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1 Introduction

Due to their reactive nature, unsaturated linear hydrocarbons, such as acetylene and polyynes (general structure $\mathrm{HC}_{2n}\mathrm{H}; \, \mathrm{X}^1 \mathrm{\Sigma_g}^{+}) ,$ are prevalent in combustion chemistry, $1-4$ plasma processes, $5,6$ chemical synthesis,⁷⁻¹⁰ chemistry of planetary atmospheres,¹¹⁻¹⁹ and interstellar gas-phase chemistry.²⁰⁻²⁵ They are of particular interest for astronomers, because they are believed to act as the ultraviolet (UV) shield in hydrocarbon-rich atmospheres, $26-29$ and in the formation and destruction of polycyclic aromatic hydrocarbons (PAHs), $18,30-33$ a major reservoir of carbon in the universe. In astronomical environments, the formation of long chain polyynes from acetylene is believed to occur through polymerization reactions,19,34,35

 $HC_{2n}H + C_2H \rightarrow HC_{2n+2}H + H,$ (1a)

$$
HC_2H + C_{2n}H \rightarrow HC_{2n+2}H + H, \qquad (1b)
$$

 $HC_{2n}H^{+} + HC_{2}H \rightarrow HC_{2n+2}H_{2}^{+} + H,$ (2a)

$$
HC_{2n+2}H_{2}^{+} + e^{-} \rightarrow HC_{2n+2}H + H. \tag{2b}
$$

Theoretical investigation of the infrared spectrum of small polyynes†

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The full cubic and semidiagonal quartic force fields of acetylene (C_2H_2) , diacetylene (C_4H_2) , triacetylene (C_6H_2) , and tetraacetylene (C_8H_2) are determined using CCSD(T) (coupled cluster theory with single and double excitations and augmented by a perturbative treatment of triple excitations) in combination with the atomic natural orbital (ANO) basis sets. Application of second-order vibrational perturbation theory (VPT2) results in vibrational frequencies that agree well with the known fundamental and combination band experimental frequencies of acetylene, diacetylene, and triacetylene (average discrepancies are less than 10 cm⁻¹). Furthermore, the predicted ground state rotational constants (B_0) and vibration–rotation interaction constants (x_i) are shown to be consistent with known experimental values. New vibrational frequencies and rotational parameters from the presented theoretical predictions are given for triacetylene and tetraacetylene, which can be used to aid laboratory and astronomical spectroscopic searches for characteristic transitions of these molecules. PAPER

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Although long carbon chain molecules (e.g., HC_n and HC_nN for $n \leq 9$ ^{36–39} and small polymes (HC_{2n}H for $n \leq 3$) have been detected in carbon-rich astronomical sources, $20,21,39$ tetraacetylene has yet to be observed. One limiting factor is that as centrosymmetric molecules, polyynes lack a permanent dipole moment, and cannot be detected by radioastronomy using pure rotational transitions, unlike, $e.g., \text{ HC}_nN$. Therefore, ro-vibrational spectra in the infrared (IR) region are the most important spectroscopic tools to detect polyynes both in the laboratory and in space. In particular, detection of acetylene, diacetylene, and triacetylene in planetary atmospheres and protoplanetary nebulae has been realized primarily through observation of the strongest perpendicular band (ν ₅, ν ₈, and ν ₁₁, respectively, at \sim 13–17 µm) and the second strongest parallel band ($\nu_4 + \nu_5$, $\nu_6 + \nu_8$, and $\nu_8 + \nu_{11}$, respectively, at \sim 8 µm).^{11,12,20,21} However, accurate line positions for tetraacetylene are lacking, from either laboratory or theoretical studies.

Extensive theoretical and experimental studies have been carried out for acetylene and diacetylene in the past few decades, including high-resolution spectroscopic studies of all the fundamental bands and a significant number of the combination bands, $29,40-50$ and high level *ab initio* calculations that take into account anharmonic effects. $51-54$ The combination of these studies shows that current quantum chemical theory, particularly coupled cluster theory with single and double excitations and augmented by a perturbative treatment of triple excitations $(CCSD(T))$,⁵⁵ is able to accurately reproduce equilibrium geometries, experimental vibrational frequencies, vibration–rotation interaction constants (α_i) , and ground state rotational constants (B_0) .

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Triacetylene and tetraacetylene are not as thoroughly studied, notably in terms of rotational information. While all of the fundamental vibrational modes of triacetylene have been measured, there is only rotational information for the IR active fundamental modes,⁵⁶ and the strongest IR combination band $(\nu_8 + \nu_{11})$.^{57–61} However, theoretical studies of triacetylene do give rotational information for the remaining modes from CCSD(T) calculations of the vibration–rotation interaction constants⁶² and the equilibrium geometry.⁶³ In addition, the harmonic frequencies of triacetylene were calculated using partial fourth-order many-body perturbation theory $[SDQ-MBPT(4)]^{63}$ Conversely though, to the authors' knowledge, there is almost no rotational information for tetraacetylene. There has been only one low-resolution spectroscopic study of tetraacetylene, which measured three of the fundamentals (ν_6 , ν_8 , and ν_{14} at 3329.4, 2023.3, and 621.5 cm^{-1} , respectively), and one combination band $(\nu_{10} + \nu_{14}$ at 1229.7 cm⁻¹), and gives an estimate for the electronic ground state rotational constant, B_0 .⁶⁴ Unfortunately, the theoretical knowledge of tetraacetylene is equally limited, with only two studies of the equilibrium geometry (at the Hartree–Fock⁶⁵ and B3LYP⁶⁶ level of theory), and a calculation of the harmonic vibrational frequencies at the SVWN level of theory.⁶⁴ While the two modes that are most useful for astronomical identification $(\nu_{14}$ and $\nu_{10} + \nu_{14})$ were measured, the uncertainty associated with the line positions is too large to allow for an unambiguous assignment. Moreover, some high-resolution IR searches have been attempted,^{50,61,62,67} but so far no transitions have been assigned to tetraacetylene. Paper

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In this paper, we report the ab initio calculations for acetylene, diacetylene, triacetylene, and tetraacetylene. Due to the centrosymmetric nature of these molecules, observations in the laboratory and in space are most easily accomplished through their infrared spectra. As such, the properties computed and presented here are those related to that technique: fundamental vibrational frequencies, ground state rotational constants, and intramolecular interactions. The computational approach is calibrated using the well studied acetylene and diacetylene, and then extended to make predictions for triacetylene and tetraacetylene.

2 Computational methods

All calculations were carried out at the CCSD(T) level of theory, which with a sufficiently large basis set has been shown to accurately reproduce experimental values of semi-rigid molecules.52–55,68–72 Equilibrium geometries were determined using the large core-valence correlation-consistent quadruple- ζ basis set (cc-pCVQZ), which features [8s7p5d3f1g] (nonhydrogen atoms) and [4s3p2d1f] (hydrogen) of (15s9p5d3f1g) and (6s3p2d1f) primitive basis sets, respectively.⁷³⁻⁷⁵ All electron (AE)-CCSD(T)/cc-pCVQZ has been shown to give very accurate equilibrium geometries for unsaturated hydrocarbons.^{54,76-78} Optimizations were done using analytic energy derivatives,79 and were considered converged when the root-mean-square (RMS) gradient fell below 10^{-10} au.

However, it is well known that correlation-consistent basis sets, such as cc-pCVQZ, tend to underestimate the vibrational frequencies of symmetric bending modes (π_{ϱ}) of conjugated molecules, e.g., polyynes, due to their susceptibility to an intramolecular variant of basis set superposition error (BSSE).^{54,80} It has been shown that one way to avoid this problem is to use basis sets with a large number of Gaussian primitives (particularly f-type), such as the atomic natural orbital (ANO) basis set (with the primitive basis set (13s8p6d4f2g) for non-hydrogen atoms and (8s6p4d2f) for hydrogen).^{52,81,82} The basis set has two common truncations: [4s3p2d1f] for nonhydrogen atoms and [4s2p1d] for hydrogen (hereafter known as ANO1), and [5s4p3d2f1g] (non-hydrogen atoms) and [4s3p2d1f] (hydrogen) (hereafter known as $ANO2$).^{74,75,81} In addition, only the valence electrons of carbon are considered in the correlation treatment, i.e., standard frozen-core (fc) calculations. (fc)-CCSD(T)/ ANO1 has been shown to accurately reproduce experimental frequencies and intensities for small molecules.^{52,83,84} Using the (fc)-CCSD(T)/ANO1 optimized geometry, second-order vibrational perturbation (VPT2) theory calculations were determined from full cubic and the semidiagonal part of the quartic force fields obtained by numerical differentiation of analytic CCSD(T) second derivatives.^{70,85} All calculations were performed with the development version of the CFOUR program.⁸⁶

3 Results and discussion

3.1 Equilibrium structure

The AE-CCSD(T)/cc-pCVQZ equilibrium geometries are shown in Fig. 1, with comparison to experimentally derived values (in italics) when known. $53,87,88$ The theoretical equilibrium bond lengths for acetylene, diacetylene, and triacetylene all agree within 0.5% of the structures determined from experimentally measured rotational constants. As the length of the carbon chain increases, the C–H bond lengths stay essentially the same, \sim 1.062 Å, consistent with a sp-H type C–H bond. However, the $C\equiv C$ bond lengths increase (particularly the internal $C \equiv C$ bonds), while the C–C bond lengths decrease, becoming closer to that typical of CC double bonds. This suggests that the π electrons become more delocalized over the internuclear axis, and the polyyne's configuration moves from a strict triple-single bond alternation to more of a consecutive

Fig. 1 AE-CCSD(T)/cc-pCVQZ equilibrium geometries (Å) for $HC_{2n}H$ Experimentally determined equilibrium bond lengths for acetylene, 87 diacetylene, 53 and triacetylene 88 are given in italics below.

$$
B_0=B_{\rm e}-\frac{1}{2}\sum_i\alpha_i,\qquad \qquad (3)
$$

3.2 Spectroscopic properties of acetylene and diacetylene

The quality of the present calculations is checked by comparison to the experimentally well studied acetylene and diacetylene. The harmonic and VPT2 fundamental frequencies of the fundamental and combination bands are given in Table 2 and 3 for acetylene and diacetylene, respectively, and experimental values are included for comparison. The (fc)-CCSD(T)/ANO1 VPT2 fundamental frequencies show good agreement with experimental values, with most observed–calculated deviations (o–c) being less than 5 cm^{-1} and all being less than 15 $\mathrm{cm}^{-1}.$

Based on previous studies of acetylene⁵² and diacetylene,⁵³ the use of the ANO2 basis set was evaluated compared to the ANO1 basis set. For some of the vibrational modes, such as the ν_4 mode of acetylene [612.88 cm $^{-1}$ (observed)], 42 Martin *et al.* 52 showed that CCSD(T)/ANO2 can give a slightly better agreement (o-c value of \sim 2 cm⁻¹) compared to the ANO1 basis set (o-c value of \sim 12 cm⁻¹). However, the study by Thorwirth *et al.*⁵³

Table 2 Harmonic and anharmonic (VPT2) frequencies (in cm⁻¹) of acetylene fundamental and selected combination bands

	$CCSD(T)/ANO1^a$		Experimental
	ω	$\boldsymbol{\nu}$	ν
$\nu_1(\sigma_{\rm g}^+)$ $\nu_2(\sigma_{\rm g}^+)$ $\nu_3(\sigma_{\rm u}^{+})$ $\nu_4(\pi_{\rm g})$ $\nu_5(\pi_{\rm u})$ $\nu_4 + \nu_5(\sigma_u^+)$ $\nu_2 + \nu_5(\pi_{\rm u})$	3514.2(0) 2001.5(0) 3414.6(84.7) 600.5(0) 752.3(90.5) 1352.8 2753.8	3375.2(0) 1964.8(0) 3285.9(74.8) 600.6(0) 734.7(91.7) 1329.2(10.8) 2698.3(0.1)	3372.85141 1974.317 ⁴¹ 3288.5807548 612.87142 730.33242 1328.07442 2701.90743
$\nu_3 + \nu_4(\pi_{\rm u})$ $\nu_1 + \nu_5(\pi_u)$ $\nu_1 + \nu_3(\sigma_u^+)$ Anharmonic ZPE = 5760.1	4015.1 4266.5 6928.7	3878.5(0.5) 4098.9(0.5) 6551.9(2.0)	3882.4060 ⁴¹ 4091.17326 ⁹¹ 6556.46 ⁴⁰

Table 3 Harmonic and anharmonic (VPT2) frequencies (in cm⁻¹) of diacetylene fundamental and selected combination bands

PCCP	Table 1 $CCSD(T)/ANO1$ rotational constants (in cm^{-1}) of acetylene, diacetylene, triacetylene, and tetraacetylene						Table 2 Harmonic and anharmonic (VPT2) frequencies (in cm^{-1}) of acetylene fundamental and selected combination bands	Paper
	HC ₂ H	HC_4H	HC_6H	HC_8H		$CCSD(T)/ANO1^u$		Experimental
Calc.						ω	ν	ν
$B_{\rm e}$ B_0 $D_e({\times}10^8)$ 160	1.181053 1.175319	0.146248 0.146167 1.5	0.044064 0.044092 0.086	0.018823 0.018844 0.012	$\nu_1(\sigma_g^+)$ $\nu_2(\sigma_{\rm g}$) $\nu_3(\sigma_{\rm u}^+)$	3514.2(0) 2001.5(0) 3414.6(84.7)	3375.2(0) 1964.8(0) 3285.9(74.8)	$3372.851^{41}\,$ 1974.317 ⁴¹ 3288.5807548
Expt. $B_{\rm 0}$	$\frac{1.17664632(18)^{90}}{1.56825(20)^{29}}\ \frac{0.1464123(17)^{50}}{0.0441735(12)^{61}}\ \frac{0.020(3)^{64}}{0.107(7)^{61}}$ $D_0(\times 10^8)$ 159.8(9) ⁹⁰				$\nu_4(\pi_{\rm g})$ $\nu_5(\pi_\mathrm{u})$ $\nu_4 + \nu_5(\sigma_u^+)$ $\nu_2 + \nu_5(\pi_u)$	600.5(0) 752.3(90.5) 1352.8 2753.8	600.6(0) 734.7(91.7) 1329.2(10.8) 2698.3(0.1)	612.87142 730.33242 1328.07442 2701.90743
	double bond character of the CC bonds, making the overall				$\nu_3 + \nu_4(\pi_u)$ $\nu_1 + \nu_5(\pi_u)$ $\nu_1 + \nu_3(\sigma_{\rm u})$ Anharmonic ZPE = 5760.1	4015.1 4266.5 6928.7	3878.5(0.5) 4098.9(0.5) 6551.9(2.0)	3882.4060 ⁴¹ 4091.1732691 $6556.46^{40}\,$
	structure more rigid as C_2 units are added, an effect that also qualitatively acts to increase the biradical character of the molecule as the size grows. The equilibrium rotational constants, B_e , obtained from the						a Intensities in km mol ⁻¹ are given in parentheses. Table 3 Harmonic and anharmonic (VPT2) frequencies (in cm^{-1}) of	
	AE-CCSD(T)/cc-pCVQZ equilibrium geometries are summarized in Table 1, and agree well with experimental ground state					$CCSD(T)/ANO1^a$	diacetylene fundamental and selected combination bands	Experimental
	rotational constants (B_0) . As such, the equilibrium rotational						ν	ν
	constants suggest that the calculations predict the correct					ω		
	ground state geometry, because for linear molecules with more				$\nu_1(\sigma_g)$ $\nu_2(\sigma_{\rm g}^+)$	3465.8(0) 2240.2(0)	3332.5(0) 2193.1(0)	3332.1547646 2188.928544
	than three atoms the summation of vibration-rotation inter-				$\nu_3(\sigma_g^+)$	891.1(0)	859.2(0)	871.9582^{44}
	action constants (α_i) is expected to be close to zero, and from				$\nu_4(\sigma_{\mathrm{u}}^+)$	3465.9(152.7)	3333.1(135.5)	3333.663450
					$\nu_5(\sigma_{\mathrm{u}}^+)$	2054.1(0.2) 636.3(0)	2016.9(0.5) 624.2(0)	2022.241544 625.643507 ²⁹
		$B_0 = B_e - \frac{1}{2} \sum_i \alpha_i,$		(3)	$\nu_6(\pi_{\rm g})$ $\nu_7(\pi_{\rm g})$	479.8(0)	476.9(0)	482.707844
					$\nu_8(\pi_\mathrm{u})$	636.3(78.7)	624.1(78.8)	628.040776 ²⁹
$B_0 \sim B_e$.					$\nu_9(\pi_\mathrm{u})$	220.7(7.3)	219.6(7.3)	219.97713^{47} 438.4775747
	In addition, as seen for other carbon chains (e.g., HC_n ,				$2\nu_9(\sigma_g^+)$ $\nu_7 + \nu_9 (\sigma_{\rm u}^+)$	441.4 700.5	438.5(0) 696.3(0.8)	701.8939 ²⁹
	$HC_{2n+1}N$, and H_2C_n ⁸⁹ the centrifugal distortion constant (D_e)				$\nu_6 + \nu_9 (\sigma_u^+$	857.0	843.9(0.01)	845.65551329
	decreases with increasing molecular size, with a theoretical				$\nu_8 + \nu_9 (\sigma_g)$	857.0	843.9(0)	848.365918 ²⁹ $\mathbf{1111}^{45}$
	D_e = 1.6 \times 10 ⁻⁶ cm ⁻¹ for acetylene, D_e = 1.5 \times 10 ⁻⁸ cm ⁻¹ for				$\nu_7 + \nu_8(\pi_u)$ $\nu_6 + \nu_8(\sigma_{\rm u}^+)$	1116.1 1272.6	1103.1(0.6) 1244.7(21.8)	1241.06082846
	diacetylene, $D_e = 8.6 \times 10^{-10}$ cm ⁻¹ for triacetylene, and				$2\nu_6 + \nu_8(\pi_{\rm u})$	1909.0	1864.6(0.0)	1863.251244
	D_e = 1.2 × 10 ⁻¹⁰ cm ⁻¹ for tetraacetylene. These values are				$\nu_2 + \nu_9(\pi_u)$	2460.9	2410.0(0.04)	2406.425144
	consistent with those found experimentally for the respective				$\nu_5 + \nu_7(\pi_u)$ $\nu_5 + \nu_6(\pi_u)$	2533.9 2690.4	2489.0(0.01) 2637.0(0.04)	2500.645844 2643.32323^{46}
	vibrational ground states (Table 1). As noted by Thaddeus et al. ⁸⁹				$\nu_2 + \nu_8(\pi_u)$	2876.6	2810.9(0.4)	2805^{45}
	this behavior of increasing stiffness with chain length is a				$\nu_1 + \nu_9(\pi_{\rm u})$	3686.5	3551.6(0.1)	3551.5615815946
	distinguishing characteristic associated with bona fide chains.				$\nu_1 + \nu_8(\pi_u)$ $\nu_4 + \nu_6(\pi_u)$	4102.1 4102.3	3946.9(0.7) 3947.8(0.7)	393945
					$\nu_2 + \nu_5(\sigma_{\rm u})$	4294.3	4194.0(0.1)	
	Spectroscopic properties of acetylene and diacetylene				$\nu_4 + \nu_3(\sigma_{\rm u})$	4357.0	4192.3(0.1)	
3.2	The quality of the present calculations is checked by comparison				$\nu_1 + \nu_4 (\sigma_u^+)$ Anharmonic ZPE = 7966.9	6931.7	6557.2(3.4)	6565.47249

 a Intensities in km mol⁻¹ are given in parentheses.

showed that, for diacetylene, the average o–c value with $CCSD(T)/ANO2$ is comparable to that for the ANO1 basis set (\sim 6 cm⁻¹ and \sim 4 cm⁻¹, respectively). Moreover, the time cost of (fc)-CCSD(T)/ANO2 calculations compared to (fc)-CCSD(T)/ ANO1 far outweighs the minor frequency differences, and does not justify the higher computational cost of the ANO2 basis set in predicting the fundamental frequencies of longer polyynes.

The (fc)-CCSD(T)/ANO1 anharmonicity constants (x_{ii}, ESI) also accurately account for the known combination bands of acetylene and diacetylene (Tables 2 and 3, respectively). All the combination bands are within 5 cm^{-1} of their observed values. For both acetylene and diacetylene, the ANO1 basis set is able

3.3 Spectroscopic properties of triacetylene

Table 4 CCSD(T)/ANO1 vibration–rotation interaction constants (α_i , in cm $^{-1}$) of acetylene, diacetylene, triacetylene, and tetraacetylene. Experimental values are in parentheses

	HC ₂ H Mode $(\times 10^3)$	HC_4H $(\times 10^4)$	HC_6H $(\times 10^5)$	HC_8H $(\times 10^5)$
α_{1}	$6.853(6.904^a)^{51}$	$2.157(2.153)^{50}$	2.97	0.730
α_2	$6.007(6.181)^{51}$	6.608	15.20	4.91
α_3	$5.800(5.882^{a})^{51}$	$3.123(3.110^a)^{54}$	7.44	2.55
α_4	$-1.464(-1.354)^{51}$	$2.139(2.183)^{50}$	3.82	3.76
α_5	$-2.134(-2.232)^{51}$	$3.938(3.948)^{44}$	$2.99(3.58)^{61}$	0.930
α_6	0.730	$-0.700(-0.678)^{29}$	$9.91(9.15)^{58}$	0.730
α_{7}	4.06	$-2.703(-2.711)^{46}$	9.91	4.06
α_{8}	2.33	$-0.647(-0.636)^{29}$	$-1.17(-1.071)^{58}$	2.33
$\alpha_{\rm o}$	2.05	$-4.125(-4.183)^{46}$	-5.83	2.05
α_{10}			$-7.42(-7.88)^{58}$	-0.295
α_{11}			$-1.06(-1.06)^{57}$	-1.95
α_{12}			-5.07	-1.69
α_{13}			$-8.47(-8.7207)^{59}$	-2.26
α_{14}				-0.295
α_{15}				-0.163
α_{16}				-2.29
α_{17}				-2.80
	a Deperturbed.			

Table 5 Harmonic and anharmonic (VPT2) frequencies (in cm⁻¹) of triacetylene fundamental and selected combination bands

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	to most accurately reproduce the C-H asymmetric stretch mode		Table 5 Harmonic and anharmonic (VPT2) frequencies (in cm^{-1}) of		
	$(\nu_3$ and ν_4 , respectively). Significant is the agreement between the experimental and our predicted frequencies of $u_6 + u_8$		triacetylene fundamental and selected combination bands $CCSD(T)/ANO1^a$		Experimental
	$[1241.060828(38)$ cm ⁻¹ (observed) ⁴⁶ and 1244.7 cm ⁻¹ (theoretical)],				
	and $2\nu_6 + \nu_8$ [1863.2512(5) cm ⁻¹ (observed) ⁴⁴ and 1864.6 cm ⁻¹		ω	ν	ν
	(theoretical)] of diacetylene; both of which had only previously	$\nu_1(\sigma_g^+)$	3463.1(0)	3330.4(0)	331356
	been calculated with CCSD(T)/cc-pCVQZ, and had o-c values	$\nu_2(\sigma_{\rm g}^{-})$	2284.0(0)	2213.2(0)	2201^{56}
		$\nu_3(\sigma_g^+)$	2061.0(0)	2023.2(0)	201956 62556
	greater than 20 cm^{-1} . ⁵⁴ This suggests that the combination band	$\nu_4(\sigma_{\rm g}^+)$	616.1(0) 3463.1(126.4)	612.7(0) 3329.5(175.0)	3329.0533 ⁶¹
	VPT2 frequencies of polyynes determined using (fc)-CCSD(T)/	$\nu_5(\sigma_{\bf u}^+)$ $\nu_6(\sigma_{\rm u}^{~~\ast})$	2172.2(0.0)	2130.4(0.1)	2128.9163758
	ANO1 are accurate to aid identification of molecules, such as in	$\nu_7(\sigma_{\rm u}^+)$	1169.6(1.7)	1160.9(0.2)	1115.0^{59}
	astronomical surveys.	$\nu_8(\pi_g)$	633.0(0)	620.9(0)	622.3857
	The vibration-rotation interaction constants (Table 4) are	$\nu_9(\pi_{\rm g})$	489.5(0)	486.2(0)	49156
		$\nu_{\mathbf{10}}(\pi_{\mathbf{g}})$	252.0(0)	251.1(0)	258^{56}
	also determined in the course of the VPT2 calculation, and are	$\nu_{\mathbf{11}}(\pi_{\mathbf{u}})$	632.0(80.5)	619.9(83.2)	621.34011^{60}
	in good agreement with both previous theoretical studies ^{52,54}	$\nu_{12}(\pi_{\rm u})$	444.7(1.0)	441.8(1.0)	443.5^{59}
	and experimentally determined values. ^{29,44,46,50,51,54} Based on	$\nu_{13}(\pi_\mathrm{u})$	106.4(4.1)	105.9(3.5)	105.03861659
	the vibration-rotation interaction constants, the ground state	$\nu_9 + \nu_{13} (\sigma_{\rm u}^{\dagger})$	595.9	591.7(0.8)	
	rotational constants (B_0) were determined using the AE-CCSD(T)/	$\nu_{10} + \nu_{12} (\sigma_{\rm u}^+)$	696.7 1077.7	691.8(1.8) 1063.5(0.3)	
		$\nu_8 + \nu_{12} (\sigma_u^+)$ $\nu_9 + \nu_{11} (\sigma_{\rm u}^+)$	1121.5	1107.1(0.7)	
	cc-pCVQZ determined B_e values (Table 1). For acetylene,	$\nu_8 + \nu_{11} (\sigma_{\rm u}^+)$	1265.0	1237.4(31.4)	1232.90429558
	B_0 = 1.175319 cm ⁻¹ , which is a 0.1% difference compared to the	$\nu_3 + \nu_7(\sigma_{\rm u}^+)$	3230.5	3182.8(0.1)	
	experimentally determined value of $B_0 = 1.17664632(18)$ cm ^{-1.90}	$\nu_2 + \nu_7(\sigma_u^+)$	3453.6	3362.2(2.5)	
	Diacetylene shows a similar 0.2% difference between the	$3\nu_7(\sigma_{\rm u}^+)$	3508.7	3498.7(0.01)	
	theoretical value of B_0 = 0.146167 cm ⁻¹ , and the experimentally	$\nu_1 + \nu_{13}(\pi_u)$	3569.5	3436.1(0.2)	
		$\nu_5 + \nu_{10}(\pi_u)$	3715.1	3583.7(0.1)	
	determined value of $B_0 = 0.1464123(17)$ cm ^{-1.50} The consistent	$\nu_4 + \nu_5({\sigma_{\rm u}}^+)$	4079.3	3945.8(0.1)	
	accuracy of these values suggests that the method presented is	$\nu_1 + \nu_{11}(\pi_u)$	4095.2 4096.2	3940.1(0.8)	
	clearly good enough to be extrapolated to and aid high-resolution	$\nu_5 + \nu_8(\pi_u)$ $\nu_3 + \nu_6 (\sigma_{\rm u}$	4233.1	3943.9(0.8) 4141.1(0.1)	
	infrared spectroscopic searches for the larger polyynes.	$\nu_2 + \nu_6(\sigma_u^+)$	4456.2	4334.7(0.1)	
		$\nu_2 + \nu_5(\sigma_{\rm u}^+)$	5747.2	5548.4(0.2)	
	Spectroscopic properties of triacetylene 3.3	$\nu_1 + \nu_5(\sigma_u^+)$	6926.3	6555.1(4.6)	
		Anharmonic ZPE = 10095.8			
	The (fc)-CCSD(T)/ANO1 harmonic and VPT2 fundamental		a Intensities in km mol ^{-1} are given in parentheses.		
	frequencies along with the experimental frequencies are given				
This article is licensed under a Creative Commons Attribution 3.0 Unported Licence. Article. Published on 03 November 2017. Downloaded on 9/20/2024 5:40:47 PM.	in Table 5. Comparison between theoretical VPT2 frequencies				
	and experimental fundamentals measured with high-resolution		constants (x_{ii}, ESI) are able to reproduce the experimental value		
	techniques shows average o-c values that are smaller than		to within 5 cm^{-1} , suggesting other combination band frequencies		
Open Access	those seen for acetylene or diacetylene ($o-c \sim 2$ cm ⁻¹). For the	are of equal accuracy.			
l≿	known combination band, the (fc)-CCSD(T)/ANO1 anharmonicity		For the modes observed in low-resolution studies $(e.g.,)$		

For the modes observed in low-resolution studies (e.g., ν_1 and ν_{12}), the agreement is still good with o–c values less than 20 cm⁻¹. The notable exception is the internal $C \equiv C$ asymmetric stretch mode (ν_7) , which differs by 45 cm⁻¹. Since no rotationally resolved data can be found for this band, it is possible that the band observed at 1115.0 cm^{-1} ⁵⁹ was mis-assigned as the ν ₇ fundamental. A more likely assignment for this band is the $\nu_9 + \nu_{11}$ combination band, which has a predicted VPT2 frequency of 1107.1 cm^{-1} , a calculated intensity of 0.7 $km \, mol^{-1}$, and the same symmetry. Furthermore, the combination band is expected to be $3.5\times$ more intense than the ν_7 fundamental at 0.2 km mol⁻¹, suggesting that $\nu_9 + \nu_{11}$ is more likely of the two to be observed. However, rotationally resolved measurements of this band are clearly needed to confirm this speculation.

We note that, a resonance between the ν_5 fundamental and the $\nu_2 + \nu_7$ and $3\nu_7$ combination bands must be addressed to achieve the very small (1 cm^{-1}) o-c difference obtained for the C–H asymmetric stretch mode, ν ₅. The vibrational frequencies as a result of resonant interactions are calculated by a deperturbation-diagonalization technique followed by transformation of the deperturbed transition moments, as discussed

Table 6 Harmonic and anharmonic (VPT2) frequencies (in cm⁻¹) of tetraacetylene fundamental and selected combination bands

ω $\nu_1(\sigma_g^+)$ 3462.0(0) 2263.2(0) $\nu_2(\sigma_g)$ 2134.6(0) $\nu_3(\sigma_g)$ 1296.4(0) $\nu_4 {(\sigma_g}^{\scriptscriptstyle \top}$ 470.0(0) $\nu_5(\sigma_g^+)$ 3461.6(223.1) $\nu_6[\sigma_u]$ 2254.7(1.0) $\nu_7(\sigma_{\mathrm{u}}^+)$ 2064.3(0.3) $\nu_8(\sigma_u)$ 911.6(3.2) $\nu_9(\sigma_u^{-1})$ 632.4(0) $\nu_{\rm 10}(\pi_{\rm g})$	ν 3330.5(0) 2208.0(0) 2094.2(0) 1285.8(0) 455.3(0) 3328.8(214.2) 2227.6(0.5)	ν	Of the predicted fundamental and combination bands, there are a number of bands that are found/predicted to have sufficient
			intensity and/or relatively unique frequency range that could offer viable target transitions to use to search for tetraacetylene in future laboratory or astronomical spectra. For example, in the IR the
			ν_1 + ν_6 at 6550.8 cm ⁻¹ or ν_{12} + ν_{15} at 871.9 cm ⁻¹ combination
		3329.4^{64}	
			bands have both comparable predicted intensity to measured
	2026.6(0.6)	2023.364	bands of di- and triacetylene, and have transitions in relatively
	922.4(2.0)		clean regions of the spectrum. In terms of astronomical searches
	620.0(0)		
489.3(0) $\nu_{11}(\pi_{\rm g})$	486.0(0)		the ν_{17} mode at 60.7 cm ⁻¹ , offers a unique target transition, since
422.3(0) $\nu_{12}(\pi_{\mathrm{g}})$	419.5(0)		its low frequency makes it accessible by far-IR observations, similar
158.8(0) $\nu_{13}(\pi_{\rm g})$	157.9(0)		to the ν_2 bending mode of C_3 . ⁹²
632.6(79.6) $\nu_{14}(\pi_u)$	619.7(79.9)	621.5^{64}	
474.2(0.1) $\nu_{15}(\pi_{\rm u})$	470.9(0.2)		Based on the results discussed for the other small polyynes
267.7(3.2) $\nu_{16}(\pi_{\rm u})$	266.7(3.1)		the theoretical vibration-rotation interaction constants given in
61.0(2.3) $\nu_{17}(\pi_{\rm u})$	60.7(2.2)		
550.3 $\nu_{11} + \nu_{17} (\sigma_{\rm u}^+)$	546.5(0.6)		Table 4 are sufficient to assist in identification of ro-vibrational
$\nu_{13} + \nu_{15} (\sigma_{\rm u}^+)$ 633.0	628.5(1.7)		bands of tetraacetylene. The α_i results in a theoretical ground
$\nu_{12} + \nu_{16} (\sigma_{\rm u}^+)$ 690.1	684.8(3.1)		state rotational constant of $B_0 = 0.018844$ cm ⁻¹ that agrees
$\nu_{12} + \nu_{15} (\sigma_{\rm u}^+)$ 896.5	871.9(3.5)		
963.5 $\nu_{11} + \nu_{15} (\sigma_{\rm u}^{\dagger})$	970.5(1.3)		within errors with the experimentally determined value
$\nu_{10} + \nu_{15} (\sigma_{\rm u})$ 1106.6	1092.2(0.6)		B_0 = 0.020(3) cm ^{-1.64} Overall, for polymes the difference
$\nu_{10} + \nu_{14} (\sigma_{\rm u}^+)$ 1265.1	1236.7(37.5)	1229.7^{64}	between the experimental and calculated rotational constants
3174.8 $\nu_2 + \nu_9 (\sigma_u)$	3130.1(0.5)		
3360.7 $\nu_4 + \nu_8(\sigma_u)$	3311.4(0.5)		(ΔB_0) decreases from 0.001 to 0.00008 cm ⁻¹ as the chain length
4094.0 $\nu_6 + \nu_{10}(\pi_u)$	3939.1(0.8)		is increased, which is consistent with the trend seen for other
4094.6 $\nu_1 + \nu_{14}(\pi_u)$	3940.6(0.8)		carbon chain molecules (e.g., HC _n N, HC _n , C _n O). ⁹³ Therefore, is
6923.6 $\nu_1 + \nu_6(\sigma_u)$	6550.8(5.7)		
Anharmonic ZPE = 12218.2			the trend continues as expected then the ΔB_0 for tetraacetylene
			is equal to or smaller than that seen for triacetylene, and the
a Intensities in km mol ⁻¹ are given in parentheses.			determined ground state rotational constant is a good approxi-
			mation of the true value.

in the work of Vázquez and Stanton and Matthews et $al.^{85}$ This combination of Fermi and Darling–Dennison interactions shifts the ν_5 predicted frequency from 3333.1 to 3329.5 $\mathrm{cm}^{-1},$ which is able to reproduce the experimentally observed frequency $[3329.0533(2)$ $\text{cm}^{-161}]$ with the same accuracy seen for diacetylene (o -c ~ 0.5 cm⁻¹). The combination bands involved are similarly shifted: $\nu_2 + \nu_7$ from 3329.5 to 3362.2 cm $^{-1}$, and 3 ν_7 from 3526.7 to 3498.7 cm^{-1} . Since the shift is most pronounced for the two combination bands, future experimental work to observe either of these bands is required to confirm this prediction.

The vibration–rotation interaction constants for triacetylene are given in Table 4, and are consistent with the previous $CCSD(T)/cc$ -pCVQZ theoretical study⁶² and experimentally determined values.^{57-59,61} Consequently, the calculated ground state rotational constant $B_0 = 0.044092$ cm⁻¹ is within 0.2% of the experimentally observed $B_0 = 0.0441735(12) \text{ cm}^{-1}$.⁶¹

3.4 Spectroscopic properties of tetraacetylene

The (fc)-CCSD(T)/ANO1 harmonic and VPT2 frequencies of the fundamental and combination bands for tetraacetylene are given in Table 6, and the (fc)-CCSD(T)/ANO1 anharmonicity constants (x_{ii}) are given in the ESI.[†] For the four experimentally observed bands, agreement of the observed and calculated frequencies is good at 7 cm^{-1} , which is comparable to the uncertainty of the low resolution measurements. Furthermore,

4 Conclusions

Accurate equilibrium geometries have been determined at the AE-CCSD(T)/cc-pCVQZ level of theory, and the full cubic and semidiagonal quartic force field have been determined at the (fc)-CCSD(T)/ANO1 level of theory for acetylene and the three smallest polyynes. No scaling or adjustments had to be included to match theoretical values with those determined by experiments. The resulting VPT2 fundamental vibrational frequencies and vibration–rotation interaction constants agree with known experimental values, showing about a 5 cm^{-1} deviation in frequencies for bands with high-resolution infrared information. For bands with only low-resolution data, the theoretical frequencies are able to confirm mode assignments or suggest a reassignment, as in the case of the observed band at 1115.0 cm^{-1} of triacetylene to the ν_{9} + ν_{11} combination band, which has previously been attributed to the ν ₇ fundamental. The provisional *ab initio* method used here is also able to accurately reproduce the observed frequencies of combination bands.

The calculated fundamental frequencies for triacetylene and tetraacetylene give insight as to why tetraacetylene has not yet been observed in space. Observation of centrosymmetric molecules in astronomical environments is mainly through infrared detection of the high intensity bending modes; e.g., ν_8 [628.040776(36) cm⁻¹]²⁹

and $\nu_6 + \nu_8$ [1241.060828(38) cm⁻¹]⁴⁶ of diacetylene, or ν_{11} [621.34011(42) cm⁻¹]⁶⁰ and $\nu_8 + \nu_{11}$ [1232.904295(74) cm⁻¹]⁵⁸ of triacetylene. However, the analogous modes for tetraacetylene are the ν_{14} [621.5(5) $\rm cm^{-1}]^{64}$ and ν_{10} + ν_{14} [1229.7(5) $\rm cm^{-1}]^{,64}$ and are predicted to be significantly weaker in intensity due to lower column densities.34,35 Consequently, at these frequencies and resolutions of the previous infrared observations where polyynes were detected,^{20–22,24,25} the transitions of tetraacetylene are blended with those of triacetylene. Other bands of tetraacetylene would be more suitable for identification, such as $\nu_1 + \nu_6$, $\nu_{12} + \nu_{15}$, or ν_{17} that are expected to be equally strong as bands already used to identify di- and triacetylene. Paper Workstein on 2013

(at 3-30) (at 3-40) (at 3-

Overall, the resulting computed geometries lead to equilibrium rotational constants (B_e) , which when corrected for vibrational zero-point effects give ground state equilibrium constants (B_0) that agree with experimental values (0.2%). Based on the small o–c values for acetylene, diacetylene, and triacetylene, we are confident that the fundamental frequencies and spectroscopic constants determined here offer an accurate guide for spectroscopic searches focused on detection of ro-vibrational bands of triacetylene and tetraacetylene. Such work is underway in our laboratory.

Conflicts of interest

There are no conflicts to declare.

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