Cyclo[18]Carbon: Insight into Electronic Structure, Aromaticity and Surface Coupling

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ABSTRACT: Cyclo[18]carbon (C18) is studied computationally at density functional theory (DFT) and ab initio levels to obtain insight into its electronic structure, aromaticity, and adsorption properties on a NaCl surface. DFT functionals with a small amount of Hartree-Fock exchange fail to determine the experimentally observed polyyne molecular structure revealing a cumulene-type geometry. Exchange-correlation functionals with a large amount of Hartree-Fock exchange as well as ab initio CASSCF calculations yield the polyyne structure as the ground state while the cumulene structure as a transition state between the two inverted polyynes structures through a Kekule distortion. The polyyne and the cumulene structures are found to be doubly Hückel aromatic. The calculated adsorption energy of cyclo[18]carbon on the NaCl surface is small (37 meV/C) and almost the same for both structures implying that the surface does not stabilize a particular geometry.

Carbon is one of the most diverse elements of the periodic table with respect to possible allotropes. Among them sp2-hybridized diamond, multilayered sp2-hybridized graphite and monolayer graphene, fullerenes and nanotubes, sp-hybridized carbyne, and linear polyynes are most attractive materials and unique objects for practical applications and theoretical surveys. The number of hypothetically predicted carbon allotropes reaches more than five hundred different structures in accordance with the SACADA database last updated on May 2017. Nevertheless, carbon allotropes are still in focus of synthetic attempts and theoretical predictions. The most impressive recent achievements in this field is the successful synthesis and characterization of the closed-circle polyyne containing 18 sp-hybridized carbon atoms, called cyclo[18]carbon. It is notable that cyclo[18]carbon possesses the special "double" aromaticity because of delocalization of two π-electrons oriented in the plane of C18 ring (18ππ) and perpendicular to it (18σσ, Figure 1) that was first mentioned by Diederich et al. and developed further by Fowler et al.

Numerous electronic structure calculations of cyclo[18]carbon molecule have been carried out at the different levels of theory including the HF, SCF, MCSCF, MP2, DFT, and CCSD quantum Monte Carlo (QMC) and CCSD levels. Surprisingly, only the HF, QMC and CCSD methods yield the polyyne structure that experimentally has been observed in high-resolution atomic force microscopy (AFM) measurements of cyclo[18]carbon adsorbed on bilayer NaCl on Cu(111) surface at the temperature of 5 K. Most of the common DFT functionals (BLYP, PBE0, etc.) as well as MP2 calculations yield the non-bond-length alternating cumulene structure of cyclo[18]carbon. Torelli and Mitas explained that DFT fails to produce correct results due to the overestimation of electronic correlation effects at the expense of exchange, while Houk and Planner proposed that the incorrect MP2 results are caused by an RHF→UHF instability (i.e., MP2 energies are inaccurate, because the RHF wave function is unstable with respect to the UHF wave function). Here, we support the notion of Torelli and Mitas by demonstrating that increasing the Hartree-Fock exchange part in an exchange-correlation functional leads to the experimentally observed polyyne structure of cyclo[18]carbon; ab initio CASSCF calculations also exhibit the correct polyyne structure. We utilized the gauge-including magnetically induced currents (GIMIC) approach33 to investigate the magnetically induced current density and the "double" aromaticity of cyclo[18]carbon beyond the previously published NICS software and isopercenctric CTODC-DZ (continuous transformation of origin of current density – diamagnetic zero) studies. The idea that the cumulenic structure of cyclo[18]carbon is the ground-state structure in the gas phase in isolated form, while surface interaction may induce stabilization of the polyyne-type structure has been investigated and disproved by simulations of cyclo[18]carbon adsorption on the NaCl (100) surface within periodic boundary conditions (PBC).

Figure 1. Polyyne and cumulene structure of cyclo[18]carbon.
The following exchange-correlation functionals (see supporting information (SI) for details) with increased percentage of Hartree-Fock exchange (HFE) relative to the common B3LYP functional (20% HFE) have been selected to probe the ground state structure of cyclo[18]carbon: BMK (42% HF), M06-2X (54% HFE), BHandHLYP (50% HFE) and wB97XD (22% short-range HFE, range separation parameter γ=0.2). All of them provide the polyyne-type ground state structure of cyclo[18]carbon of the C₀₀ symmetry point group with alternating single and triple C-C bonds (the corresponding structural parameters are collected in Table 1). No significant bond angles alternating θ₁ and θ₂ was observed. The typical deviations are less than 1°. All calculated vibrational frequencies were found to be real (ω₁ is real and positive in Table 1) for all cases of polyyne cyclo[18]carbon indicating the true energy minimum whereas one imaginary frequency was obtained for the cumulene structure.

In favor of the hypothesis that increased contribution of HFE affects the ground-state geometry of large cyclo[4N+2]carbon molecules (N=4-8), we refer to the study by Remya and Suresh. They used the meta-GGA M06L local functional with zero 0% HFE, which revealed a cumulenic ground-state structure for the studied cyclo[4N+2]carbons (N=4-8) including cyclo[18]carbon. In order to estimate the threshold HFE value sufficient to obtain the correct bond alternation in cyclo[18]carbon in the hybrid functional, we performed structural relaxations using two DFT levels (B3LYP/6-311++G(d,p) and HSE06 with plane wave (PW) basis set as described in details in SI) varying the HFE part from 0 to 100%. We found that 37% HFE contribution at the B3LYP/6-311++G(d,p) and 32% HFE within HSE06/PW levels are enough to reproduce the experimentally observed polyyne bond-length alternation of cyclo[18]carbon. Other hybrid functionals used in this work reproduce the HFE trend obtained with the B3LYP and HSE06 reference functionals (Figure S1).

The attempts to optimize the cumulenic-type geometry of cyclo[18]carbon within BMK, M06-2X, BHandHLYP, wB97XD functionals and CASSCF(8,8) calculations with different symmetry constraints always yielded a saddle point with the one or few imaginary frequencies. The calculations suggest that the cumulene-type geometry might be a transition state structure for a bond shift between two polyyne forms of cyclo[18]carbon with inverted ordering of the single and triple bonds (both of C₀₀ symmetry, Figure 2). Indeed, QST3 optimization of the transition state structure for such a bond shift resulted in a bond-angle alternating cumulene-type geometry (Table 1) of the same C₀₀ symmetry with one imaginary frequency corresponding to the Kekule-type in-plane C-C stretching vibrations (Figure 2). The calculated energy barrier for the inversion process depends on the method. The BMK functional underestimates the bond shift barrier in comparison to the other levels of theory, demonstrating a barrier of only +3.5 kcal mol⁻¹, while the rest of the functionals (M06-2X, BHandHLYP, wB97XD) as well as CASSCF(8,8) calculations, yield very similar barrier heights of 8.8-12.1 kcal mol⁻¹ (Figure 2). The CASSCF calculations show that the predominant weight of a closed-shell singlet determinant for the polyyne and cumulenic structures is 0.95, implying that single-reference approaches should be applicable.

Using the M062X/6-311++G(d,p) optimized ground-state polyyne structure and the transition-state cumulene geometry, we performed calculations of the magnetically induced ring current strength using the GIMIC approach (see details in the SI). We found that applying an external magnetic field perpendicular to the molecular plane of cyclo[18]carbon induces a strong diatropic ring current of 29 nA T⁻¹ showing that polyyne cyclo[18]carbon is aromatic in agreement with NICS and ipsoconcentric CTOCD-DZ calculations and the Hückel (4N+2) rule. For comparison, benzene sustains a net diatropic ring current about 12 nA T⁻¹. The signed modulus of the magnetically induced current density (Figure 3) shows that diatropic ring current in polyyne-type cyclo[18]carbon originates from the two scaffolds of 18π-electrons oriented in- and out-of molecular plane proving the double aromaticity. Calculating the ring current strength within reduced contours that cover particular in- and out-of-plane delocalization areas (see details in SI) shows the efficient delocalization of the 18πout electrons (I₀out = 2.18 nA T⁻¹) and less efficient delocalization of 18πin electrons (I₀in = 7.2 nA T⁻¹) due to the perpendicular alignment of these orbitals. The cumulene-type TS structure sustains much higher net diatropic current (73.2 nA T⁻¹) due to the more efficient overlap and delocalization of the πout and πin orbitals. The strengths of the I₀ and I₀out currents for the cumulene structure were also found to be not the same (32.2 and 41 nA T⁻¹, respectively) meaning that the πin and πout orbitals are not identical with respect to the orbital delocalization. The reason for that is the same as for polyyne-type cyclo[18]carbon: the angular overlapping of 18πout electrons results in less efficient delocalization comparing to the perpendicularly oriented 18πin electrons. Despite the higher aromaticity of transition state cumulene-
type cyclo[18]carbon its total energy is considerably higher relative to the ground state polyyne-type cyclo[18]carbon. The reason for that argued in terms of the second-order Jahn-Teller effect\textsuperscript{29,32} that overcomes the aromatic stabilization of cumulenic-Teller effect\textsuperscript{29,32} and hence results in bond-length alternated polyyne structure.

![Figure 3](image)

**Figure 3.** The signed modulus of the magnetically induced current densities for the polyyne and cumulene cyclo[18]carbon.

In order to elucidate the hypothesis that the cumulene type geometry is a gas phase global minimum structure, while the polyyne geometry can only exist on the NaCl surface due to extra stabilization effects (relative to cumulenic one), we performed calculations of the adsorption of polyyne- and cumulene-type cyclo[18]carbon on a NaCl (100) surface using the PBC approximation (see SI for details). The initial $4 \times 4$ thin-film models of NaCl superstructure consisting of three atomic layers were obtained from a well-converged NaCl unit cell and then relaxed. We found that the unit cell is large enough to simulate cyclo[18]carbon adsorption on the NaCl surface avoiding spurious interactions between the molecules with an effective separation distance of 9.58 Å between nearest neighboring cyclo[18]carbon molecules. The simulated superstructure is shown in Figure 4a. Regardless of the assumed initial cyclo[18]carbon geometry (cumulenic or polyynic), the final optimized geometry was found to be of cumulenic type when the PBE functional with 0% of HFE was utilized. This confirms once again that functionals like PBE cannot properly describe geometry and electronic structure of cyclo[18]carbon.\textsuperscript{51}

![Figure 4](image)

**Figure 4.** (a) Top (left) and side (right) views of the NaCl supercell with adsorbed cyclo[18]carbon. (b) Results of the NEB simulation of the cyclo[18]carbon mobility along the NaCl surface. The arrows depict the direction of cyclo[18]carbon molecule movement during NEB cycle.

To estimate the adsorption parameters of polyyne cyclo[18]carbon, we used its equilibrium structure obtained at the CASSCF(8,8) level and then fixed all carbon positions but allowed to completely relax the NaCl (100) substrate. Similarly we calculated the cumulene type but allowing to relax all atoms. We found, that the formation of the C\textsubscript{18}/NaCl heterostructure does not play any significant role in the stabilization of the polyyne structure. The adsorption energies were found to be almost equivalent (37 meV/C) for both cumulenic and polyynic structures. It confirms that there is no surface-induced transition between polyyne and cumulenic structures, because the adsorption energies of both structures are the same. Accounting for the very weak interaction between cyclo[18]carbon and NaCl it is evident that the molecules can freely move along the NaCl surface even at very low temperatures, which is in agreement with experimental observations.\textsuperscript{16} Indeed, calculations using the nudge elastic band (NEB) method\textsuperscript{36} yield a very small energy barrier of 30 meV for the cyclo[18]carbon mobility on the NaCl surface (Figure 4b).

In conclusion, the recently synthesized cyclo[18]carbon molecule has been studied computationally at the DFT and CASSCF levels of theory in order to confirm that the polyyne structure is the global minimum. The cumulene-type structure of cyclo[18]carbon is found to be a transition state for the single-triple bond inversion process. This interconversion is mediated by the Kekule-type stretching vibration with the estimated energy barrier of around 10 kcal mol\textsuperscript{–1}. An extended amount of Hartree–Fock exchange in the employed functional is crucial for obtaining the experimentally observed polyyne structure of cyclo[18]carbon. The double aromaticity of both polyyne and cumulenic cyclo[18]carbon has been investigated by gauge-including magnetically induced currents calculations, which predict that the bond-length alternating polyyne structure is less aromatic than non-alternating transition-state structure due to a less efficient overlap of \textit{\pi} and \textit{\sigma} orbitals of the polyyne structure. Two molecular structures of cyclo[18]carbon have identical adsorption energies on the NaCl surface, suggesting that there is no extra stabilization favoring polyyne structure as compared to the cumulenic one. NEB simulations show that cyclo[18]carbon can move almost without any barrier on the NaCl (100) surface, which is in agreement with experimental data.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.XXXXXXX. Computational details and corresponding references on the employed methods, bond length dependence on the percentage of HFE contribution within the HSE06 and B3LYP hybrid functionals (PDF).

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**Notes**

The authors declare no competing financial interests.

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