On-Surface Synthesis of a π-Extended Diaza[8]circulene

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ABSTRACT: Heterocyclic [8]circulenes are an important class of polycyclic aromatic hydrocarbon molecules because of their unique structural properties and promising applications. However, the synthesis of heterocyclic [8]circulenes is still limited and thus is an important synthetic challenge. Here we describe the first example of a π-extended diaza[8]circulene surrounded by and fused with six hexagons and two pentagons, which was successfully synthesized only by a combined in-solution and on-surface synthetic strategy. State-of-the-art scanning tunneling microscopy with a CO-functionalized tip and density functional theory calculations revealed its planar conformation and unique electronic structure.

[8]Circulene (1) (Figure 1) is a polycyclic aromatic hydrocarbon (PAH) molecule in which the central octagon is surrounded by and fused with eight benzene rings. Because of the high strain caused by the large sum of the wedge angles (480°), [8]circulene has a saddle-shaped structure and is expected to be used as a building block in bottom-up synthesis of negatively curved (nano)graphenes. Nevertheless, the synthesis of [8]circulene as well as its benzo-fused derivatives was achieved quite recently.

Heterocyclic [8]circulenes are [8]circulene molecules in which one or more of the benzene ring(s) are replaced by heteroaromatic ring(s) such as pyrrole or furan (Figure 1). They exhibit unique properties compared with the parent [8]circulene (1) and have received much attention because of their promising applications in organic semiconductors, organic light-emitting diodes, and liquid-crystalline materials. However, the variety of the core structure of heterocyclic [8]circulenes is still limited. Notable examples synthesized to date include compounds 2 bearing four hexagons (benzenes) and four pentagons (heteroles) and compounds 3 bearing eight pentagons. Their structural feature is their planar or quasi-planar structure due to the smaller sums of the wedge angles derived from pentagons. A key research target is the synthesis and structure of heterocyclic [8]circulenes having other combinations of hexagons and pentagons. Thus, we were interested in designing and synthesizing diaza[8]circulene 4, which would be the first example of a heterocyclic [8]circulene bearing six hexagons (benzenes) and two pentagons (heteroles).

The development of polycyclic aromatic azomethine ylides offers new pathways to nitrogen-containing PAHs. This 1,3-dipolar cycloaddition strategy enables the rapid construction of highly fused pyrrole structures and has been applied to the efficient synthesis of a variety of hitherto-unknown pyrrole-fused PAH molecules having planar bowl shapes as well as antiaromatic molecules. In order to construct core structures with even more varied shapes and properties, we employ on-surface synthesis, which has recently emerged as a powerful tool for the bottom-up synthesis of nanocarbon structures, as exemplified by graphene nanoribbons, carbon nanotubes, fullerenes, acenes, periacenes, and isomeric nanographenes. On-surface synthesis also provides an effective approach toward heteroatom-containing PAH molecules. Here we report a combined in-solution and on-surface synthesis of π-extended diaza[8]-circulene 4 bearing six hexagons and two pentagons. In particular, the final cyclodehydrogenation of a precursor molecule could be achieved only by on-surface synthesis on a Au(111) surface, as all of the other solution-based reaction attempts failed. Therefore, we developed a straightforward and effective method for the synthesis of π-extended diaza[8]-circulene 4 and studied its conformation, unique electronic structure, and intrinsic properties.

First, dibenz[a,e]cyclooctatetraene (5) (Scheme 1) was reacted with azomethine ylides formed from iminium salts 6. Although cycloadducts 7 can be isolated, the obtained crude mixtures were directly oxidized with 2,3-dichloro-5,6-dicyano-
p-benzoquinone (DDQ) to obtain monocycloadducts 8 in good yields. It is worth noting that the formation of dicycloadducts was not detected, although excess amounts of the azomethine ylides were reacted with 5 possessing two reactive carbon–carbon double bonds. Thus, it was necessary to repeat the same cycloaddition reaction to convert pyrroles 8 into diadducts 10 through pyrrolidine intermediates 9. All of the products were characterized by NMR spectroscopy and mass spectrometry (MS). In addition, single crystals suitable for X-ray diffraction analysis were obtained for compounds 7b, 8b, 9b, and 10b.24 The structure of 10b shown in Scheme 1 revealed 10 to have a tub-shaped structure.

In order to gain insight into the monoselectivity of the first 1,3-dipolar cycloaddition reaction, density functional theory (DFT) calculations at the B3LYP/6-31G(d) level were conducted (Scheme 2). In the first step, to form 7a, four possible stereoisomers derived from inner/outer and exo/endo structures need to be considered (Figure S24). Among them, the inner/exo addition gave the most stable transition state TS1 and the product 7a (Scheme 2A). The activation energy of the first transition state TS1 ($\Delta G^\ddagger$) was calculated to be 18.0 kcal/mol, while those of the other pathways exceeded 20 kcal/mol. This is consistent with our experimental results that only the inner/exo isomer was isolated. The activation energies of the second addition of the azomethine ylide to 7a ($\Delta G^\ddagger_2$) were also calculated (Scheme 2A). In this case, a reaction in an outer/exo manner gave the lowest energy of 31.7 kcal/mol. In contrast, the cycloaddition reaction to 8a gave a relatively
lower activation barrier ($\Delta G^\ddagger$) of 20.0 kcal/mol (Scheme 2B). The fact that $\Delta G^\ddagger$ is higher than $\Delta G^\ddagger$ could be the reason why the second addition did not occur even in the presence of an excess amount of azomethine ylide.

We devoted considerable efforts to convert 10a to 4a by a synthesis in solution, but all of those attempts failed. Therefore, we turned our focus to on-surface synthesis. Thus, 10a was deposited on Au(111) under ultrahigh vacuum at room temperature and then observed by scanning tunneling microscopy (STM) at 4.3 K (Figure 2A). The molecules were located at the elbow sites of the herringbone structure. In addition to the natural tub conformation of the octagon, the steric effect lifted the phenyl groups attached to the central octagon up from the substrate (Figure 2A inset and Figure S19). Subsequently, the sample was annealed at ca. 700 K to induce cyclodehydrogenation. Figure 2B shows the overview STM topography. It was found that some of the molecules became planar (Figure 2B inset), while some remained unchanged. Since the numbers of molecules on the surface observed before and after the annealing were almost the same, the molecular mass and the adsorption energy are high enough to prevent thermal desorption during annealing. To resolve the inner structures of the molecules, the metal tip was terminated with a CO molecule. Figure 2C,D shows the high-resolution CO-STM images in constant-height mode. Apparently, the dehydrogenative cyclization of 10a proceeded to form 4a (for the side products, see Figures S22 and S23). The pronounced sharp edges in the image confirm that the molecule on the Au(111) surface is planar. The small bilaterally asymmetric contrasts in the image may relate to asymmetric tilting of the CO tip (Figure S36). The high planarity of the central octagon allowed us to investigate the bond order variations via the apparent lengths in the CO-STM images (Figure 2C). To reduce the effect of the asymmetric tip, the mean values of the apparent lengths over equivalent sites of the central octagon in 4a were taken. We found a significant variation in the apparent lengths as 1.42, 1.56, and 1.69 Å (Figure 2D), indicating a certain variation in the bond orders as well. Our DFT calculations on Au(111) also showed a variation in the bond lengths (Figure 2E), although the differences were rather small. In order to investigate the validity of the comparison, we calculated the PP-STM image of the DFT-calculated structure. Indeed, the simulated PP-STM image (Figures 2F and S36) also shows a significant deviation in the apparent bond lengths, and the order of the apparent bond lengths is consistent with that in the experiment. Therefore, it is unambiguously concluded that the planarized octagon has a certain deviation of the bond order. In addition, the octagon looks highly distorted from a regular octagon, presumably because of the large sum of the wedge angles of the surrounding six hexagons and two pentagons. These results indicate that the central octagon is highly strained by the formation of the 4a structure as well as further planarization on Au(111). This steric frustration is also supported by DFT calculations on 4a in the gas phase, which indicated a twisted-shaped structure as the most stable state compared with the saddle-shaped (2.7 kcal/mol) and planar (12.7 kcal/mol) conformations (Scheme 1 and Figure S26).

Next we discuss the aromaticity because there has been continued interest in the aromaticity of cyclooctatetraene derivatives. Nucleus-independent chemical shift (NICS) calculations on 10a and 4a were performed in the gas phase. In 10a, the pyrrole rings and the peripheral benzene rings show large negative NICS(0) values (all in ppm) ranging from −7.7 to −16.0, while the central octagon shows a value of 3.6, which means that it is slightly antiaromatic (Figure S28). After conversion to 4a, the NICS(0) value increased to 7.4, which indicates an enhancement of the antiaromaticity upon planarization (Figure 3A). This NICS(0) value of 7.4 is comparable to those of other hetero[8]circulenes including sulfur and selenium (5.6−5.7). In order to get further insight, the anisotropy of the induced current density (ACID) of 4a was calculated in the gas phase (Figures 3B, S31, and S32). The plot clearly shows a counterclockwise $8\pi$ ring current flowing along the central octagon ring as well as a clockwise $4\pi$ ring current flowing along the outer fused rings, including the two pyrrole rings. The observation of a diatropic $4\pi$ ring current is consistent with those of other hetero[8]circulenes, which typically show internal paratropic $8\pi$ ring currents surrounded by outer diatropic $24\pi$ ring currents. Although the shapes of
the ring currents are not significantly affected by the change in the conformation of 4a (Figures S31 and S32), the efficiency of the diatropic $4\pi$ conjugation is maximized in the planar geometry, as the simulated $^1$H NMR signals in Figure S29 show larger shifts toward lower fields. Combined with the NICS calculations and bond lengths (Figure S27), these results suggested the presence of two major contributing electronic structures, one having two $6\pi$ pyrrole rings and eight $6\pi$ benzene rings (left in Figure 3C) and the other having inner $8\pi$, outer $40\pi$, and two $6\pi$ conjugations (right in Figure 3C). It should be noted that metal surfaces were not considered in the above NICS and ACID calculations, but we consider the discussion to be valid for 4a on the metal surface because of the similarity of the frontier orbitals in the gas phase and on the metal surface (as shown by comparison of Figures 4G and S35) and the consistency with the experimental $dI/dV$ curves (Figures 4D). In addition, the charge transfer plot (Figure S34) strongly suggested that the charge transfer between 4a and the metal surface is marginal.

Finally, we investigated the electronic structure of 4a on Au(111) with $dI/dV$ mapping at various energies (Figure 4A−C) and scanning tunneling spectroscopy at specific positions (Figure 4D). The lowest unoccupied molecular orbital (LUMO), at 2.2 V, was clearly observed at positions 1 and 2 in Figure 4D. This is consistent with the on-gold calculated simulation in Figure 4F and the vacuum-calculated LUMO in Figure 4G, in which the LUMO is mainly localized at the peripheral benzene rings. In contrast, the highest occupied molecular orbital (HOMO), at −0.9 V, and HOMO−1, at −1.2 V, were clearly observed at positions 3 and 4, which is also supported by the simulated HOMOs (Figure 4E,G). Thus, the HOMO−LUMO gap was determined to be 3.1 eV, which is consistent with the value of 3.08 eV calculated in vacuum (Figure 4G). The on-gold DFT calculations also show peaks for HOMOs and LUMOs near the Fermi Level, and their electron densities (Figure S35) are in agreement with gas-phase-calculated wave functions.

In summary, a combined in-solution and on-surface approach resulted in the successful synthesis of $\pi$-extended diaza[8]circulene 4. High-resolution CO-STM elucidated that the molecule adopts a planar conformation on Au(111) and has a unique electronic structure due to the introduction of multiple nitrogen atoms into the PAH framework. DFT calculations fully supported the experimental results. This work demonstrates the power of a combined in-solution and on-surface approach for the synthesis of unprecedented nitrogen-containing PAH molecules that could not otherwise be synthesized.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c02534.
Experimental procedures, compound characterization, spectra, details of DFT calculations, CO-STM images, and dI/dv simulations (PDF)
X-ray crystallographic data for 7b (CIF)
X-ray crystallographic data for 8b (CIF)
X-ray crystallographic data for 9b (CIF)
X-ray crystallographic data for 10b (CIF)

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Notes
The authors declare no competing financial interest.

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■ REFERENCES


(24) For details, see the Supporting Information.


