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An Entrant of Smaller Fullerene: C₅₆ Captured by Chlorines and Aligned in Linear Chains

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Since the first celebrated fullerene, $C_{60}(I_h)$, was macroscopically synthesized in 1990, a growing family of fullerenes has come into reality. The number of novel fullerenes is still increasing even at the present time. However, most of the synthesized fullerenes are larger cages containing carbon atoms more than 60. The fullerenes smaller than C_{60} are predicted to have unusual electronic properties because of their geometric structures with higher curvature, and have aroused the long-attention of chemists in either theoretic computations or experimental invesitigations.^{2,3} These smaller fullerenes, however, inevitably consist of structural abutting pentagons and violate the well-known isolated pentagon rule (IPR), leading to their instability and elusion in standard laboratory conditions under ambience atmosphere. Recent experiments have demonstrated the IPR-violating fullerenes could be stabilized either by endohedral encapsulation of metal clusters3a,5 or exohedral derivatization.3c-g,6 By the latter method, three kinds of smaller fullerenes are stabilized and unambiguously characterized heretofore, that is, $C_{20}Br_nH_{20-n}$, $^{3e,f}C_{50}Cl_{10}$, 3c,g and $C_{58}F_{18}$. 3d Among them, only C₅₀Cl₁₀ was the species directly captured during the fullerene growth in carbon arc plasma, so as to be helpful for understanding fullerene formation. In the paper about $C_{50}Cl_{10}$, 3c we mentioned the mass spectrometry observation of another smaller chlorofullerene, C₅₆Cl₁₀, as one of the byproducts. Following this observation, some theoretical predictions about the structures of C₅₆Cl₁₀ and pristine C₅₆ (#913)⁷ have been reported.⁸ However, authentic structure, assembly, and properties of this new entrant of smaller fullerene will need to be experimentally investigated. In this Communication we focus our efforts on isolation and characterization of the C₅₆Cl₁₀. Its structure, with implications on stabilization, chemical reactivity, and alignment in crystal has been unambiguously identified by X-ray diffraction.

 $C_{56}Cl_{10}$ was produced in a modified Krätschmer–Huffman arc-discharge reactor. ^{1b,9} Crude soot was extracted with toluene in a supersonic bath, followed by separation and purification using multistage recyclic high performance liquid chromatography (HPLC) on Shimadzu LC-6AD HPLC instrument with a Cosmosil Buck-yprep column (10 i.d. \times 250 mm) eluted by toluene at a flow of 4.0 mL/min under the temperature of 40 °C (The chromatograms for $C_{56}Cl_{10}$ isolation are shown in Supporting Information). The purified $C_{56}Cl_{10}$ is soluble in carbon disulfide, chloroform, and toluene with absorption around 304, 364, 444, and 509 nm (Supporting Information). By solvent evaporation from its carbon disulfide solution, its dark red single crystals suitable for X-ray diffraction were obtained.

As revealed by X-ray diffraction (Figure 1), the overall symmetry of $C_{56}Cl_{10}$ is $C_{2\nu}$, the same as its parent cage. The most outstanding feature of the C_{56} structure is the existence of four pairs of adjacent pentagons, that are respectively bonded with eight of the chlorines. For a spherical carbon cage, the local surface strain can be defined

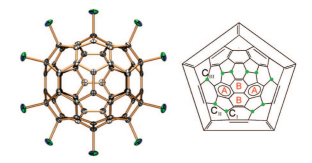


Figure 1. Structures of $C_{56}Cl_{10}$: ORTEP drawing with thermal ellipsoids at 50% probability level (left) and Schlegel diagram with chlorines indicated by green dots (right).

in terms of pyramidalization angle, that is, the π -orbital axis vector (POAV), at the involved carbon atom. 10 The POAVs of the two sets of carbon atoms (i.e., $C_{\rm I}$ and $C_{\rm II}$ in Figure 1) at the adjacent pentagons of the pristine C₅₆ cage are 14.63 and 16.32°, 8b respectively, sharply higher than that of the regular POAV in I_h - C_{60} cage (11.64°). Such higher POAVs lead to the significantly reduced angles between the mean planes of the involved rings and subsequently the high reactivity of the bare C₅₆ cage. However, the enhanced curvature stain can be released as the result of chlorination to transfer the carbon atoms at the pentagon adjacencies as sp³-hybriderized. As shown in the structural data of C₅₆Cl₁₀ (Supporting Information), the average C-C_I-Cl and C-C_{II}-Cl bond angles are respectively 3.72° and 4.43° deviated from the regular tetrahedron (109.48°). Therefore, adding chlorines at adjacent-pentagons sites is well favorite geometrically, so as to stabilize this IPR-violating smaller fullerene.

Another factor in contributing to the overall stability of the C₅₆ is the aromaticity of the remaining carbon skeletons after chloroderivatization. Besides the eight chlorines bonded at the abuttingpentagons sites, the remainder of the chlorines are linked at two hexagon-hexagon-pentagon vertexes (i.e., the C_{III} in Figure 1) to complete a close "belt" of chlorine-bonded sp³-carbons cutting the cage into two aromatic fragments of C₁₆ and C₃₀. The chlorination pattern in C56Cl10 agrees well with proposed principles for fullerene reactivity. 11 In the pyrene portion of C₁₆ there are two sorts of hexagons, rings A and B (Figure 1). Ring A with fairly equalized C-C bond lengths ranging from 1.395 to 1.411 Å suggests a good electronic delocalization, whereas ring B is less aromatic with the alternating C-C/C=C bond lengths ranging from 1.363 to 1.455 Å. This fits well with our theoretical computations about the nucleus-independent chemical shifts (NICS) values at the centroids of the hexagons, with values of -14.36 and -8.87 ppm for ring A and B, respectively. The larger segment of C₃₀ is just a half of I_h -C₆₀ with alternating single-double bonds ranging from 1.350 to 1.446 Å.

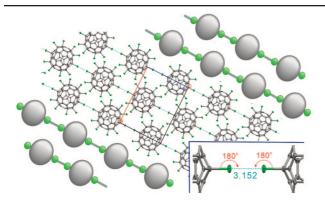


Figure 2. Alignment of $C_{56}Cl_{10}$ molecules along c axis. The skeletons of the C₅₆Cl₁₀ molecule on both sides are indicated as gray/green balls for a clearer view of the pearl-necklace-shaped alignment. Inset shows the distance and angles of the short C-Cl···Cl-C contact.

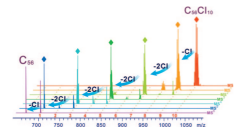


Figure 3. Dechlorination of C₅₆Cl₁₀. Multistage mass spectrometry (MSⁿ. n = 1-6) showing the formation of C₅₆ by progressive dechlorination of $C_{56}Cl_m$ (m = 1-10) (m value is indicated as number in red on the bottom line, and the species selected for next stage of MS fragmentation are marked with colored rhombus).

Of interest is the assembly of the $C_{56}Cl_{10}$ molecule in the crystal. There are three kinds of intermolecular contacts among C₅₆Cl₁₀ molecules in the crystal: $\pi - \pi$, C···Cl, and Cl···Cl contacts (see Supporting Information). The most intriguing intermolecular interaction is the linear C-Cl-···Cl-C short contact along the c axis in the crystal of C₅₆Cl₁₀ (Figure 2). According to the symmetry, C₅₆Cl₁₀ has three different C-Cl bonds (1.779, 1.785, and 1.810 Å). Interestingly only Cl atoms which are involved in the highly elongated C-Cl bonds form short Cl···Cl contacts, although strong repulsive forces are expected due to accumulation of negative charge on the corresponding Cl atoms. 12 This Cl···Cl distance is rather short (3.152 Å), which is, to our knowledge, one of the shortest Cl···Cl contacts observed in the chlorofullerenes family. 13 Unexpectedly, the C₅₆Cl₁₀ molecules in the crystal are aligned as entirely straight chains with the angle of C-Cl···Cl-C being 180°, just like pearl necklaces (Figure 2).

Suggested by regioselectivity of IPR-violating C₆₀Cl₈ (#1809) in Friedel-Crafts-type and nucleophilic reactions where the substitution reactions prefer to take place for the chlorines in longer C-Cl bonds of 1.81 Å (the unreacted chlorines are located in the C-Cl bond with distance of 1.78 Å), 14 linear fullerene polymers of C₅₆ are expected to be synthesized because the two chlorines involved in the linear chains have longer C-Cl distance (1.81 Å vs the others of 1.78 Å) and are sterically ready for reaction with bidentate donors to construct one-dimensional architectures. Such kind of one-dimensional fullerene polymer may be applicable as electronic or mechanical materials.

Dechlorination of C₅₆Cl₁₀ was conducted by multistage mass spectrometry (MSⁿ), where stepwise dissociation of chlorine was achieved by collision of $C_{56}Cl_m$ (m = 1-10) molecules with helium buffer gas in ion trap (Figure 3). The eventual formation of bare C₅₆ units suggests the possibility of studying electronic properties of its bare cages by, for example, photoelectron spectroscopy.

In summary, our experiments have demonstrated the existence of C₅₆ (#913), a new entrant in the family of smaller fullerenes. The otherwise unstable IPR-violating C₅₆ has been macroscopically stabilized by chlorination as C₅₆Cl₁₀, and thus provides significant experimental opportunities into the new world of fullerenes smaller than C₆₀. The crystallographic data of C₅₆Cl₁₀ reveal an unusual Cl···Cl short contact and the alignment of pearl-necklace-like molecular chains, which can be potentially converted into a class of useful one-dimensional fullerene polymer through previously established regioselective reactions.

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Supporting Information Available: Crystallographic information file (CIF), intermolecular contacts in C₅₆Cl₁₀ crystal, HPLC chromatograms, and UV-Vis-NIR spectrum of C56Cl10. This material is available free of charge via the Internet at http://pubs.acs.org.

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