N. Spectroscopic
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7 assignments were backed with density functional theory calculations of electronic and vibrational
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9 energy levels.
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16 METHODS

17 A. Experimental

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20 Propyne was commercially available from Aldrich (purity > 97%). The second precursor,
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22 cyanodiacetylene, was prepared using the method developed by Trolez & Guillemin.⁴⁷ The
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24 compounds were purified before the experiments by pumping while kept at T<200 K (HC₅N) or
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26 by several freeze-pump-thaw cycles with freezing at T<200 K (CH₃C₂H).
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33 Krypton (purity 4.0, Messer), mixed with the precursor chemicals, was trapped onto a sapphire
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35 substrate held at 30 K inside a closed-cycle helium cryostat. Photochemical transformations were
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37 induced with a 193 nm ArF excimer laser. All details concerning the experimental set-up and
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39 spectroscopic instrumentation were given in a paper by Szczepaniak et al. (this issue).³⁹
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41 Experiments were primarily aimed at identification of the created photoproducts and
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43 characterization of their electronic transitions, by measuring dispersed phosphorescence spectra,
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45 as well as phosphorescence excitation spectra, and phosphorescence lifetime.
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50 An experiment involving CH₃C₂H + HC₃N mixture (HC₃N obtained following the Miller and
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52 Lemon⁴⁸ method), expected to yield CH₃C₅N, was first performed to verify the validity of the
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54 applied approach. The test was positive; not only the phosphorescence of CH₃C₅N, but also that
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3 of HC₅N could be detected in Kr/CH₃C₂H/HC₃N (1000/1/1) matrices previously subjected to 193
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5 nm photolysis.
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8 Two different gas mixtures were used in the main CH₃C₂H + HC₅N experiment.
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10 Cyanodiacetylene was deposited in the amount either the same or twice smaller than that of
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12 propyne. The latter mixture (Kr/CH₃C₂H/HC₅N ratio of 1000/2/1) noticeably favored the
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14 formation of CH₃C₇N over HC₇N.
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25 Theoretical procedures applied here coincide with those described in more detail in a recent paper
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27 by Szczepaniak et al.³⁹ GAUSSIAN 09 (Rev. B. 01)⁴⁹ implementations of the density functional
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29 theory (DFT)⁵⁰ were mostly applied. The B3PW91⁵¹ functional was employed in the derivation
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31 of molecular geometries, harmonic vibrational frequencies, IR absorption intensities, and Raman
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33 activities for the ground and the lowest triplet state, while the time-dependent B3PW91 approach
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35 served for predicting transitions from the ground to excited singlet electronic states. Basis set
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37 aug-cc-pVTZ^{52,53} was applied in all these computations.
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42 Geometry optimizations carried out for excited singlet states of CH₃C₇N led to the structures
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44 (genuine potential energy minima) very slightly deviating from the C_{3v} symmetry. Based on our
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46 former CH₃C₅N study,²⁹ we suppose this effect to be of a purely numerical nature.
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51 Influence of the Herzberg–Teller effect on intensity of the vibronic $\tilde{B}^1\Delta - \tilde{X}^1\Sigma^+$ transitions was
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53 estimated by carrying out oscillator strength calculations for a HC₇N molecule slightly distorted,
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55 mimicking, case by case, the effect of various bending vibrations. The distortion vector was
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3 always parallel to a respective normal coordinate and had a length of 0.1 Å. The coupling effect
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5 was considered as important when the described simulation produced a noticeable oscillator
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7 strength ($f \geq 0.0001$).
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10 11 12 13 14 RESULTS AND DISCUSSION 15

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20
21 Despite obviously different molecular symmetries, HC₅N and CH₃C₅N exhibit strikingly alike
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23 electronic spectra, as noticed by Turowski et al.²⁹ Such similarities, stemming from closely
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25 related chromophores, were expected also for the HC₇N/CH₃C₇N pair. Quantum chemical
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27 predictions for the lowest electronic energy levels of HC₇N and CH₃C₇N are presented in Table 1;
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29 general patterns of electronic state distribution are indeed very similar for both species.
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31 Nevertheless, the states arising from HOMO-LUMO electronic excitations have slightly higher
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33 energies for HC₇N than for CH₃C₇N (resembling what was observed for HC₅N and CH₃C₅N).²⁹
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35 For both molecules, $\tilde{E} - \tilde{X}$ is predicted as the strongest transition. The expected $\alpha - \tilde{X}$ transition
36
37 energy of CH₃C₇N (2.23 eV) is practically identical to that of HC₇N (2.25 eV), just as previously
38
39 calculated and measured for CH₃C₅N and HC₅N (2.90 eV vs. 2.92 eV at CAM-B3LYP/aug-cc-
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41 pVTZ level of theory, 2.94 vs. 2.92 eV in Ar matrix).²⁹
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Table 1. Energy [eV], wavelength [nm], and oscillator strength for transitions involving the ground and excited electronic states of HC₇N and CH₃C₇N, as calculated at B3PW91/aug-cc-pVTZ level of theory. Electronic configurations of the molecules are the following. HC₇N, $X^1\Sigma^+$: [core] $(1\sigma)^2 (2\sigma)^2 (3\sigma)^2 (4\sigma)^2 (5\sigma)^2 (6\sigma)^2 (1\pi)^4 (7\sigma)^2 (2\pi)^4 (8\sigma)^2 (9\sigma)^2 (3\pi)^4 (4\pi)^4 (1\pi^*)^0 (2\pi^*)^0 (1\sigma^*)^0$; CH₃C₇N, X^1A_1 : [core] $(1a_1)^2 (2a_1)^2 (3a_1)^2 (4a_1)^2 (5a_1)^2 (6a_1)^2 (7a_1)^2 (1e)^4 (2e)^4 (8a_1)^2 (3e)^4 (4e)^4 (5e)^4 (1e^*)^0 (1a_1^*)^0 (2e^*)^0$.

HC ₇ N					CH ₃ C ₇ N				
State	Dominant orbital excitation	Vertical energy (λ)	f^a	$0-0$ transition energy (λ)	State	Dominant orbital excitation	Vertical energy (λ)	f^a	$0-0$ transition energy ^c (λ)
$A^1\Sigma^-$	$4\pi \rightarrow 1\pi^*$	3.29 (377)	0	2.84 (437)	A^1A_2	$5e \rightarrow 1e^*$	3.23 (384)	0	2.75 (451)
$B^1\Delta$	$4\pi \rightarrow 1\pi^*$	3.40 (365)	0	2.96 (419)	B^1E	$5e \rightarrow 1e^*$	3.34 (371)	0	2.9 (428)
$C^1\Sigma^-$	$3\pi \rightarrow 1\pi^*$	5.10 (243)	0	4.79 (259)	C^1A_2	$4e \rightarrow 1e^*$	5.05 (246)	0	4.73 (262)
$D^1\Delta$	$3\pi \rightarrow 1\pi^*$	5.27 (235)	0	4.98 (249)	D^1E	$4e \rightarrow 1e^*$	5.22 (238)	0.0001	4.92 (252)
$E^1\Sigma^+$	$4\pi \rightarrow 1\pi^*$	5.84 (212)	2.9	5.63 (220)	E^1A_1	$5e \rightarrow 1e^*$	5.61 (221)	2.9	5.40 (230)
					F^1E	$5e \rightarrow 1a_1^*$	6.18 (201)	0	
					G^1A_2	$5e \rightarrow 2e^*$	6.2 (200)	0	
$F^1\Sigma^+$	$3\pi \rightarrow 1\pi^*$	6.23 (199)	0.78	5.86 (212)	H^1A_1	$4e \rightarrow 1e^*$	6.25 (198)	1.2	
					I^1E	$5e \rightarrow 2e^*$	6.25 (198)	0.0002	
$\alpha^3\Sigma^+$	$4\pi \rightarrow 1\pi^*$			2.25 (551)	α^3A',b	$5e \rightarrow 1e^*$			2.23 (556)

^aValue of zero indicates $f < 5 \cdot 10^{-5}$. ^b C_s geometry uncertain; C_{3v} (leading to A_1 triplet state symmetry) also possible, albeit less likely. ^cDerivation of missing $0-0$ energy entries was impeded by numerical problems (inherent to the applied method).

The nature of $E - X$ transitions is in both molecules similar to that of $A - X$ or $B - X$, in the sense that they all involve mainly the HOMO–LUMO excitations. $A - X$ transitions (either $\Sigma^- - \Sigma^+$ or $A_2 - A_1$) are formally not allowed for both molecules, while $B - X$, forbidden for HC₇N ($\Delta - \Sigma$) and formally allowed for CH₃C₇N ($E - A_1$), is in fact effectively forbidden also for this latter molecule. One may nevertheless expect to see the $B - X$ absorption of CH₃C₇N due to the

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3 Herzberg–Teller vibronic coupling of electronic excitation with one of the vibrational bending
4 modes, a phenomenon reported for shorter cyano-^{22,54,55} and methylcyanopolyacetylenes^{24,29}. The
5 Herzberg-Teller effect was supposed to occur if the electronic transition dipole moment
6 significantly increased from its equilibrium value after distorting a molecule along a particular
7 normal vibrational coordinate. In our simulations performed for HC₇N (see *Computational*
8 *details*), only the ν_{10} mode coordinate produced such an effect. $\mathcal{A} - \mathcal{X}$ bands are expected to have
9 very low intensity; merely traces of that system were observed for CH₃C₅N²⁹ and for HC₅N.⁵⁵

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20 Molecular geometries for the states \mathcal{X} , \mathcal{B} , \mathcal{E} , and α , involved in the experimentally explored
21 electronic transitions of HC₇N and CH₃C₇N, are depicted in Figure 1 (predictions for the other
22 states of Table 1 can be found in Figure S1). Ground-state HC₇N and CH₃C₇N possess conjugated
23 carbon-nitrogen backbones of closely similar geometries. Noteworthy, heavy-atom skeleton
24 changes accompanying $\mathcal{B} - \mathcal{X}$ transition are, for both molecules, almost identical to those of
25 $\alpha - \mathcal{X}$. One may therefore expect the main vibronic progressions in $\mathcal{B} - \mathcal{X}$ absorption and in
26 phosphorescence spectra to be shaped by vibrational modes of similar nature, just as observed for
27 the HC₅N/CH₃C₅N pair.²⁹ Conversely, the geometry change occurring upon the transition from \mathcal{X}
28 to \mathcal{E} is much different from those characterizing $\alpha - \mathcal{X}$ or $\mathcal{B} - \mathcal{X}$.

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42 It is worth noting that our optimizations of CH₃C₇N geometry formally indicate, for the α state,
43 the point group C_s , rather than C_{3v} ; the effect comes primarily from a small lowering of the CH₃
44 group symmetry (see Figure 1). We currently cannot conclude whether this deviation from C_{3v} is
45 or is not a numerical artifact. However, even if it were real, it would have not, (considering its
46 magnitude) show up in the low-resolution spectra produced in this study.

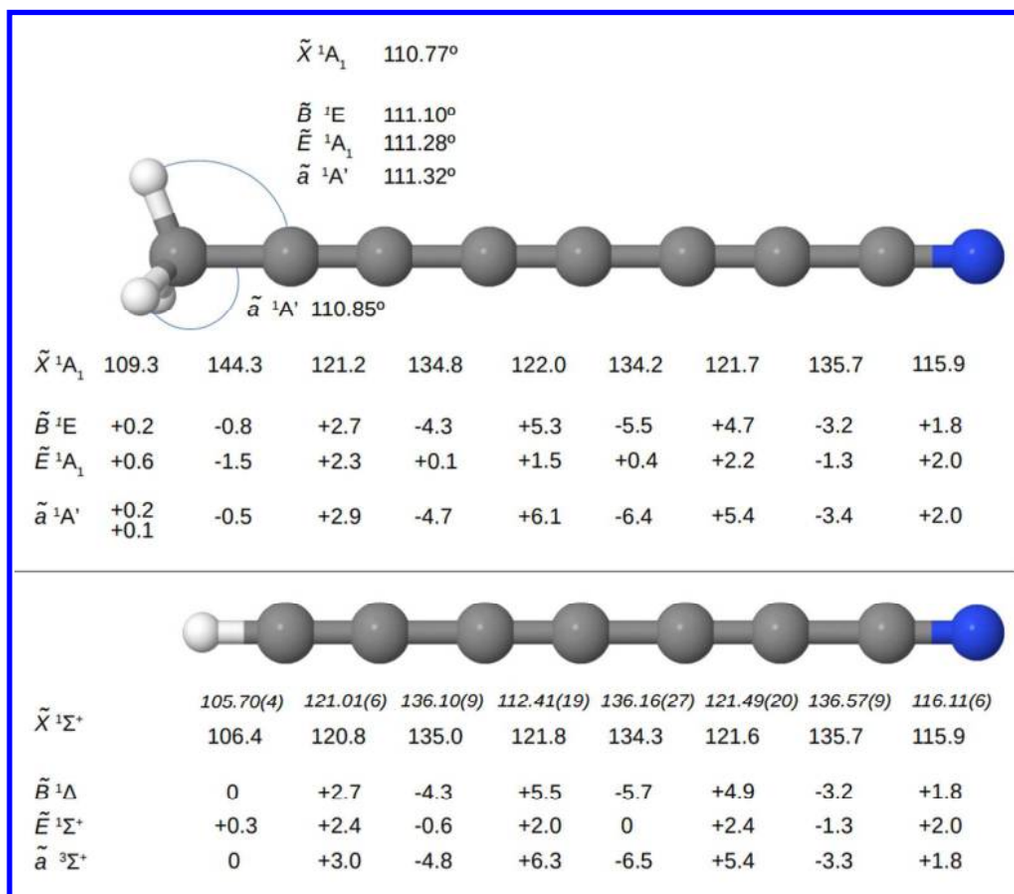


Figure 1. $\text{CH}_3\text{C}_7\text{N}$ and HC_7N geometry in the ground and excited electronic states, as derived at the B3PW91/aug-cc-pVTZ level of theory. Experimental parameters of HC_7N (in italics) are cited after McCarthy et al., Ref. 56. Interatomic distances in pm. Values listed for the excited electronic states are relative, calculated with respect to \tilde{X} . For higher electronic states, see Figure S1 of Supporting Information.

Table 2. Harmonic wavenumbers of vibrational modes (ν ; scaled by 0.96) for the ground and selected excited electronic states of HC₇N and CH₃C₇N, together with ground-state infrared intensities (*IR*) and Raman activities (*R*), as derived at the B3PW91/aug-cc-pVTZ level of theory. Mode symmetry descriptions refer strictly to the ground electronic states. See Tables S1 and S2 of the Supporting Information for the visualization of vibrations.

HC ₇ N						CH ₃ C ₇ N					
Mode	\tilde{X}			\tilde{B}	\tilde{E}	Mode	\tilde{X}			\tilde{B}	\tilde{E}
	ν cm ⁻¹	<i>IR</i> km/mol	<i>R</i> Å ⁴ /amu	ν cm ⁻¹	ν cm ⁻¹		ν cm ⁻¹	<i>IR</i> km/mol	<i>R</i> Å ⁴ /amu	ν cm ⁻¹	ν cm ⁻¹
σ symmetry						a_1 symmetry					
ν_1	3324	156	18	3316	3290						
						ν_1	2910	8.9	1223	2882	2841
ν_2	2253	88	1600	2117	2172	ν_2	2260	7.9	100	2114	2180
ν_3	2197	133	9800	2033	2073	ν_3	2208	410	10000	2038	2132
ν_4	2154	5	1600	1907	1949	ν_4	2187	290	8500	1968	1969
ν_5	2049	0	445	1819	1855	ν_5	2093	0.014	7.3	1848	1917
						ν_6	1364	1.3	89	1460	1323
						ν_7	1333	7.7	78	1332	1295
ν_6	1295	0	24	1393	1252	ν_8	1066	0.7	13	1082	1061
ν_7	902	3	0.41	926	884	ν_9	758	6.7	0.88	767	745
ν_8	464	0	2.7	477	456	ν_{10}	402	1.0	1.9	412	395
π symmetry						e symmetry					
						ν_{11}	2976	1.9	370	2892	2894
						ν_{12}	1398	21	27	1384	1352
						ν_{13}	991	4.3	12	1022	953
ν_9	643	80	14	522 ^a	541						
ν_{10}	602	1	57	553 ^a	517	ν_{14}	574	2.3	33	523	490
ν_{11}	523	2	0.001	483	434	ν_{15}	518	4.4	0.092	476	437
ν_{12}	461	10	4.0	434	412	ν_{16}	463	2.2	5.3	419	417
						ν_{17}	340	6.4	4.7	289	212
ν_{13}	287	0	1.7	290	211	ν_{18}	226	0.036	0.28	172	193
ν_{14}	162	13	1.0	153	71	ν_{19}	124	12	0.27	92	57
ν_{15}	61	0	1.8	60	41	ν_{20}	47	0.0010	0.072	43	39

^aReversed order; ν_9 corresponds to the H-C vibration.

Vibrational frequencies computed for the selected electronic states (of interest to the analysis of IR spectra or of vibronic features) are collected in Table 2. The only available IR absorption spectrum of CH₃C₇N was measured over 60 years ago by Bohlmann and Mannhardt³¹ for a CCl₄ solution; digitization of the published figure points to band maxima at approximately 2290, 2240, 2210, and 2190 cm⁻¹. Our DFT results predict two very intense IR transitions, ν_3 and ν_4 , at,

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3 respectively 2208 cm^{-1} and 2187 cm^{-1} , i.e. matching the two of those observed. Other bands are
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5 expected to be much weaker. The 2240 and 2290 cm^{-1} bands observed in CCl_4 could correspond
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7 to the ν_2 band, possibly split by an anharmonic interaction (such interactions, involving triple-
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9 bond stretching modes, were recognized²⁸ for $\text{CH}_3\text{C}_5\text{N}$). In our study³⁷ of the photochemically
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11 generated non-methylated analogue, HC_7N , the IR absorption spectroscopy of solid Ar-isolated
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13 substance revealed traces of ν_3 (2202.0 cm^{-1}) and ν_1 (3323.1 cm^{-1}) bands, close to the
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15 theoretically predicted values. However, in the present experiments, aimed at $\text{CH}_3\text{C}_7\text{N}$, no IR
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17 absorption of this compound could be detected, and the identification was based solely on the
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19 analysis of electronic luminescence, just as in our recent study on UV-assisted synthesis of HC_9N
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21 from smaller chain precursor molecules.
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30 B. Dispersed phosphorescence

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33 In line with our previous studies, the aim here was to detect $\text{CH}_3\text{C}_7\text{N}$ by means of its
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35 phosphorescence, predicted to appear (Table 1) in the vicinity of the $\alpha \rightarrow \mathcal{X}$ bands of HC_7N .
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37 However, the use of HC_5N as one of the precursors implied the observation of its own intense
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39 phosphorescence, partly coinciding with anticipated $\text{CH}_3\text{C}_7\text{N}/\text{HC}_7\text{N}$ emissions. Consequently, all
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41 detected HC_5N bands had to be thoroughly assigned. A listing of these, appending our former
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43 report,⁵⁵ is available as the Supporting Information, Table S3 and Figures S2 and S3.
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49 Luminescence from photoproducts generated in a $\text{Kr}/\text{CH}_3\text{C}_2\text{H}/\text{HC}_5\text{N}$ matrix was detected
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51 especially in spectral windows around 19 700 cm^{-1} (508 nm, 2.44 eV; see Figure 2) and 17 500
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53 cm^{-1} (571 nm, 2.17 eV), corresponding to the most intense phosphorescence bands of HC_7N .³⁷
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55 An emission/excitation graph (for the first of the above-mentioned spectral ranges) is shown in
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Figure 2a. Two closely spaced emission bands are clearly discerned there; their individual excitation patterns indicate the presence of two disparate products. Another strong doublet, with components similarly spaced by $\sim 60 \text{ cm}^{-1}$, appeared in the $17\,500 \text{ cm}^{-1}$ region. The stronger component shown in Figure 2b corresponds to a lower wavenumber, and is coincident with an already known^{37,57} HC_7N emission band. It further gains intensity when excited at $46\,430 \text{ cm}^{-1}$ (215.4 nm , 5.76 eV ; see Figure 2a), in agreement with an excitation maximum previously observed for HC_7N . The other band, with a different excitation spectrum (maximal signal observed upon excitation at $43\,150 \text{ cm}^{-1}$, 231.7 nm , 5.35 eV), can be assigned to $\text{CH}_3\text{C}_7\text{N}$. Indeed, theory predicted (Table 1) for HC_7N almost identical phosphorescence origin energy as for its methylated analogue. It should be noted that the applied computational method is not precise enough to rely on the exact magnitude of the difference between these two values.

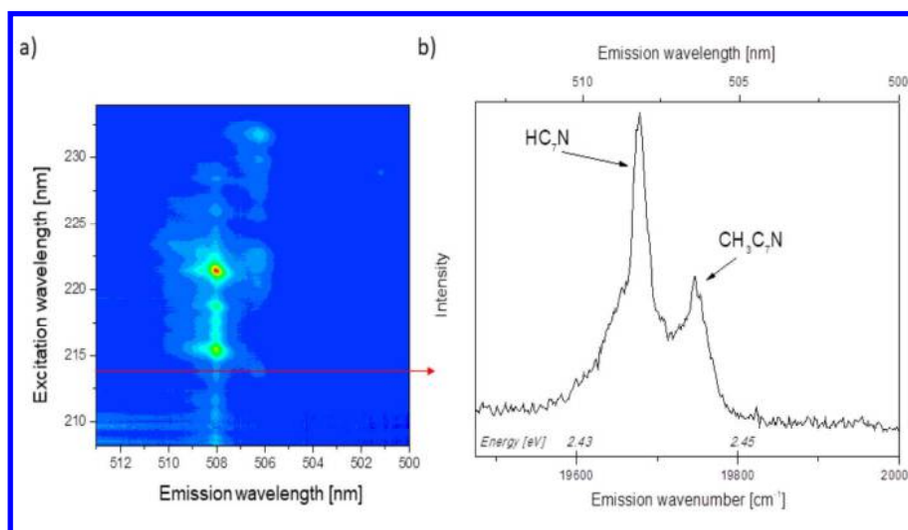


Figure 2. The excitation-emission map of a photolyzed (193 nm) $\text{Kr}/\text{CH}_3\text{C}_2\text{H}/\text{HC}_5\text{N}$ ($1000/1/1$) matrix showing the spectral features of HC_7N and $\text{CH}_3\text{C}_7\text{N}$ phosphorescence (a). Red color represents the highest and deep blue the lowest phosphorescence intensity. Spectrum (b) corresponds to cross-sectioning the map along the red line, i.e. to phosphorescence excited at 213.5 nm ($46\,840 \text{ cm}^{-1}$).

The upper trace of Figure 3 presents HC₇N phosphorescence for a photolyzed Kr/CH₃C₂H/HC₅N matrix, excited at a maximum of UV absorption (215.4 nm, Figure 2). The respective wavenumbers of vibronic bands are listed in Table 3, together with the values measured in previous Ar/HC₅N/C₂H₂³⁷ and Kr/HC₅N/C₂H₂⁵⁷ experiments; an additional, fairly weak band, identified as 3₃⁰, has been presently detected.

Table 3. HC₇N phosphorescence (α ³Σ⁺ → $\bar{\chi}$ ¹Σ⁺) bands, as measured in cryogenic matrices. Wavenumber (ν) values in cm⁻¹. Relative wavenumbers ($\Delta\nu$) calculated with respect to the preceding element of the vibronic progression.

Ar (Ref. 37)		Kr (Ref. 57)		Kr (this work)		Assignment
ν	$\Delta\nu$	ν	$\Delta\nu$	ν	$\Delta\nu$	
19764	-	19700	-	19690	-	0 ₀ ⁰
17565	2199	17500	2200	17490	2200	3 ₁ ⁰
15378	2187	15310	2190	15310	2180	3 ₂ ⁰
				13140 ^a	2170 ^a	3 ₃ ⁰

^a±20 cm⁻¹; the band is weak and broad.

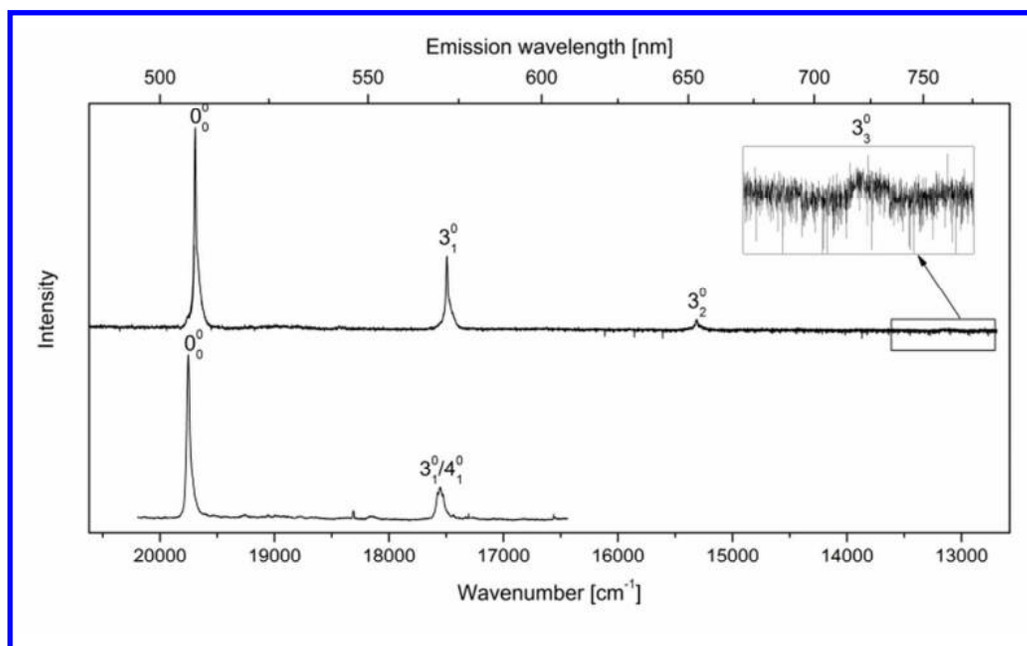


Figure 3. Top: HC₇N phosphorescence spectrum of a photolysed (193 nm) Kr/CH₃C₂H/HC₅N (1000/2/1) matrix excited at 46 430 cm⁻¹ (5.76 eV, 215.4 nm). Bottom: CH₃C₇N phosphorescence of a photolyzed (193 nm) Kr/CH₃C₂H/HC₅N (1000/2/1) matrix excited at 43 150 cm⁻¹ (5.35 eV, 231.7 nm).

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5 Just as indicated by Figure 2a, the excitation at 231.7 nm led preferably to the phosphorescence
6 of CH₃C₇N (lower curve of Figure 3), with prominent 19 750 cm⁻¹ and 17 550 cm⁻¹ bands, very
7 similar to that of HC₇N. Of note, the observed CH₃C₇N phosphorescence was less intense than
8 that of HC₇N, preventing the detection of lower frequency features. The position (corresponding
9 to 2.45 eV) of the most intense of the said bands matches reasonably well (the same quality of
10 agreement as for HC₇N) with what is expected (2.23 eV, Table 1) for the vibrationless origin of
11 the $\alpha^3A' \rightarrow X^1A_1$ system. The bands are 2200 cm⁻¹ apart, in acceptable agreement with the
12 present predictions (see Table 2) for either ν_4 or ν_3 X -state frequencies. Using the HC₇N case as a
13 guide, preference could possibly be given to the pseudosymmetric mode ν_4 (2187 cm⁻¹; see Table
14 S2 in Supporting Information for the visualization). Similar reasoning was used²⁹ in the analysis
15 of the CH₃C₅N spectrum, based on its resemblance to that of HC₅N. However, constancy of
16 HC₇N/CH₃C₇N “splitting” observed in the consecutive elements of the vibronic progression
17 suggests that the vibrational quantum governing the progression of CH₃C₇N is very close to 2200
18 cm⁻¹, i.e. to the value found for the analogous progression of HC₇N. This points to the mode ν_3
19 (2208 cm⁻¹) rather than ν_4 . Additionally, ν_3 shows the highest Raman activity, and the stretchings
20 of such a character have already been shown to shape the main vibronic progressions of
21 cyanopolyacetylenic molecules.^{40,57,58} Broadness of the 17 550 cm⁻¹ band may in fact suggest the
22 presence of overlapping ν_3 and ν_4 progressions (the Raman activity of ν_4 is also high).
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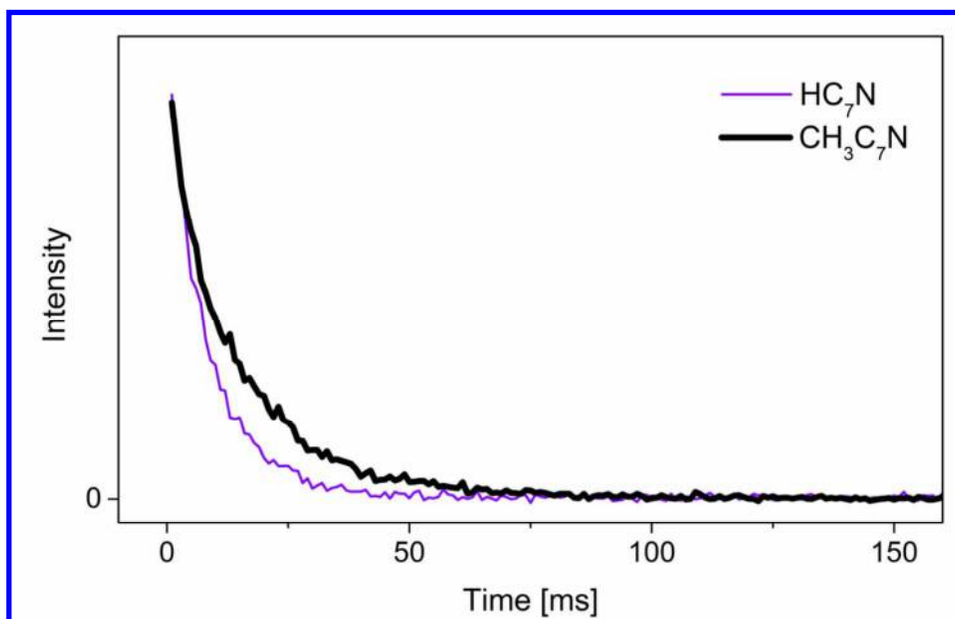


Figure 4. Phosphorescence lifetime measurements for HC_7N and $\text{CH}_3\text{C}_7\text{N}$, excited at $46\,430\text{ cm}^{-1}$ (215.4 nm) and $43\,150\text{ cm}^{-1}$ (231.7 nm), respectively.

HC_7N and $\text{CH}_3\text{C}_7\text{N}$ emission decays were registered, as demonstrated in Figure 4. The band assigned to the origin of HC_7N phosphorescence showed a mono-exponential decay with $\tau = (8 \pm 0.5)\text{ ms}$. This Kr-matrix value is by far smaller than $\tau = (34 \pm 1.5)\text{ ms}$ reported³⁷ for solid Ar. The band marking the origin of $\text{CH}_3\text{C}_7\text{N}$ emission was decaying with $\tau = (15 \pm 1)\text{ ms}$.

Table 4. Phosphorescence lifetime (ms) for solid Kr-isolated cyanopolyacetylenes and methylcyanopolyacetylenes.

HC_nN	τ	$\text{CH}_3\text{C}_n\text{N}$	τ
HC_5N^a	40	$\text{CH}_3\text{C}_5\text{N}^b$	150
HC_7N	8	$\text{CH}_3\text{C}_7\text{N}$	15
HC_9N^c	3.9		

^aAfter Ref. 57. ^bAfter Ref. 29. ^cAfter Ref. 39.

These data, together with those reported for other $\text{H}(\text{C}\equiv\text{C})_n\text{CN}$ and $\text{CH}_3(\text{C}\equiv\text{C})_n\text{CN}$ molecules isolated in solid Kr, are collected in Table 4. Within each of these series, there is a pronounced decrease of lifetime as the length of a polyacetylenic backbone increases. Such a regularity most

likely originates in the diminishing singlet-triplet gap and in the increasing number of chain-deforming vibrational modes; both factors may enhance non-radiative relaxation channels. H-capped cyanopolyacetylenes are characterized by shorter phosphorescence lifetimes than their methylated derivatives. It remains an open question whether this effect is of an intramolecular or of an environmental nature. One may speculate that matrix trapping sites occupied by $C_{\infty v}$ molecules are tighter than those available to their C_{3v} analogues (less Kr atoms need to be removed from an fcc lattice to accommodate an H-capped than a CH_3 -capped molecule), and hence the interaction of $C_{\infty v}$ species with lattice phonons is stronger, promoting the non-radiative electronic relaxation.

C. Singlet excited electronic states

Resemblance of the chromophores of HC_7N and $\text{CH}_3\text{C}_7\text{N}$ is illustrated by the predicted geometries (Figure 1). It is also reflected in the corresponding DFT-derived electronic transition energies (Table 1), characterized by similarities already seen within the pairs of methyl- and hydrogen-capped cyanopolyacetylenes $\text{CH}_3\text{C}_3\text{N}/\text{HC}_3\text{N}$ ^{22,24,54} or $\text{CH}_3\text{C}_5\text{N}/\text{HC}_5\text{N}$ ²⁹. Singlet-singlet electronic absorption features of $\text{CH}_3\text{C}_7\text{N}$ and HC_7N (accessed here by means of phosphorescence excitation) were therefore expected to partially overlap.

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3 yet another promoting π -symmetry mode, tentatively identified here as ν_9 (B -state wavenumber
4 prediction: 522 cm^{-1}). Given that the lowest-wavenumber doublet was detected at $27\ 620/27\ 640$
5 cm^{-1} , the $B - X$ origin is expected around $27\ 100$ or $27\ 120\ \text{cm}^{-1}$, i.e. $3.36\ \text{eV}$. Further profiting
6 cm^{-1} , the $B - X$ origin is expected around $27\ 100$ or $27\ 120\ \text{cm}^{-1}$, i.e. $3.36\ \text{eV}$. Further profiting
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8 from analogies with shorter $\text{H}(\text{C}\equiv\text{C})_n\text{CN}$ species (which, compared to HC_7N , feature similar
9 molecular distortions upon $B - X$ excitation, similar patterns of the excited electronic states
10 distribution, and show similarities in phosphorescence spectra), the spacing of the main vibronic
11 progression can be attributed to the B -state mode ν_2 (predicted wavenumber $2117\ \text{cm}^{-1}$),
12 analogous to the ν_3 stretching of X (see Table S1). Wavenumbers of HC_7N bands identified in
13 Figure 5 are listed in Table 5.
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25 The analogous, low energy region of the $\text{CH}_3\text{C}_7\text{N}$ phosphorescence excitation spectrum showed
26 just two weak bands attributable to the $B\ ^1\text{E} - X\ ^1\text{A}_1$ system, at $29\ 330\ \text{cm}^{-1}$ ($3.64\ \text{eV}$) and $31\ 400$
27 cm^{-1} (i.e. spaced by the value of $2070\ \text{cm}^{-1}$, characteristic of triple bond stretching frequencies).
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29 The known spectra of smaller (i.e. $n = 1$ or 2) $\text{H}(\text{C}\equiv\text{C})_n\text{CN}$ and $\text{CH}_3(\text{C}\equiv\text{C})_n\text{CN}$ molecules indicate
30 that the forbiddance of $B - X$ transitions (and hence the lack of vibrationless origin bands),
31 strictly valid for $\text{H}(\text{C}\equiv\text{C})_n\text{CN}$ species, is in practice effective also for the methyl-capped
32 derivatives. It should be recalled that theory predicts a shift of approx. $500\ \text{cm}^{-1}$ (see Table 1)
33 between the $B - X$ origins of HC_7N and $\text{CH}_3\text{C}_7\text{N}$. A very similar value of $475\ \text{cm}^{-1}$ was
34 experimentally found for this shift in the analogous case of the HC_5N , $\text{CH}_3\text{C}_5\text{N}$ pair of
35 molecules.^{29,55} Therefore, the $29\ 330\ \text{cm}^{-1}$ and $31\ 400\ \text{cm}^{-1}$ features can be related to the HC_7N
36 doublets around $29\ 750$ and $31\ 860\ \text{cm}^{-1}$, respectively. The detected bands are presumably due to
37 the coupling of $B - X$ with a ‘zig-zag’ distortion of the carbon-nitrogen chain, combined with a
38 chain stretching mode. This suggests the placement of $B\ ^1\text{E} - X\ ^1\text{A}_1$ origin at approx. $3.32\ \text{eV}$, an
39 estimation coming from the appropriate lowering of the above-mentioned value $29\ 330\ \text{cm}^{-1}$, by
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subtracting from it the frequencies of (i) the stretching that governs the main vibronic progression (measured value: 2070 cm^{-1}) and (ii) the vibronic-coupling mode (most likely the ν_{14} zig-zag chain-bending; predicted value: 523 cm^{-1}).

Table 5. Wavenumbers (cm^{-1}) of $\mathcal{B}^1\Delta - \mathcal{X}^1\Sigma^+$ bands tentatively identified in the phosphorescence excitation spectrum of HC_7N in solid krypton.

ν	Assignment
27 620	9_0^1
27 640	10_0^1
29 460	$2_0^1 14_0^1 15_0^1$
29 520	$2_0^1 13_0^1$
29 730	$2_0^1 9_0^1$
29 770	$2_0^1 10_0^1$
30 190	$2_0^1 11_0^2$
30 390	$2_0^1 12_0^2 14_0^2$
31 600	$2_0^2 14_0^2 15_0^2$
31 670	$2_0^2 13_0^1$
31 840	$2_0^2 9_0^1$
31 900	$2_0^2 10_0^1$
31 950	Site / $2_0^2 10_0^1 15_0^1$
32 030	Site / $2_0^2 10_0^1 14_0^1$
32 320	$2_0^2 11_0^2$
32 540	$2_0^2 12_0^2 14_0^2$
34 030	$2_0^3 9_0^1$
34 090	$2_0^3 10_0^1$

Figure 6 presents the dependence of experimentally found $\mathcal{B} - \mathcal{X}$ origin wavelength on the molecule chain length, for two homologues series: $\text{H}(\text{C}\equiv\text{C})_n\text{CN}$ and $\text{CH}_3(\text{C}\equiv\text{C})_n\text{CN}$. Also shown, for comparison, are the corresponding theoretical predictions, separately for vertical and vibrationless transitions. All sets of data follow linear patterns and, for a given n value, the wavelengths found for $\text{H}(\text{C}\equiv\text{C})_n\text{CN}$ and $\text{CH}_3(\text{C}\equiv\text{C})_n\text{CN}$ are very close to one another. Moreover,

experimental values are consistently lower than the theoretically derived ones, with a fortuitous agreement between the wavelengths of experimental vibrationless transitions and of theoretical vertical transitions (as already reported for $\text{H}(\text{C}\equiv\text{C})_n\text{CN}$ molecules).³⁹

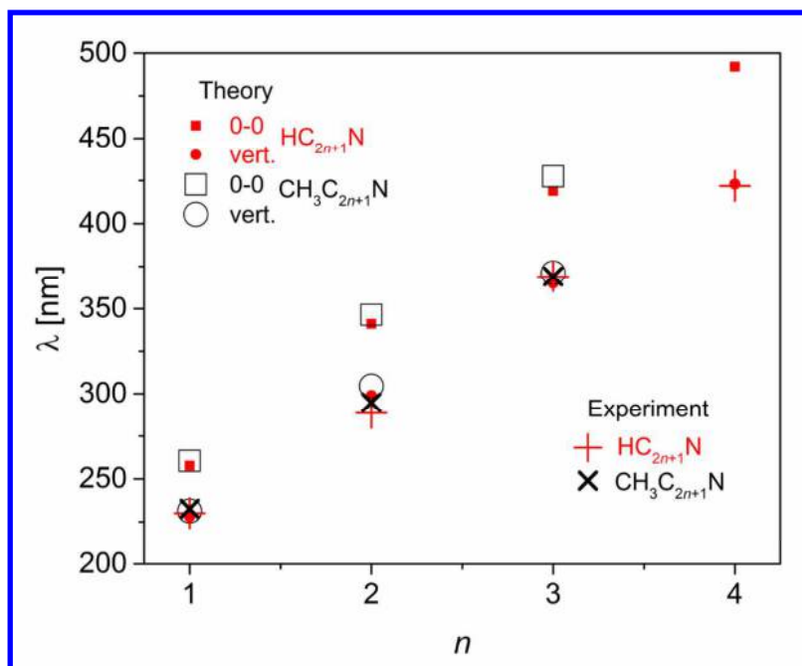


Figure 6. Dependence of $\tilde{B}-\tilde{X}$ transition wavelength (λ) on the carbon chain length (n), for HC_{2n+1}N and $\text{CH}_3\text{C}_{2n+1}\text{N}$. Theoretical values are those of this work and Refs. 28, 38. Experimental data correspond to $\tilde{B}-\tilde{X}$ origin wavelengths in solid Kr (this work and Refs. 29, 55), with the exception of $n = 1$: for HC_3N the value found in solid Ar⁵⁹ is given (Ar-to-Kr shift being negligible in the scale of the plot), and the $\text{CH}_3\text{C}_3\text{N}$ entry corresponds to the corrected gas-phase value²⁴ gas-to-Kr shift is estimated as -500 cm^{-1} .

The high-energy region of phosphorescence excitation spectra, above $43\,000\text{ cm}^{-1}$, gave access to fully allowed $\tilde{E} - \tilde{X}$ transitions of HC_7N and $\text{CH}_3\text{C}_7\text{N}$. Just as predicted (Table 1), the two $\tilde{E} - \tilde{X}$ systems are located very close to one another.

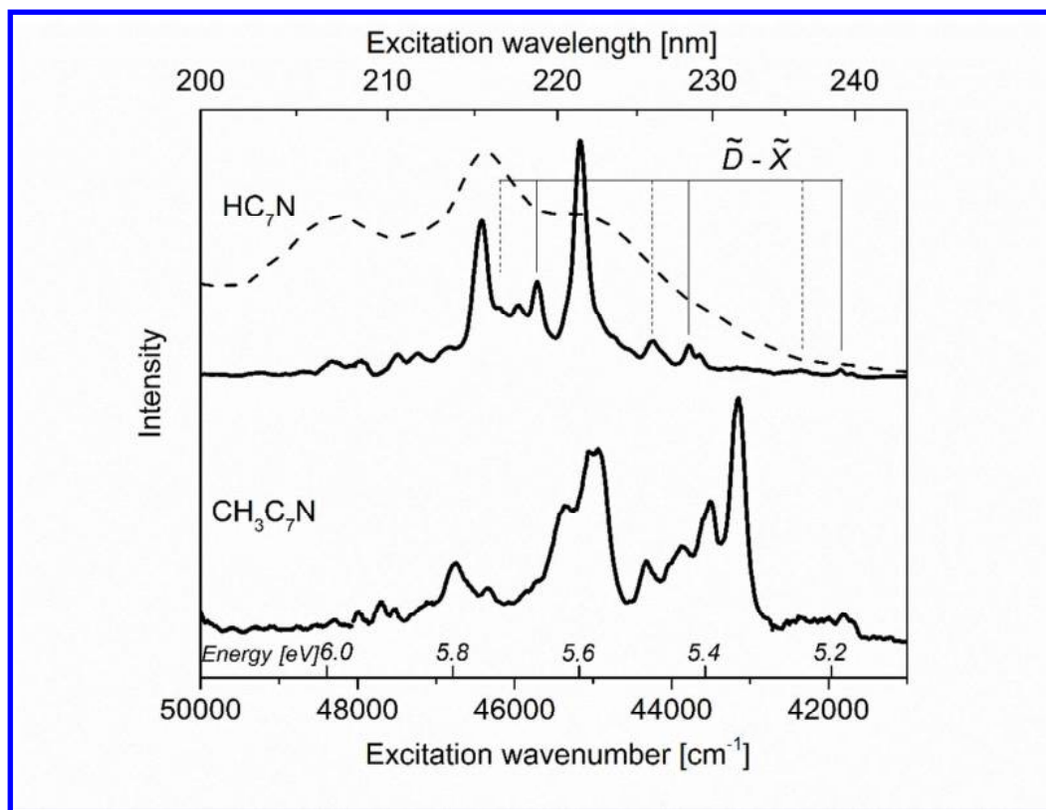


Figure 7. Top: $\tilde{E} - \tilde{X}$ transitions in the excitation spectrum of HC_7N phosphorescence (detection at the $0-0$ band, $19\,690\text{ cm}^{-1}$) emitted from a photolyzed (193 nm) $\text{Kr}/\text{CH}_3\text{C}_2\text{H}/\text{HC}_5\text{N}$ ($1000/1/1$) matrix. Dashed trace (reproduced from Ref. 43) corresponds to the absorption spectrum of HC_7N in acetonitrile. Tentatively identified elements of the $\tilde{D} - \tilde{X}$ vibronic progression are indicated. **Bottom:** Phosphorescence excitation spectrum (detection at the $0-0$ band, $19\,750\text{ cm}^{-1}$) assigned to the $\tilde{E} - \tilde{X}$ electronic system of $\text{CH}_3\text{C}_7\text{N}$, coming from a $\text{Kr}/\text{CH}_3\text{C}_2\text{H}/\text{HC}_5\text{N}$ ($1000/2/1$) matrix irradiated at 193 nm . Wavelength-dependent laser intensity variations might distort the phosphorescence intensity patterns.

Figure 7 (bottom) presents the relevant system of bands identified as due to $\text{CH}_3\text{C}_7\text{N}$. A UV absorption feature reported by Bohlmann³¹ for an ether solution of $\text{CH}_3\text{C}_7\text{N}$ (centered at $44\,740\text{ cm}^{-1}$), remains in reasonable agreement with present data. The first strong band of the excitation spectrum is located at $43\,150\text{ cm}^{-1}$ (5.35 eV or 231.7 nm), very close to the predicted origin of $\tilde{E}^1\text{A}_1 - \tilde{X}^1\text{A}_1$ transitions (5.40 eV , see Table 1). Several other bands of this system are

tentatively identified, as proposed in Table 6. The second strong maximum, around 45 000 cm^{-1} , is broad and may correspond to more than one band. Therefore, the main vibronic spacing, given by a distance between the two major spectral features, is not well defined. Its value of $\sim 1800 \text{ cm}^{-1}$ is smaller than the closest DFT prediction for \tilde{E} state modes listed in Table 2: ν_5 of 1917 cm^{-1} or ν_4 of 1969 cm^{-1} . It should be noted that the excitation at 43 150 cm^{-1} , which leads to the strongest phosphorescence, was also found to potentially destroy the molecule over time (an effect that was not significant for the H-capped analogue). This contributed to a relatively high intensity of detected HC_7N bands compared to those of $\text{CH}_3\text{C}_7\text{N}$.

Table 6. Vibronic band attributions proposed for the $\tilde{E} - \tilde{X}$ electronic system of $\text{CH}_3\text{C}_7\text{N}$. Wavenumbers (cm^{-1}) rounded to the nearest 10. Relative wavenumbers give the distance from the vibrationless 0_0^0 band.

ν	$\Delta\nu$	Tentative assignment
43 150	0	0_0^0
43 520	370	10_0^1
43 880 ^a	730	$9_0^1 / 10_0^2$
44 330	1180	$9_0^1 10_0^1$
45 000 ^b	1850	5_0^1
45 370	2220	$5_0^1 10_0^1 / 2_0^1$
46 350	3200	$5_0^1 6_0^1 / 2_0^1 8_0^1$
46 770	3620	5_0^2

^a This broad band is shifted by $\sim 2000 \text{ cm}^{-1}$ from another broad feature at lower energy, assigned to $\tilde{D} - \tilde{X}$ (see text), it may therefore also belong to the $\tilde{D} - \tilde{X}$ system, rather than to $\tilde{E} - \tilde{X}$.

^b Broad band, possibly comprising a 44940 / 45070 cm^{-1} doublet, tentatively due to 5_0^1 coupled with 13_0^2 .

The top part of Figure 7 shows the analogous region of the phosphorescence excitation spectrum, as measured for HC_7N (qualitatively similar data, for the compound produced in electric discharges through a $\text{HC}_3\text{N}/\text{Kr}$ mixture, were reported by Turowski et al.⁴⁰). The first intense

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3 band, at $45\,180\text{ cm}^{-1}$ (221.3 nm), almost matches the theoretical prediction of 5.63 eV (Table 1).
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5 As can be seen in Figure 7, a much less resolved UV absorption of HC₇N in acetonitrile, reported
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7 in Wakabayashi *et al.*,⁴³ appears in the same spectral region. The resemblance of HC₇N and
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9 HC₉N excitation spectra, reported by Szczepaniak *et al.*,³⁹ helps in the assignment of bands. More
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11 precisely, at the low energy side, the spectra of both of HC₇N and HC₉N show weak features,
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13 most likely due to the $\mathcal{D} - \mathcal{X}$ system, with a vibronic spacing of $\sim 1930\text{ cm}^{-1}$ (a typical triple bond
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15 stretching frequency). The $45\,180\text{ cm}^{-1}$ band does not belong to the $\mathcal{D} - \mathcal{X}$ progressions (marked
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17 in Figure 7) and is assigned here to the origin of the $\mathcal{E} - \mathcal{X}$ system. Table 7 lists tentative
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19 identifications of the most intense bands.
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25 As previously mentioned, theoretical predictions pointed to the similarities between the first
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27 excited states of HC₇N and CH₃C₇N (Table 1). However, as depicted by Figure 7, the measured
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29 patterns of $\mathcal{E} - \mathcal{X}$ bands differ for the two molecules, and the presence of $\mathcal{D} - \mathcal{X}$ transitions in
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31 the CH₃C₇N spectrum is less obvious (even if possibly manifested with unresolved bands around
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33 $42\,000\text{ cm}^{-1}$). Long vibronic progressions are not expected in $\mathcal{E} - \mathcal{X}$ spectra, considering that the
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35 respective electronic excitation is accompanied by little change in molecular geometry. Our
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37 tentative band assignments, provided in Tables 6 and 7, do not involve, for the two molecules, the
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39 same types of vibrational modes; it is mainly a C-C stretching (ν_6) for HC₇N and a triple bond
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41 stretching (ν_5) for CH₃C₇N. No close equivalent to ν_6 of HC₇N does in fact exist for CH₃C₇N (see
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43 Tables S1 and S2 of the Supporting Information), what may explain the differences in band
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45 patterns observed for these spectra. Moreover, the vibronic pattern of HC₇N spectrum, more
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47 difficult to interpret than that found for CH₃C₇N, might result from the energetic proximity of the
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49 two allowed electronic systems: $\mathcal{E} - \mathcal{X}$ and $\mathcal{F} - \mathcal{X}$. In the case of CH₃C₇N, the separation of \mathcal{E}
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and \dot{H} states (analogous to \dot{E} and \dot{F} of HC_7N) is substantially larger, probably reducing their mutual interaction.

Table 7. Wavenumbers (cm^{-1}) of the most intense bands observed in the $\dot{E} - \dot{X}$ region of the HC_7N excitation spectrum. $\dot{D} - \dot{X}$ bands listed in italics. Relative wavenumbers give the distance from the vibrationless (0_0^0) band of the $\dot{E} - \dot{X}$ system.

ν	$\Delta\nu$	Tentative assignment
41 720		<i>\dot{D}</i> (minor band)
41 860		<i>\dot{D}</i> (main band)
43 660		<i>\dot{D}</i> (minor band) + ν (triple bond stretch)
43 790		<i>\dot{D}</i> (main band) + ν (triple bond stretch)
44 250		<i>\dot{D}</i> (broad band)
45 180	0	0_0^0
45 720		<i>\dot{D}</i> (main band) + 2ν (triple bond stretch)
45 960	780	<i>\dot{D} / \dot{E}^a</i>
46 210		<i>\dot{D}</i> (broad band) + ν (triple bond stretch)
46 430	1250	6_0^1
46 840	1660	$6_0^1 8_0^1$
47 230	2050	3_0^1
47 510	2330	6_0^2
47 960	2780	$6_0^2 8_0^1$
48 340	3160	$6_0^2 8_0^2$

^a An unassigned band belonging either to $\dot{E} - \dot{X}$ (possibly perturbed by coupling to a $\dot{D} - \dot{X}$ transition) or to $\dot{D} - \dot{X}$.

D. Chain-coupling mechanisms

Studies on the synthesis of HC_7N and HC_9N in rare gas matrices suggested that the dominant path may consist in the attack of a (cyano)acetylenic radical on the other, intact precursor molecule.^{37,39} Consequently, the following scheme may be at work once $\text{C}_5\text{N}^{\bullet}$ becomes available:



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3 Although the above scheme is aimed at explaining the reactions occurring in cryogenic matrices,
4 not in the interstellar medium, it is nevertheless in accordance with what Cherchneff *et al.*
5 proposed for monocyanopolyacetylene formation in the circumstellar shell IRC +10 216.^{60,61}
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10 The propyne photolysis may also take place, and then the reactions of $\text{CH}_3\text{C}_2^\bullet$ could include:



17 Considering that the detection of products relied on phosphorescence, and that the applied
18 excitation laser beam, not harmful to HC_7N , efficiently photolyzed $\text{CH}_3\text{C}_7\text{N}$, it was not possible
19 to propose even a rough estimation of relative yields for the reaction channels leading to the two
20 discussed triacetylenic molecules. In this context, it is of interest to note, that our introductory
21 experiments with a matrix-isolated, photolyzed 1:1 mixture of $\text{CH}_3\text{C}_2\text{H}$ and HC_3N led to
22 approximately equally intense emissions from the two cyanodiacetylenes: $\text{CH}_3\text{C}_5\text{N}$ and HC_5N . In
23 this case, however, both photoproducts were stable during phosphorescence measurements.
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34 The attempted reactions $\text{CH}_3\text{C}_5\text{N} + \text{C}_2\text{H}_2$ and $\text{CH}_3\text{C}_3\text{N} + \text{C}_2\text{H}_2$ did not yield any evidence for the
35 elongation of a carbon chain. This suggest that it is easier to detach the methyl group from
36 propyne (Eq. 1) than from a methylcyano(poly)yne molecule.
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42 The discussed processes could possibly occur with a precursor molecule electronically excited
43 rather than converted into a free radical (i.e. the separate dissociation step might not be
44 distinguishable).
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53 CONCLUSIONS

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3 CH₃C₅N and CH₃C₇N molecules were successfully formed from respective smaller
4 (cyano)acetylenic precursors in UV-stimulated cryogenic processes. Phosphorescence of
5 CH₃C₇N, observed here for the first time, was strong enough to uncover the trace amounts of this
6 compound, below the detection level for IR absorption. HC₇N and CH₃C₇N (both products appear
7 simultaneously) turned out to have almost coincident $\alpha - \bar{X}$ origin wavelengths, as expected for
8 large carbon-nitrogen chain species where the presence or absence of a terminating methyl group
9 does not strongly affect the chromophore. Present studies provide good prospects for using the
10 same approach to synthesize thus far unexplored CH₃C₉N molecule; phosphorescence emissions
11 of the tetraacetylenes HC₉N and CH₃C₉N are however expected to be even more similar than
12 those of their presently studied triacetylenic analogues.
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27 Previous investigations of similar compounds^{37,38,40,57,62,63} were indicating the vibrational mode of
28 the highest Raman activity as the one responsible for the leading progression in phosphorescence
29 spectra. This trend seems to be preserved for HC₇N and CH₃C₇N.
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35 The resemblance of HC₇N and CH₃C₇N chromophores is well marked in their singlet-singlet
36 transitions. As usual for the homologous series of chain compounds, the HOMO-LUMO (either π
37 $- \pi^*$ or $e - e^*$) energy gap decreases with the molecular length.⁶⁴ Two main electronic systems
38 could be distinguished: a fully allowed one and a strictly (for HC₇N) or effectively (for CH₃C₇N)
39 forbidden one. The forbidden $\bar{B} - \bar{X}$ transitions ($\Delta - \Sigma$ and $E - A_1$, for HC₇N and CH₃C₇N,
40 respectively) were rendered detectable through coupling with a chain-bending vibration of the
41 C₇N chromophore. This bending, further coupled to a triple-bond stretching mode, promoted the
42 main vibronic progression, recognized in the phosphorescence excitation spectrum.
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3 SUPPORTING INFORMATION
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6 Geometry of HC₇N and CH₃C₇N in several electronic states; visualization of vibrational modes in
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8 selected electronic states of HC₇N and CH₃C₇N; phosphorescence spectra and newly observed
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10 band assignment for HC₅N.
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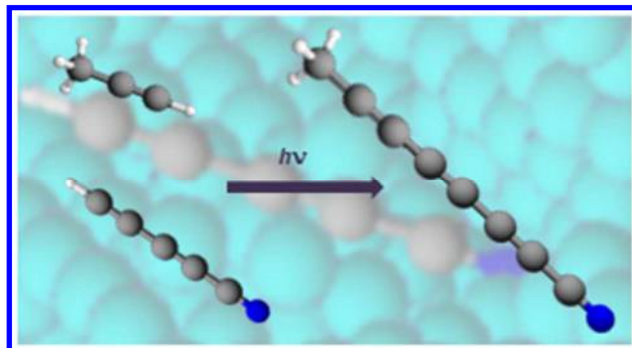
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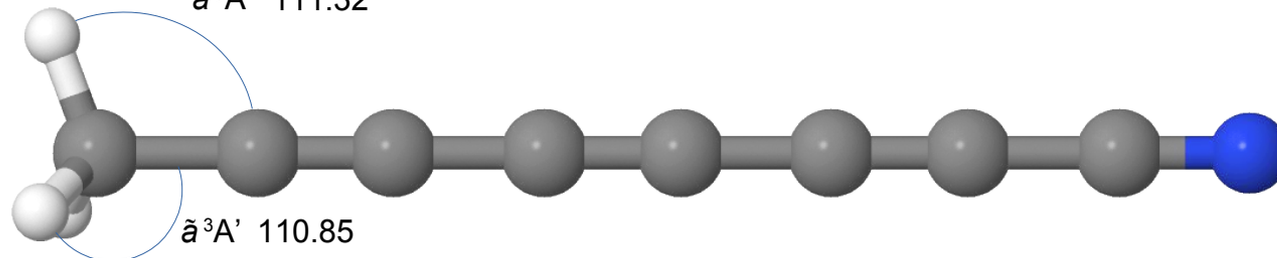
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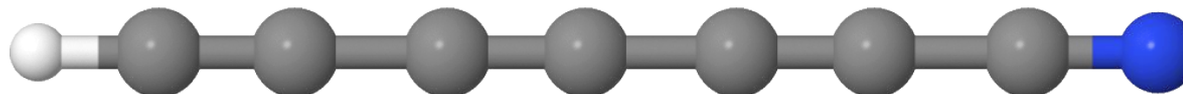
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TOC Graphic

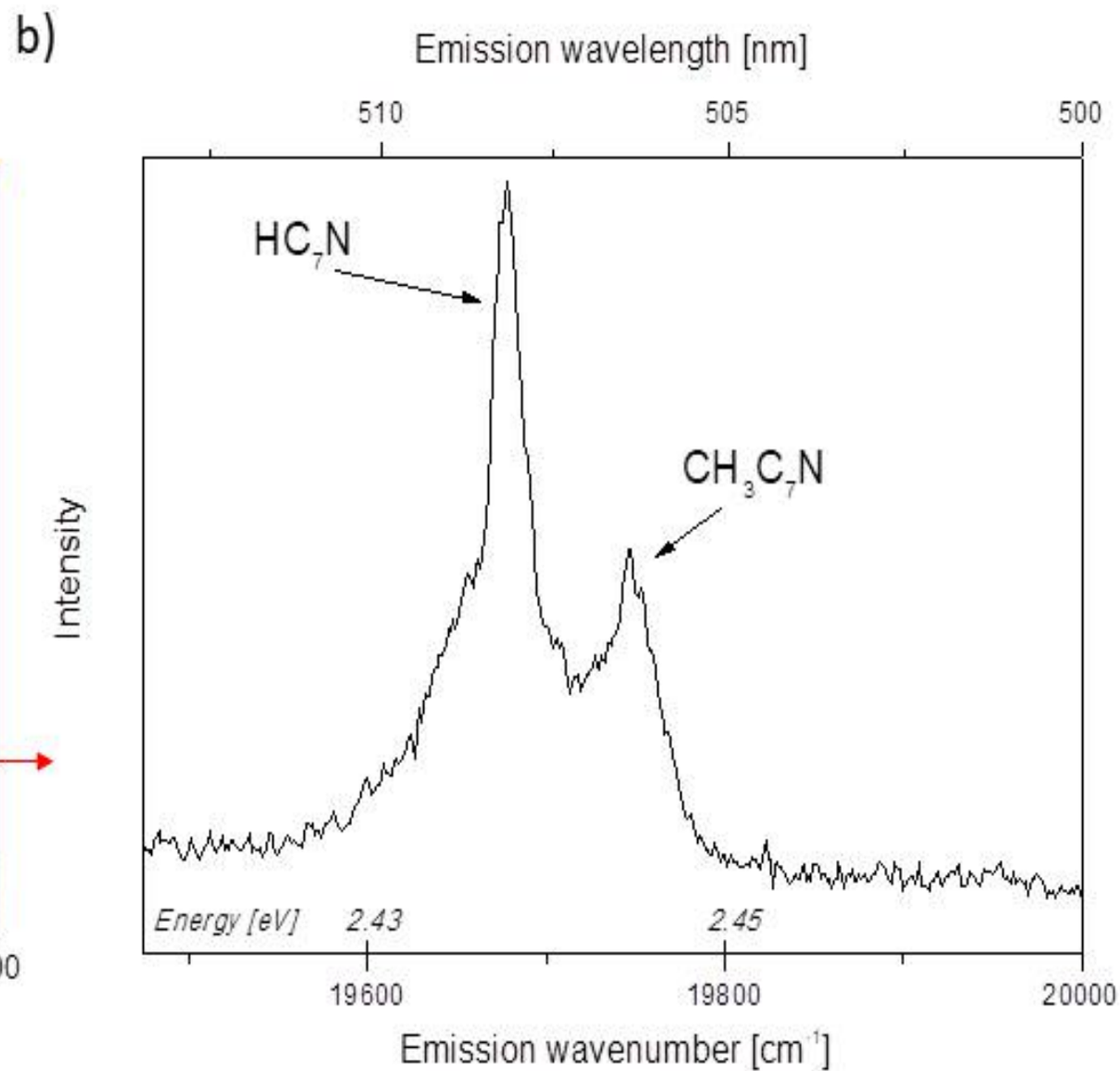
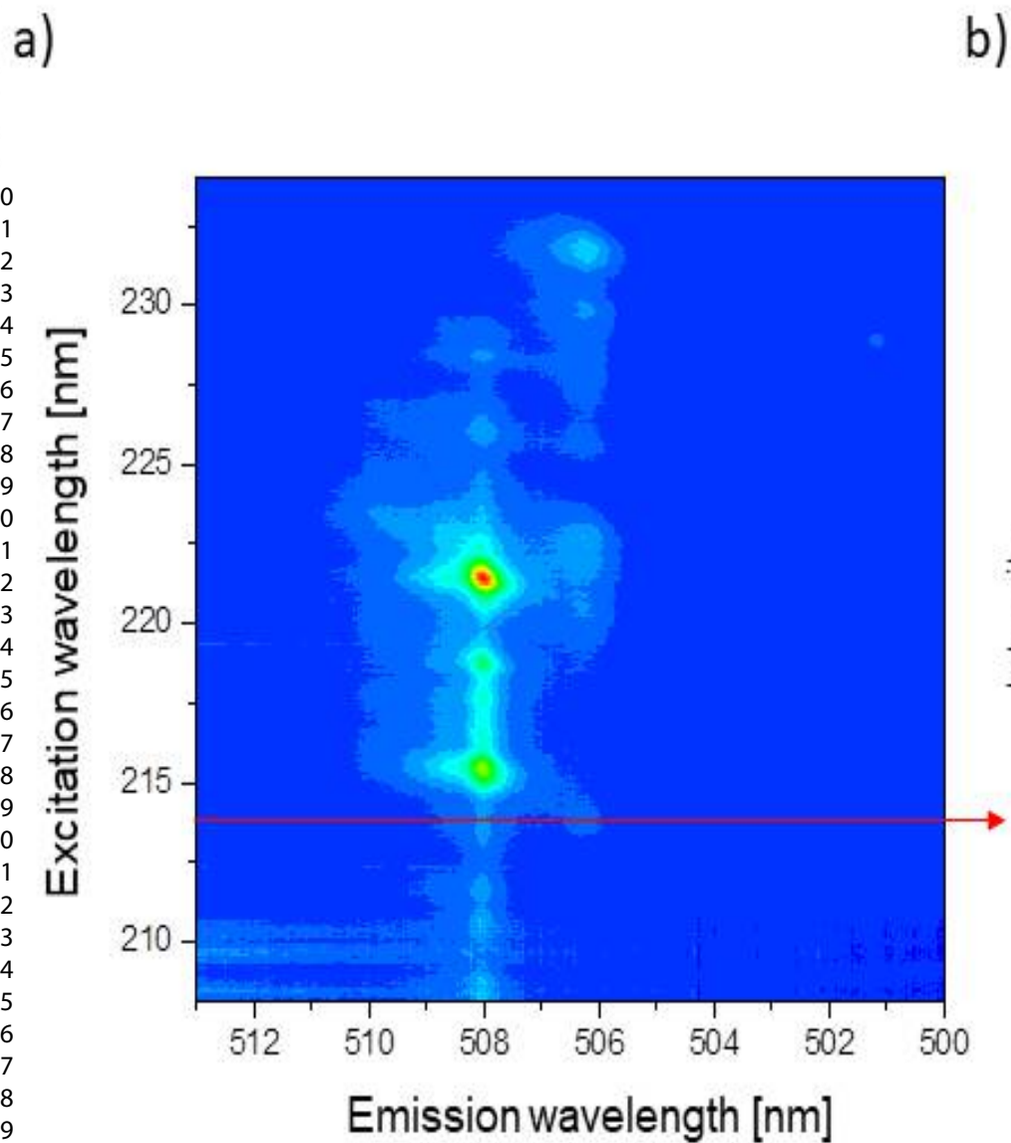


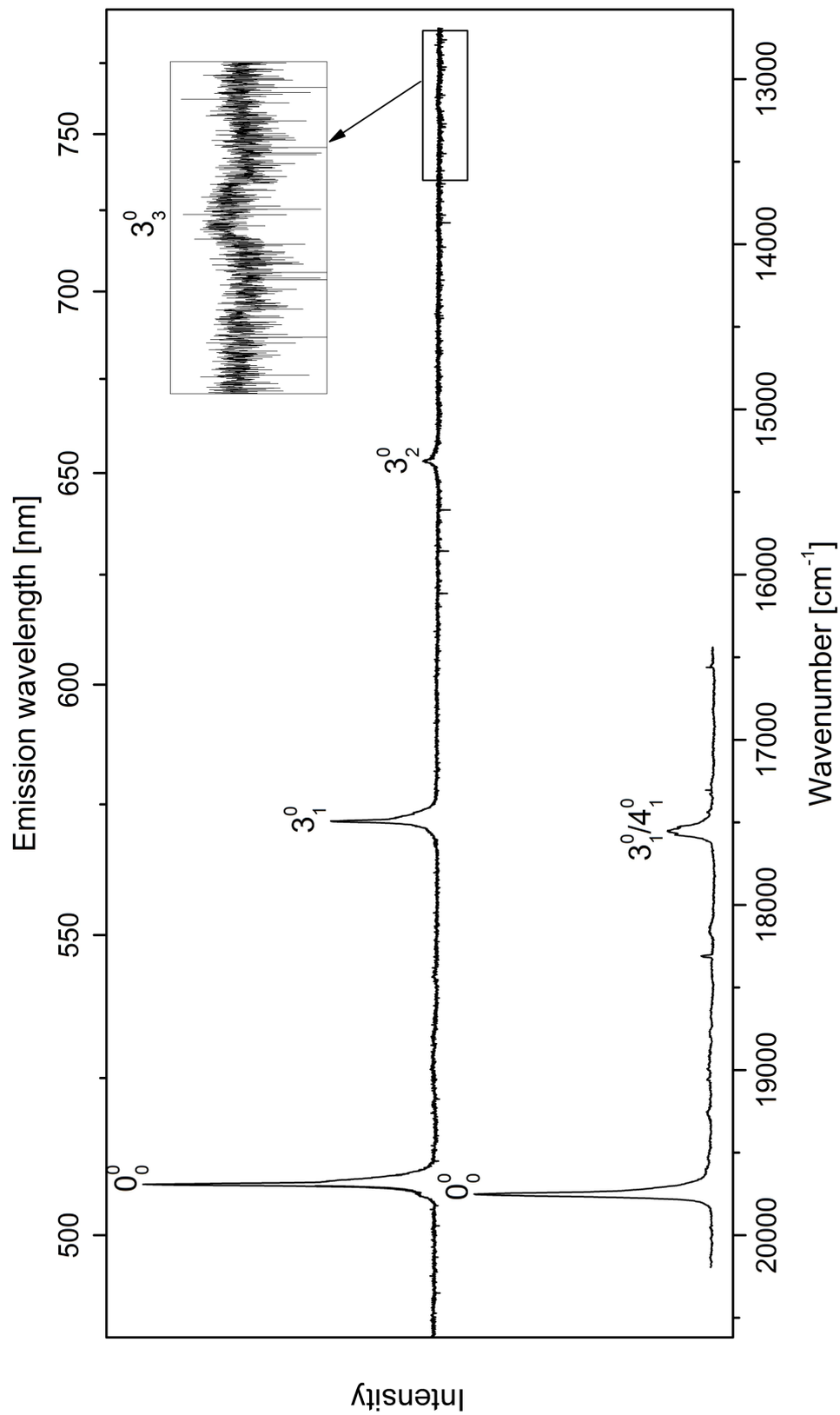
\tilde{X}^1A_1 110.77 \tilde{B}^1E 111.10 \tilde{E}^1A_1 111.28 \tilde{a}^3A' 111.32

\tilde{X}^1A_1	109.3	144.3	121.2	134.8	122.0	134.2	121.7	135.7	115.9
\tilde{B}^1E	+0.2	-0.8	+2.7	-4.3	+5.3	-5.5	+4.7	-3.2	+1.8
\tilde{E}^1A_1	+0.6	-1.5	+2.3	+0.1	+1.5	+0.4	+2.2	-1.3	+2.0
\tilde{a}^3A'	+0.2 +0.1	-0.5	+2.9	-4.7	+6.1	-6.4	+5.4	-3.4	+2.0



$\tilde{X}^1\Sigma^+$	1.0570(4)	1.2101(6)	1.3610(9)	1.1241(19)	1.3616(27)	1.2149(20)	1.3657(9)	1.1611(6)
	106.4	120.8	135.0	121.8	134.3	121.6	135.7	115.9
$\tilde{B}^1\Delta$	0.0	+2.7	-4.3	+5.5	-5.7	+4.9	-3.2	+1.8
$\tilde{E}^1\Sigma^+$	+0.3	+2.4	-0.6	+2.0	0.0	+2.4	-1.3	+2.0
$\tilde{a}^3\Sigma^+$	0.0	+3.0	-4.8	+6.3	-6.5	+5.4	-3.3	+1.8





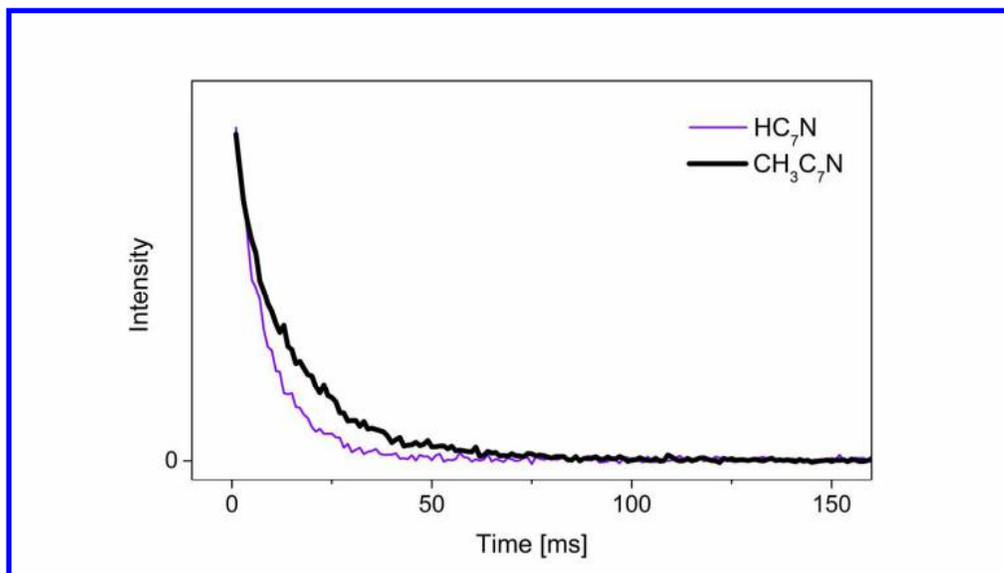
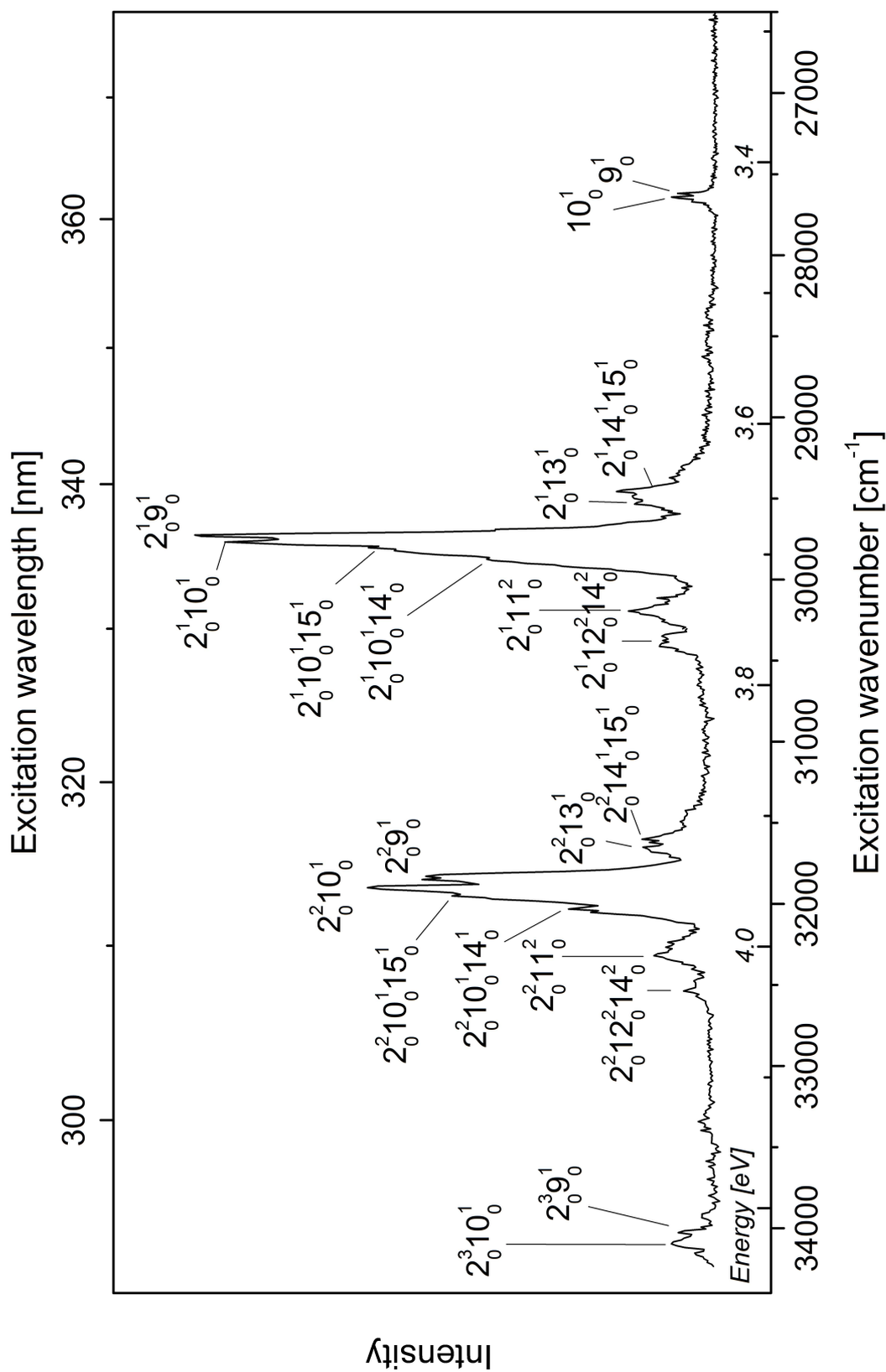


Figure 4.

199x112mm (300 x 300 DPI)



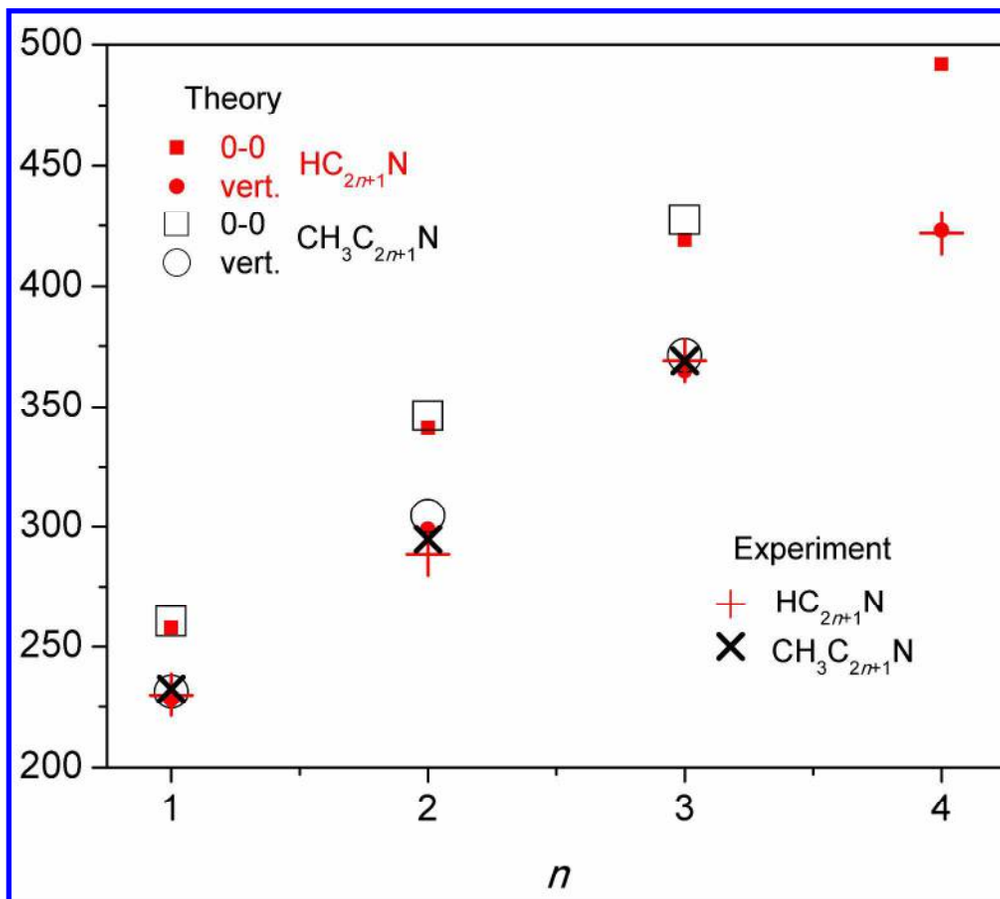


Figure 6.

289x257mm (300 x 300 DPI)

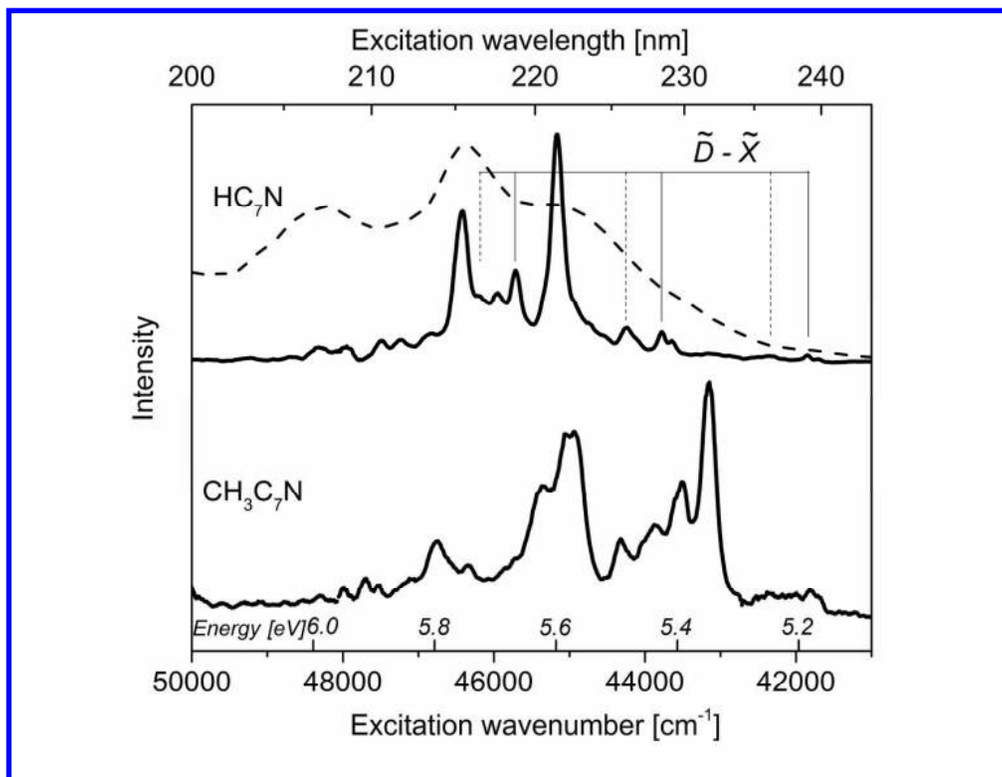


Figure 7.

208x159mm (300 x 300 DPI)

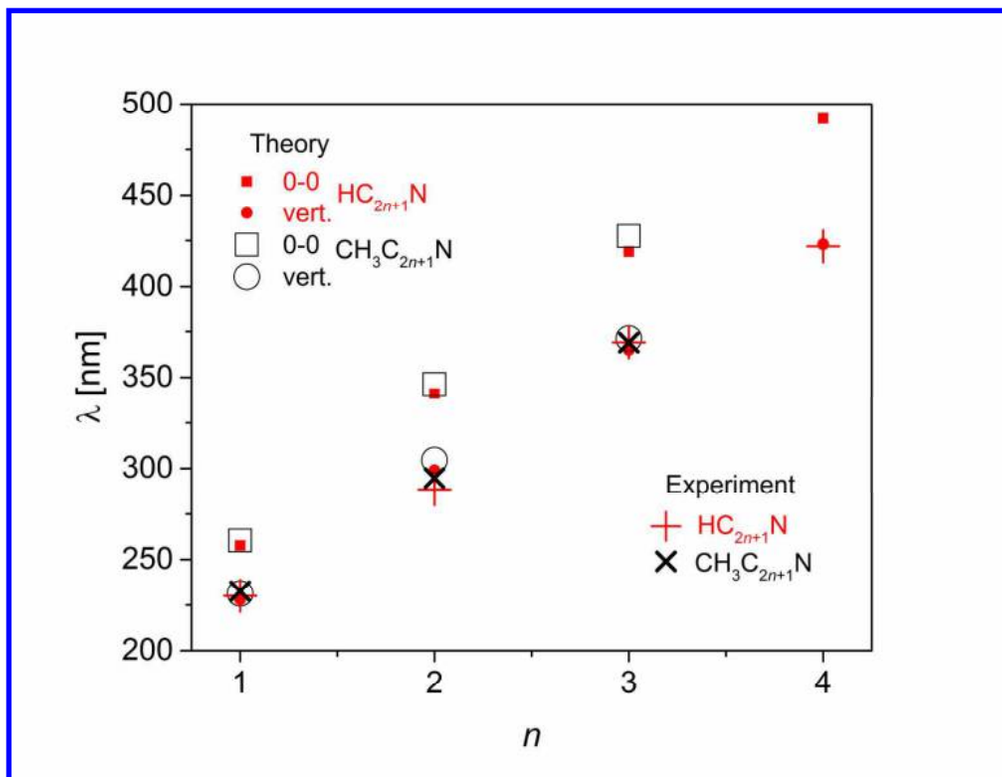


Figure 6.

208x159mm (300 x 300 DPI)