Large-Cavity Coronoids with Different Inner and Outer Edge Structures

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ABSTRACT: Coronoids, polycyclic aromatic hydrocarbons with geometrically defined cavities, are promising model structures of porous graphene. Here, we report the on-surface synthesis of C168 and C140 coronoids, referred to as [6]- and [5]coronoid, respectively, using 5,9-dibromo-14-phenylbenzo[⌊]tetraphene as the precursor. These coronoids entail large cavities (>1 nm) with inner zigzag edges, distinct from their outer armchair edges. While [6]coronoid is planar, [5]coronoid is not. Low-temperature scanning tunneling microscopy/spectroscopy and noncontact atomic force microscopy unveil structural and electronic properties in accordance with those obtained from density functional theory calculations. Detailed analysis of ring current effects identifies the rings with the highest aromaticity of these coronoids, whose pattern matches their Clar structure. The pores of the obtained coronoids offer intriguing possibilities of further functionalization toward advanced host–guest applications.

Coronoids are macrocyclic conjugated hydrocarbons formed by circularly fused benzenoid rings, featuring a cavity.⁴ Kekule is a representative coronoid with a single layer of benzene rings around the cavity, belonging to the family of cycloarenes. The first synthesis of kekulene in 1978⁵ provided fundamental information about the nature of macrocyclic conjugation and aromaticity, pointing out that kekulene is better represented by Clar’s model rather than the so-called Kekulé structure. Other examples of cycloarenes are cyclo[de,de,e,e,d,e,e,d,e,e,e,d,e,e,d,e,e,d,e,e]decakisbenzene,⁷ predicted to have intriguing electronic and magnetic properties,⁸-¹⁰ septulene,¹¹ and octulene.¹² In 2016, we reported a C216 coronoid,¹ a macrocyclic molecule with 216 sp² carbon atoms distributed in more than one layer of circularly fused benzene rings (Figure 1). The C216 coronoid can also be regarded as a nanographene with a cavity arising upon removal of the central benzene ring from a C222 nanographene (Figure 1). Detailed characterization of C216 was compromised by its vanishing solubility, which also limited further synthetic studies of large coronoids. Very recently, Tan and co-workers reported a synthesis of mesityl-substituted C108 coronoid, which formed molecular bilayers.¹³

On the other hand, on-surface chemistry offers an alternative method to synthesize nanographenes and to explore their structure and electronic properties via state-of-the-art scanning probe microscopy and spectroscopy.¹⁴-¹⁵ Recently, kekulene was studied using noncontact atomic force microscopy (nc-AFM), which corroborated its structure in accordance with Clar’s model.¹⁶ Additionally, the on-surface synthesis of a C108 coronoid (Figure 1) was achieved,¹⁷ which represents a smaller homologue of the C216 coronoid. However, to date, no other coronoids have been synthesized on surfaces. Coronoids can also serve as models for graphene with nanoscale cavities, namely nanoporous graphene, which, depending on the size and structure of the pores, has intriguing electronic and magnetic properties.¹⁸-²⁰ Nanoporous graphene with zigzag-edge cavities stands out since it is related to nanographenes and graphene nanoribbons with zigzag edges.²¹,²² However, all reported coronoids have inner and outer armchair edges, and coronoids hosting a cavity with zigzag edges have never been synthesized.

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In this work, we targeted an exemplary coronoid (C168) with outer armchair edges and a large pore (1.45 nm, C–C distance) featuring inner zigzag edges. Conceptually, this molecule can be obtained via removal of a kekulene from the aforementioned C216 coronoid (Figure 1). Using S,9-dibromo-14-phenylbenzo[m]tetraphene (1) as a precursor, we achieved a thermally activated on-surface synthesis of the C168 coronoid through dehalogenative aryl–aryl coupling and subsequent cyclodehydrogenation on Au(111) (Figure 2a). C168 is obtained by the coupling of six molecules 1 and is hereafter called [6]coronoid. The on-surface synthesis also led to the formation of a coronoid consisting of five molecules 1, namely C140 ([5]coronoid), which adopts a strain-driven nonplanar conformation. The on-surface structural and electronic characterization of these coronoids was performed by scanning tunneling microscopy/spectroscopy (STM/STS) and nc-AFM, which were supplemented by density functional theory (DFT) calculations supporting the experimental results and shedding light on the aromaticity of [5]- and [6]-coronoids.

For the synthesis of [6]coronoid we have designed the U-shaped precursor 1, having a benzo[m]tetraphene core with a preinstalled zigzag edge as well as two bromo groups carefully positioned to allow for the macrocyclization on surface (see the SI for the synthesis of characterizations of 1). When 1 was deposited onto a Au(111) surface held at 200 °C, the homolytic cleavage of C–Br bonds readily took place, initiating aryl–aryl coupling.24 A temperature increase to 380 °C promoted the cyclodehydrogenation of the obtained nanostructures.25–27 The final products of this sequence consisted of various chainlike structures, as well as some macrocyclic molecules (Figure S1) including highly symmetrical ones (Figure 2b,f). The yields of [5]- and [6]coronoids could be increased by applying high-dilution conditions,28 with the highest yields of 30% and 6%, respectively, when the deposition rate of 1 onto Au(111) was set at 0.23 monolayer/hour (see Figure S2 for further details). The higher yield of [5]coronoid is most likely due to the ability of pentamic intermediates to undergo intramolecular head-to-tail coupling before the incorporation of the sixth building block, prevented by the high-dilution conditions.

To assess the exact chemical structure of the observed macrocycles, we acquired constant-height frequency-shift nc-AFM images using a CO-functionalized tip,29 which clearly revealed the formation of [6]- and [5]coronoid (Figure 2c,g). [6]Coronoid is planar on the Au(111) surface and displays the expected armchair/zigzag configurations of its outer/inner edges, as further confirmed by the excellent agreement of the experimental results with simulated nc-AFM images (Figure 2d) based on the DFT-optimized structural model (Figure 2e). The size of the inner cavity is measured to be 1.40 ± 0.05 nm (C–C distance of two opposing zigzag edges), in accordance with the DFT-calculated value of 1.45 nm (see the SI for details). The [5]coronoid also reveals an armchair/zigzag configuration of outer/inner edges (Figure 2f,g), while the size of the inner cavity is reduced to 1.10 ± 0.05 nm (diameter of the inscribed circle through the central carbon atoms of the zigzag edges measured from the nc-AFM data). Again, this value agrees with the pore size of 1.09 nm estimated from the DFT-optimized geometry of the [5]coronoid on Au(111) (Figure 2i). For the [5]coronoid, two out of five inner edges appear as brighter/higher features in both the STM and nc-AFM images (Figure 2f,g). Such a nonplanar structure is in line with a partial flattening of the more distorted gas-phase geometry of [5]coronoid (Figure S3b) upon adsorption. For quantification, we acquired z-dependent nc-AFM images (Figure 2j) and recorded the frequency shift (Δf) of the CO-functionalized tip while it approached the molecule to a specified location (Figure 2k). This allowed us to extract a height difference between the highest (blue) and lowest (red)
part of the molecule of 180 ± 10 pm, in good agreement with a DFT-calculated value of 170 pm (determined from the optimized geometry in Figure 2i). This height difference is substantially smaller than the value calculated for the [5]coronoid in the gas phase (360 pm, Figure S3b), which reflects the effect of van der Waals interactions with the Au(111) surface upon flattening the molecule.

To characterize the electronic properties of the [5]- and [6]coronoids, we performed constant-height differential conductance $dI/dV$ measurements and constant-current $dI/dV$ mappings at the detected positive (PIR) and negative ion resonances (NIR) (Figure 3a−j). Gas-phase orbital calculations (Figure 3k) show for [6]- and [5]coronoid, respectively, 6 and 5 close-lying frontier orbitals both for occupied and unoccupied levels. This is reflected in DFT calculations of the molecules on Au(111), where both the HOMO and LUMO are hybridized with the respective close-lying molecular orbitals into singular frontier ion resonances (see Figure S4 for details) that can be assigned to the experimentally measured PIR and NIR signatures. Hence, the experimental HOMO−LUMO gaps of the [6]- and [5]-coronoids on Au(111) are 2.2 and 2.5 eV, respectively. DFT calculations predicted the gas-phase HOMO−LUMO gaps of the [5]- and [6]coronoid, C216 and C222, to be 2.30, 2.25, 2.15, and 1.77 eV, respectively (Figure 3k), highlighting the trend of increasing gap with increasing pore size and decreasing number of carbon atoms.

The [6]coronoid could potentially constitute a building block for extended nanoporous graphene with larger cavities than those reported so far, and thus be of interest in the framework of graphene antidot lattices. We calculated the electronic properties of such hypothetical periodic layer via tight binding (TB) simulations (see Figure S5 for details), which reveal a band gap of 0.63 eV (between frontier bands localized at each cavity rim), considerably reduced compared to the [6]coronoid HOMO−LUMO gap (TB) of 1.51 eV.

Carbon macrocycles are known to have peculiar aromaticity properties, leading to intriguing concepts, such as superaromaticity, and represent an ideal playground for studying fundamental aspects of the π-electron clouds. To display the aromaticity pattern of the [6]- and [5]coronoid, we performed nucleus-independent chemical shift (NICS) calculations at 1 Å above the center of each carbon ring (NICS(1), Figure 4a,d). The results qualitatively match between similar locations of [6]- and [5]coronoids, indicating that the nonplanarity of the
latter does not significantly affect its aromaticity. Moreover, the NICS$_s$(1) patterns resemble the Clar structure of these coronoids (compare Figures 1 and 4), with each Clar sextet showing strong aromatic character (from $-22$ ppm to $-27$ ppm), similar to benzene ($-29$ ppm). The hexagons connecting the aromatic phenanthro[2,3,4,p-qrab]-perylene substructures (at the “bridges” between the fused precursor units) exhibit positive NICS value (16 ppm, Figure 4a,d) reminiscent of perylene. 35 In this regard, the [6]- and [5]-coronoids can be well represented by alternatingly fused perylene and pyrene constituents (Figure S6). To shed further light on the observed aromaticity pattern, we performed a harmonic oscillator model of aromaticity (HOMA) analysis (Figure 4b,e), which qualitatively matches the NICS$_s$(1) patterns of [6]- and [5]-coronoid. Further, we studied the induced currents due to an applied magnetic field by analyzing the anisotropy of the induced current density (ACID) (Figure 4c,f). The valence electrons are delocalized throughout the [6]- and [5]-coronoids. Each phenanthro[2,3,4,p-qrab]-perylene substructure displays a macroscopic diatropic current on its outer edge (red arrows in Figure 4c,f). Within these substructures, the most aromatic sextets exhibit local diatropic currents.

In conclusion, we have achieved the successful on-surface synthesis of large, planar [6]- and nonplanar [5]-coronoids featuring outer armchair and inner zigzag edges. Our study provides a complete characterization of their fundamental structural and electronic properties, revealing a HOMOLUMO gap of 2.2 and 2.5 eV on Au(111), respectively. Both coronoids present similar aromaticity pattern and homogeneously distributed π-electron clouds, and the location of the most aromatic sextets matches their Clar structure. The [6]- and [5]-coronoid have unprecedentedly large pores of 1.40 ± 0.05 and 1.10 ± 0.05 nm, respectively. Our synthetic strategy opens new avenues toward coronoids diversified outer and inner edge structures and offers a rich playground to investigate fundamental electronic properties of coronoids. Further, lateral fusion of these coronoids potentially leads to nanoporous graphene.

**ASSOCIATED CONTENT**

* Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c05268.

Methods, precursor synthesis and characterizations, additional computational details, and additional experimental and theoretical results (PDF)

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**Notes**

The authors declare no competing financial interest.

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