Theoretical Calculations for the Acidity of Cyanopolyynes $HC_{2n+1}N$ (n = 0-5) in Gas and Aqueous Phases Using Ab initio Methods

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Abstract—Cyanopolyynes have been found in the interstellar medium, cold dust cloud Taurus Molecular Cloud-1, and the Titan's atmosphere. Theoretical calculations are carried out to predict gas and aqueous phase acidities of a series of cyanopolyynes acids. Two levels of theory were used in this study, with the combination of density functional theory, and Møller–Plesset perturbation (MP2) theory, MP2 methods with two types of basis set, namely, Pople's 6-311++g(d, p) basis set and Dunning's aug-cc-pVTZ basis set. The calculations of these molecules reveal that pKa values varying from 12.25 to 17.25 and indicate that the acidity of these molecules in aqueous phase increases whereas the acidity in gas phase decreases with an increasing chain length of these acids.

Index Terms—Ab initio, Acidity, Cyanopolyynes, Density functional theory, Møller–Plesset perturbation.

I. INTRODUCTION

Cyanopolyynes with the general formula, $HC_{2n+1}N$ (n = 0-5), have been detected in the interstellar medium. The larger $HC_{11}N$ was observed in cold dust cloud Taurus molecular cloud (TMC)-1 (Bell et al., 1997). Cyanopolyynes are also discovered in the Titan's atmosphere, which is rich in elements such as carbon, hydrogen, and nitrogen (Kunde et al., 1981). The origin of these large carbon chains with alternating single and triple carbon-carbon (C-C) bonds is still unclear but ion-molecule reactions especially those involving C+ are presumed to be the main source. However, recent studies indicate that neutral-neutral reactions may also lead to such carbon chains (McCarthy and Thaddeus, 2001; Mendoza et al., 2018; and Skomorowski et al., 2018).

The detection of long linear carbon-chain molecules in interstellar and circumstellar clouds has come as a surprise. Ab initio calculations show that long linear species

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are usually more energetic than their ring or ring-chain counterparts and were observed to be less stable in the earthbound environment. In contrast, many linear chains such as the cyanopolyynes $HC_{2n+1}N$ were found to be widespread and abundant in the cold circumstellar and interstellar clouds (Kunde et al., 1981 and McCarthy and Thaddeus, 2001). The abundance of these long chains and the scarcity of their ring counterparts in the cold ultraviolet-shielded clouds yield keys to the formation of large molecules in interstellar space and can potentially help us in understanding the origin of diffuse interstellar bands (Botschwina, 2003).

Carbon chain molecules can be classified into several families, depending on a few factors such as the structure of their linear backbone (acetylenic backbone with alternating single and triple C-C bonds, or cumulenic backbone with double bonds), their electronic ground state (open shell or closed shell) and the groups at the end of the chain (H, CN, CH_2 , or CH_3). The two most widespread families are the polyynes ($HC_{2n+1}N$ and $CH_3C_{2n+1}H$) and the polyacetylene radicals (C_nH). The most abundant is presumably the family of polyacetylenes ($HC_{2n}H$), which unfortunately are non-polar and hence not detectable at radio wavelengths. Note, however, that these species have been detected in mid-infrared spectrum with the infrared space observatory (Botschwina and Horn, 1997).

It has been proposed that the polyynes and carbon-chain radicals formed directly in the gas phase through reactions of CCH with polyynes, polyacetylenes, and/or polyacetylene ions (for example, $HC_{2n+1}N + CCH \rightarrow HC_{2n+3}N + H$) (Botschwina et al., 1997). Such reactions tend to insert two triply bonded carbon atoms into the carbon backbone, uncoupling the formation of chains with an odd number of C-atoms from those with an even number, which could explain the alternating high and low abundances observed as the length of the C backbone increases. For example, in the molecular shell around IRC+10216 and the dark cloud TMC-1, the two main astronomical sources of carbon chain molecules, the abundance of the C₄H radicals with an even number of C-atoms is about 30 times larger than that of radicals of similar size with an odd number of carbon atoms. In contrast, the decrease in abundance between successive species within the even (or odd) number of C-atoms families,

 $C_nH/C_{n+2}H$, is found to be only 4–6 (Botschwina, 2003 and McCarthy and Thaddeus, 2001).

Electronic structure calculations in cyanopolyynes were also reported and guided the discovery of some cyanopolyynes in the interstellar medium (McCarthy and Thaddeus, 2001). The HCoN molecule was already investigated with the Coupled Cluster Theory, at the CCSD (T)/cc-pVTZ level of theory (Botschwina et al., 1997), and the results were employed to determine cyanopolyvne column densities in TMC-1 (Bell et al., 1997). This CCSD (T)/cc-pVTZ study presents the equilibrium geometry and the dipole moment for HCoN (Botschwina et al., 1997). Another study compares dipole moments obtained for small cyanopolyynes with perturbation theory, Møller-Plesset perturbation (MP2), and density functional theory (DFT), through the BP86 functional, using a small basis set, 6–31G*. The results showed that MP2 yields more accurate dipole moment values than BP86 (Moliner et al., 1996). More recent articles also deal with the electronic structure of some cvanopolyvnes using DFT and CCSD (T) (Botschwina, 2003, Qi et al., 2009, Woon and Herbst, 2009, and Scemama et al., 2002). Other works deal only with the stretching modes of a few cyanopolyynes such as HCN, HC₃N (Botschwina et al., 1995), HC₅N (Botschwina et al., 1997), and HC₇N (Botschwina et al., 1997) using the CCSD (T) method with triple-zeta basis sets or some combined treatment.

Moreover, DFT calculations also indicated that the alternation between C-C bonds that maintain characteristics of single and triple bonds persists in polyynes as large as $HC_{30}H$ (Scemama et al., 2002). Interestingly, the difference between the bond lengths for either single and triple C-C bonds near the center of polyynes ($HC_{2n}H$) and cyanopolyynes are predicted as being equal to only 0.1 Å. Another interesting characteristic of these molecules has large molecular dipole moments that facilitate their detection (Arnau et al., 1990). Unfortunately, the experimental values of this quantity are only limited to the smallest cyanopolyynes.

The deprotonation energies of organic acids and the proton affinities of the corresponding conjugate bases are widely used for the prediction of gas phase and aqueous phase Bronsted acidities (Smith and March, 2007, Dewar and Dieter, 1986, Siggel et al., 1988, Kass, 1990, Burk and Koppel, 1993, Koppel et al., 1994, and Burk et al., 1996). Strong acids have small values of deprotonation energy (that is, the release of the proton is easier) whereas strong bases have large values of proton affinity (that is, the binding to the proton is stronger). Several works on the prediction of the acidity of organic and inorganic acids can be found in literature. For instance, Smith and Radom (Smith and Radom, 1995 and Smith and Radom 1995) have shown that the G2 and G2 (MP2) methods provide excellent results for both deprotonation enthalpies and proton affinities of small molecules.

Catalan and Palomar (Catalan and Palomar, 1998) have investigated gas-phase acidities of a number of species and have shown that the calculations at B3LYP/6–311+G (d) and 6–311+G (3df, 3pd) level of theory correlate well with the experimental data. Good correlations have been obtained between experimental pKa values of a wide range of organic Bronsted acids and their calculated gas-phase deprotonation enthalpies (Choho et al., 1996). Correlations between theoretical results and gas-phase acid-base equilibrium constants of organic compounds have been reported for amines, alcohols, and thiols (Yang and Mortier, 1986, Contreras et al., 1999, and Pérez et al., 2000). An excellent correlation was obtained between the aqueous-phase acidity calculated with the HF/3–21G (d) method and experimental pKa values for a series of nitrogen bases (D'Souza et al., 2000).

Recently, a reasonable correlation was obtained between the deprotonation energies of the compounds calculated at the HF/3–21G and B3LYP/6–31G (d) levels of theory and their aqueous pKa values (Rezende, 2001 and Rezende 2001). The ability to predict acidity using a coherent, well-defined theoretical approach, without external approximation or fitting to experimental data would be very useful to chemists. Due to the rapid development of computational chemistry, the acidity of small molecules in the gas phase can be calculated with equivalent or greater accuracy than can be obtained experimentally. However, the current situation is less satisfactory in solution, mostly due to the difficulty of calculating solvation energies with adequate accuracy.

In this study, the acidy of six cyanopolyynes acids is predicted in the gas and aqueous phases using DFT and MP2 methods with two basis sets, 6-311++g (d, p) and aug-cc-pVTZ. The charge distribution and the effect of the length of the cyanopolyynes on the acidy of these compounds are elucidated.

II. COMPUTATIONAL METHODS

A. Gas-phase Acidity Calculations

Calculations are performed with the Gaussian 09 W software package (Frisch et al., 2009). The geometries of the neutral and deprotonated species are fully optimized using DFT and MP2 theory, MP2 level of theory at the 6–311++G (d, p) and aug-cc-pVTZ basis sets. Frequency calculations are performed at the same level of theory to characterize the stationary points obtained. The gas-phase Gibbs free energy change (ΔG_{gas}°) of Scheme 1 is calculated using Eq. (1). For ΔG_{gas}° (H⁺) the experimental value of –6.28 kcal/mol is used (Liptak et al., 2002).

$$AH_{(\text{gas})} \to A^{-}_{(\text{gas})} + H^{+}_{(\text{gas})}$$
$$\Delta G^{\circ}_{\text{gas}} = G^{\circ}_{\text{gas}} \left(A^{-}\right) + G^{\circ}_{\text{gas}} \left(H^{+}\right) - G^{\circ}_{\text{gas}} \left(AH\right) \tag{1}$$

B. Aqueous-phase Acidity Calculations

Solvent effects are taken into account by means of the polarizable continuum model (PCM) (Miertus et al., 1981, Miertus and Tomasi, 1982, and Cossi et al., 1996) through the full optimization calculations at the B3LYP/6–311++G (d, p) level of theory (using the gas-phase optimized geometries). The PCM calculations employ the UAHF atomic

radii when constructing the solvent cavity for the calculation of the Gibbs free energy of solvation. A common practice to calculate the aqueous Gibbs free-energy change of an acid dissociation (ΔG_{aq}°), see Eq. (2) is by combining the ΔG_{gas}° of the deprotonation process with the change in Gibbs free energy of solvation $\Delta \Delta G_{solv}^{\circ}$ using the thermodynamic cycle of Scheme 1 and Eq. (3). For ΔG_{solv}° (H⁺) the experimental value of -264.61 kcal/mol is used (Liptak et al., 2002). The aqueous pKa is calculated according to Eq. (5). All the calculations and experimental data reported in this paper are at 298.15 K.

$$\Delta G_{aq}^{\circ} = G_{aq}^{\circ} \left(A^{-} \right) + G_{aq}^{\circ} \left(H^{+} \right) - G_{aq}^{\circ} \left(AH \right)$$
(2)

$$\Delta G_{aq}^{\circ} = \Delta G_{gas}^{\circ} + \Delta \Delta G_{solv}^{\circ}$$
⁽³⁾

$$\Delta G_{aq}^{\circ} = G_{gas}^{\circ}(A^{-}) + G_{gas}^{\circ}(H^{+}) - G_{gas}^{\circ}(AH) + \Delta G_{solv}^{\circ}(A^{-}) + \Delta G_{solv}^{\circ}(H^{+}) - \Delta G_{solv}^{\circ}(AH)$$
(4)

$$pK_{a} = \frac{\Delta G_{aq}^{\circ}}{RT \ln 10}$$
(5)

Charif and his group (Charif et al., 2007) have found a correlation between these two Gibbs free energy changes with



Scheme 1. $G_{solv}^{\circ}(X)$ represents the Gibbs free energy of solvation of species X.

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 $R^2 = 0.947$ that might be very helpful to be used as a scale factor to get values closer to the experimental results. They found a correlation between the experimental and calculated aqueous pKa values. The regression equation with the corresponding standard deviation and correlation coefficient (R^2) values is given in Eq. (6).

$$pK_a = 0.8755 \ pKa \ (calc.) - 1.7563 (R^2 = 0.934, \ SD = 4.80)$$
(6)

III. RESULTS AND DISCUSSION

A. Optimized Geometry

The molecules under study were optimized to obtain their ground state structures, with the levels of theory aforementioned in the methods of calculations. Fig. 1 shows the optimized geometry of the studied cyanopolyynes. The optimized structures were at the ground state with no imaginary frequencies. The bond lengths of the studied molecules are listed in Table I.

The vibrational modes of the optimized structures were carefully examined. The most important vibrational mode belongs to the CH stretching. The frequencies of the stretching vibration are listed in Table II. In addition, the values of the dipole moment of these compounds are also listed. It is clear that the value of the stretching frequency is increasing from HCN to the HC₅N, whereas for the molecules beyond that the value is constant. This indicates that the strength of the CH bond increases with the length of the chain, resulting in difficulty for the proton to dissociate as the chain becomes longer until it becomes constant at a certain length. On the other hand, the dipole moments of the molecules show a different trend whereby the dipole moment is increasing as the length of the molecule increases.



Fig. 1. The optimized structures of the studied cyanopolyynes.

TABLE I Bond Lengths in a of the Optimized Structures of the Studied Cyanopolyynes in the Gas-Phase and the Values Between Brackets are for the Aqueous Phase

Bond	HCN	HC ₃ N	HC ₅ N	HC ₇ N	HC,N	HC ₁₁ N
H–C	1.078 (1.081)	1.076 (1.078)	1.066 (1.068)	1.066 (1.068)	1.066 (1.068)	1.066 (1.068)
$C_1 \equiv C_2$		1.236 (1.236)	1.226 (1.227)	1.228 (1.228)	1.229 (1.229)	1.229 (1.229)
C ₂ -C ₃		1.386 (1.385)	1.363 (1.363)	1.360 (1.360)	1.358 (1.358)	1.358 (1.358)
$C_3 \equiv C_4$			1.232 (1.232)	1.238 (1.238)	1.241 (1.241)	1.242 (1.242)
$C_4 - C_5$			1.368 (1.367)	1.352 (1.352)	1.347 (1.347)	1.344 (1.344)
$C_{5} \equiv C_{6}$				1.236 (1.236)	1.243 (1.243)	1.247 (1.246)
$C_{6} - C_{7}$				1.365 (1.364)	1.348 (1.347)	1.341 (1.341)
$C_8 \equiv C_9$					1.238 (1.238)	1.246 (1.246)
$C_{9} - C_{10}$					1.364 (1.363)	1.346 (1.345)
C ₁₀ ≡C ₁₁						1.239 (1.239)
C ₁₁ -C ₁₂						1.364 (1.362)
C≡N	1.183 (1.181)	1.193 (1.192)	1.184 (1.183)	1.186 (1.184)	1.186 (1.185)	1.186 (1.185)

TABLE II Dipole Moment (m) and the Stretching Frequencies in the Gas and Aoueous Phases

		L				
Bond	HCN	HC ₃ N	HC ₅ N	HC ₇ N	HC ₉ N	HC ₁₁ N
μ (gas) (debye)	3.333	4.273	5.021	5.657	6.186	6.636
μ (aq) (debye)	4.103	5.560	6.599	7.395	7.978	8.412
C-H str. (gas) (cm ⁻¹)	3452.76	3472.67	3487.72	3486.33	3486.02	3486.25
C-H str. (aq) (cm ⁻¹)	3428.35	3450.98	3467.33	3466.35	3466.01	3466.14

TABLE III Density Functional Theory results of the Gibbs Free Energy in (kcal/mol), PKa Values in the Gas and Aqueous Phases and the Scaled Values of PKa in the Aqueous Phase

Bond	Basis set	ΔG_{gas}°	ΔG_{aq}°	pK _a -gas	pK _a -aq	pK _a -aq-sc
HCN	6–311++g (d, p)	341.993	21.963	250.705	16.000	12.251
	aug-cc-pVTZ	340.110	20.708	249.577	15.096	11.460
HC ₃ N	6–311++g (d, p)	339.483	29.493	248.917	21.694	17.236
-	aug-cc-pVTZ	337.600	28.238	247.560	20.540	16.226
HC ₅ N	6–311++g (d, p)	333.208	29.493	244.563	21.853	17.376
	aug-cc-pVTZ	331.953	28.865	243.453	20.956	16.591
HC ₇ N	6–311++g (d, p)	329.443	30.120	241.570	21.890	17.408
	aug-cc-pVTZ	328.188	28.865	240.446	21.128	16.741
HC₀N	6–311++g (d, p)	326.305	30.120	239.229	22.075	17.570
[^]	aug-cc-pVTZ	325.050	28.865	238.275	21.276	16.871
HC ₁₁ N	6–311++g (d, p)	324.423	30.120	237.953	22.246	17.720
	aug-cc-pVTZ	323.168	29.493	236.877	21.710	17.251

A. Gas-phase Acidities

The calculated gas-phase ΔG_{gas}° of the acid dissociation and aqueous pK_a values (relative to water) at 298.15 K is reported in Tables III and IV. Besides that, the last column is for the scaled values which were calculated according to the Equation (6).

A satisfactory linear correlation is obtained showing that the gas-phase Gibbs free energy change of deprotonation generally increases when the acidity of these compounds decreases. It should be noted that very accurate aqueous pK_a values (relative to water) can only be obtained for acids weaker than the hydronium ion ($pK_a = -1.74$) and stronger than water



Fig. 2. The correlation between the pK_a values and chain length of the cyanopolyynes in the gas phase.

 $(pK_a = 15.74)$ (Bell et al., 1997). It is well known that in the gas phase, acidity orders may be much different than in aqueous solution for certain families of compounds. Simple alcohols are an example of this: Whereas tertiary alcohols are more acidic than secondary and primary alcohols in the gas phase, in aqueous solution the acidity order is reversed because the much bulkier tertiary anion is poorly solvated. A similar pattern is found in simple aliphatic carboxylic acids and when analyzing the basicity order of amines (Bell et al., 1997).

From the results we have obtained, we can infer that the aqueous phase acidity order of the family of cyanopolyynes acids is very similar to that in the gas phase. Hence, it might be possible to use gas-phase calculations to predict the aqueous pK_a values of carbon acids.

As shown in Fig. 2, the pK_a values of the studied cyanopolyynes are generally decreasing with the increase of the chain length in the gas phase in parallel behavior with the values of the ΔG_{gas}° . Fig. 2 shows that the calculated pK_a values using DFT method are lower than the MP2 method. The general behavior of the plot may be attributed to the

effect of the solvent molecules on the acidity of these compounds. In conclusion, in the gas phase, the acidity of these compounds increases with increasing chain length; however, this may be attributed to the method and the type of the basis set used in the calculations.



Fig. 3. The correlation between the pK_a values and chain length of the cyanopolyynes in the aqueous phase.

B. Aqueous-phase Acidities

The calculated aqueous phase ΔG_{aq}° of the acid dissociation and pK_a values of the compounds under study (using Eq. 4 and 5) at 298.15 K are reported in Tables III and IV using DFT and MP2 methods, respectively. According to both

TABLE IV Møller–Plesset Perturbation Results of the Gibbs Free Energy in (kcal/mol), pKa Values in Gas and Aqueous Phases and the Scaled Values of PKa in the Aqueous Phase.

Bond	Basis set	ΔG_{gas}°	ΔG_{aq}°	pK _a -gas	pK _a -aq	pK _a -aq-sc.
HCN	6–311++g (d, p)	343.875	23.845	252.213	17.409	13.485
	aug-cc-pVTZ	340.110	20.080	249.457	14.935	11.319
HC ₃ N	6–311++g (d, p)	342.620	32.631	251.424	23.704	18.997
5	aug-cc-pVTZ	338.228	28.238	248.235	20.844	16.493
HC ₅ N	6–311++g (d, p)	338.228	33.258	248.084	24.333	19.547
-	aug-cc-pVTZ	334.463	30.120	245.472	22.150	17.636
HC ₇ N	6–311++g (d, p)	334.463	33.258	245.198	24.407	19.612
	aug-cc-pVTZ	329.443	28.865	241.699	21.154	16.764
HC₀N	6–311++g (d, p)	332.580	33.886	244.002	24.724	19.890
<i>,</i>	aug-cc-pVTZ	328.815	30.748	241.178	22.400	17.855
HC ₁₁ N	6–311++g (d, p)	330.698	33.886	242.468	24.730	19.895
	aug-cc-pVTZ	325.678	29.493	238.788	21.659	17.206



Fig. 4: The charge distribution on atoms in the gas and aqueous phases as found in the natural bond orbital analysis.

methods, the values have an opposite trend to the values in the gas phase in which the change in Gibbs free energy increases with the increase in the length of the molecule. Tables III and IV also show the results of the scaled values of the pK_a . The scaled values are smaller than those calculated with Equation (6) which indicates that the calculated change in Gibbs free energy of solvation might be slightly overestimated. The reason for this lies in the inadequacy of continuum solvation models to accurately calculate solvation energies since explicit solute–solvent interactions are ignored. The inclusion of continuum solvent effects in the geometry optimizations and frequency calculations might improve these results slightly. The plot of the pK_a values versus the studied molecules is displayed in Fig. 3.

A satisfactory correlation is obtained between the length of the carbon chain and the values of pK_a . However, there are some points to be highlighted. The scaled values are lower than the calculated raw values. Furthermore, the DFT method has shown lower values of pK_a than the MP2 method. Finally, the pK_a values are slightly increasing with the increase of the chain length starting from HC₃N and beyond and hence the acidity is decreasing with increasing the chain length.

For more understanding to the properties of these molecules, natural bond orbital method was used to calculate the charge distribution on each atom of the molecules and shown in Fig. 4 at the gas and solution phases. As shown in Fig. 4, there is no single trend for the charge distribution on the atoms. However, in general, the N and H atoms have the highest negative and positive charges, respectively.

IV. CONCLUSIONS

In the present work, we have shown satisfactory values of aqueous pK_a values of cyanopolyynes in the gas and aqueous phases through the calculation of the change in Gibbs free energy for these molecules and their deprotonated form. The correlation equation between the experimental aqueous pK_a values and the calculated values was used to predict better pK_a values. To assess the validity of our results, further experimental work is required. The difference in the acidity of the compounds studied can be interpreted through the charge distribution and conjugation effects. In the gas phase, the pK_a values decrease and the acidity increase with the increase of conjugation or the molecule length, whereas, in the aqueous phase, the pK_a values increase and the acidity decrease with increasing the chain length.

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