On-surface route for producing planar nanographenes with azulene moieties

Jeremy Hieulle,1 Eduard Carbonel-Sanromà,1 Manuel Vilas-Varela,2 Aran Garcia-Lekue,3,4 Enrique Guitián,2 Diego Peña,2 and Jose Ignacio Pascual1,4

1CIC nanoGUNE, 20018 San Sebastián-Donostia, Spain
2CIQUS, Universidad de Santiago de Compostela, Spain
3Donostia International Physics Center, 20018 Donostia-San Sebastian, Spain
4Ikerbasque, Basque Foundation for Science, Bilbao, Spain

Large aromatic carbon nanostructures are cornerstone materials due to their increasingly active role in functional devices, but their synthesis in solution encounters size and shape limitations. New on-surface strategies facilitate the synthesis of large and insoluble planar systems with atomic-scale precision. While dehydrogenation is usually the chemical zipping reaction building up large aromatic carbon structures, mostly benzenoid structures are being produced. Here, we report on a new cyclodehydration reaction transforming a sterically stressed precursor with conjoined cove regions into planar carbon platform by incorporating azulene moieties in their interior. Submolecular resolution STM is used to characterize this exotic large polycyclic aromatic compound on Au(111) yielding unprecedented insight into a dehydrogenative intramolecular ary1aryl coupling reaction. The resulting polycyclic aromatic carbon structure shows a [18]annulene core hosting peculiar pore states confined at the carbon cavity.

In recent years, solution chemistry has been successfully employed to obtain graphene nanostructures with fascinating structures following a bottom-up approach [1–4]. The precise control of their size and shape regarded these carbon nanostructures as technologically relevant materials in electronic and optoelectronic applications, in which large electron mobility and robust structure are stringent requirements [5, 6]. However, the large size and the extremely low solubility of these molecules severely complicates their manipulation and characterization by standard methods. Recently, combined submolecular atomic force microscopy (AFM) and scanning tunneling microscopy (STM) have emerged as methods to study insoluble nanographenes prepared by solution chemistry [7–10]. An important drawback of these methods is that they require sublimation of the material on an adequate surface, which limits the size of the molecules that can be studied. To overcome this limitation, an alternative approach relies on the preparation by solution chemistry of key building blocks that can be sublimated onto a surface, and act as precursors for the final on-surface preparation of the corresponding nanomaterial [11–15]. This approach is usually based on a sequence of thermally activated Ullmann couplings (UC) to polymerize dibromo-substituted monomers, followed by surface-induced cyclodehydrogenation reactions (CDH). The last CDH step is assisted by the metal surface to planarize the reaction products, turning molecular sites with high steric hindrance into benzenoid rings [16]. This protocol have been extensively used for the on-surface preparation of graphene nanoribbons (GNRs) [17, 18].

Based on this strategy, we aimed at combining UC reactions with a CDH cascade to produce large nanographene structures on a surface. In particular, we strived for covalently coupling three graphene blades into a trimer-like shape with a central [18]annulene pore (structure 3 in Figure 1) using the 12-ring dibromo polycyclic aromatic compound 1 as precursor. The [18]annulene pore is the smallest possible that can be created by means of C-C coupling strategies [20, 21], and the prediction is that it can host peculiar electronic features confined at the pore [22]. We found that the resulting structure 3 suffers strong intramolecular torsion at the carbon blades due to the presence of six conjoined cove regions with high steric hindrance. The structure spontaneously transforms into a new type of planar nanographene incorporating six azulene moieties (i.e. a seven-membered ring fused to a five-membered ring).

The structure of the produced nanographenes implies the discovery of a novel CDH reaction: while it is well documented the CDH of fjord cavities to form six-membered rings [16], as well as the CDH of cove regions to afford pentagons [19] (Fig. 1a), here we describe the CDH of conjoined cove regions (CCR) to obtain azulene fragments (Fig. 1a). Notably, since the azulene motif is the basic distortion of a graphenoid structure, named as Stone-Wales (SW) defect [23, 24], this novel CDH of conjoined cove regions provides a synthetic tool to introduce SW defects in nanographene structures.

The on-surface process followed two steps: first, a submonolayer amount of precursor 1 was sublimated under ultra-high vacuum conditions onto a clean Au(111) substrate (see Methods). During deposition, the metal sample was held at (200 °C) to favour the Ullmann-coupling polymerization of 1 upon arrival to the surface, rather than their packing in more complex π-stacked ensembles. Still, STM inspection of the sample after this intermediate step finds that the reacted monomers formed
FIG. 1: Bottom-up strategy to fabricate large carbon nanostructures: Chemical structure of the molecular precursor synthesized by aryne cycloadditions (see Supporting Note 1). Structure 2 is the expected nanographene after Ullmann coupling three precursors, to form an [18]annulene core, while structure 3 is the expected product after cyclodehydrogenation of the peripheral phenyls. Red arrows indicate the conjoined cove regions with high steric repulsion, described later in the manuscript. The new bonds after each step are highlighted in pink (UC) and in blue (CDH). Square A shows previous C-C zipping CDH reactions reported on a metal surface, creating six-membered rings [16] and five-membered rings [19] from fjord and cove regions, respectively. Square B shows the reported CDH reaction starting from a conjoined cove region, resulting in the formation of azulene moieties (dashed blue lines show new bonds).

closed-packed islands of different sizes and shapes (Figure 2a) but did not undergo CDH yet [16]. Second, the precovered sample (as in Fig. 2a) was annealed to temperatures around 375 °C. This converted the packed molecular islands into a disperse set of clusters of interconnected structures (Fig. 2b). The flat appearance of the structures in the STM images suggests that most of the polymers underwent a substrate-induced cyclodehydrogenation, which fused all their phenyl moieties into a graphenoid carbon structure (a very small amount of unreacted structures can still be distinguished).

A fraction of the reacted polymers corresponds to structures with a trimer-like shape, composed of three flat blades around a central pore, which we attribute to the [18]annulene core (Figure 2c). These shapes resemble the expected product of the two step-reaction, sequentially producing 3 C-C bonds by Ullmann coupling and 15 C-C bonds by intramolecular cyclodehydrogenation. However, a closer inspection reveals that although these trimers are flat they deviate from structure 3: they show a faint helical shape endowed by the small bend of all three blades with the same orientation with respect to the pore. Additionally, we observe other trimers even deviating from the C3-symmetric forms, in which one of the blades bends opposite to the others (Figure 2d). We did not find in our experiments any trimer with shape resembling 3, but only with (C3-symmetric) propeller shape and asymmetric trimers, as in Figs. 2c and 2d.
The observation of these unexpected shapes suggested that an alternative cyclization process occurred during the reaction.

To find out the origin of the asymmetric shapes, we resolved the intramolecular structure of the trimers with high-resolution STM images using a CO functionalized tip (see Methods). The presence of a CO molecule at the tip apex allows approaching the tip to the onset of repulsive Pauli forces and transduce variations of atomic-scale forces into changes of tunneling current and differential conductance \(dI/dV_s\) [25–29] with high resolution.

In figure 3a, we show a constant height \(dI/dV_s\) map of a C3-symmetric trimer, resolving with high detail its intramolecular ring structure. Each of the three planar blades is composed of 21 rings instead of the expected 17 rings in 3. Hence, the resulting trimers 4 possess in total 64 carbon rings, 12 more than compound 3. Furthermore, the conjoined cove regions of structure 3 are missing and, instead, two azulene moieties appear in each of the three blades.

Five- and seven-membered rings are typically observed as defects in the graphene structure [30–35], and appear commonly at the connecting boundaries between two graphene sheets with different alignment [36]. They are also allotropic forms of graphene, which maintain the \(sp^2\) character of the carbon sheet. In the current case, the two azulene moieties appearing inside each of the three planar blades show a well-defined and repetitive configuration, which refrain us from calling them defects. They always appear forming a perylene isomeric motif, where two azulenes are fused to the same benzene ring as depicted in Fig. 3b. We are not aware of previous synthesis strategies to produce this moiety, which turns out to endow flexibility to flat graphenoid sheets [23, 24].

The resulting azulene moieties are the origin of the propeller shape and asymmetric structure of the trimers. They show two possible (mirror-symmetric) orientations with respect to the axis of the carbon blade (the radius of the trimer), imposing the bending distortion apparent already in Figs. 2c and 2d. Three equally oriented azulene pairs in each blade lead to propeller-shaped C3-symmetric trimers 4 (Figure 3a), while a reversed orientation in one of the trimer blades affords the asymmetric structure 5 shown in Figure 3c. After STM inspection of the structure of 127 trimers, we found 27 species of the C3-symmetric form 4, while the rest are asymmetric trimers 5. The ratio of 4 vs. 5 species is close to the 1 to 3 proportion of possible configurations of each type. Therefore, the formation of azulene moieties in the core of the blades is not affected by the orientation of the neighbour blades in the trimer, indicating a non-cooperative reaction. Furthermore, the CDH reaction does not require the presence of contacting blades to succeed, since similar azulene moieties were found in monomers 1 as well as in dimers (see Supporting Note 3).

It is interesting to note that the 7-membered rings are
FIG. 3: High resolution STM images revealing a new flattening reaction: (a) Constant height $dI/dV$ map with a CO functionalized tip of a propeller shape C3-symmetric trimer (structure 4) showing its ring structure. Each of the three blades possesses two azulenes fused to a benzene ring (highlighted in colour in 4), which share the same orientation respect to the centre of the trimer. (b) Scheme showing the formation of two azulene moieties from two conjoined cove regions in each of the blades of 3, with new bonds indicated with dashed lines. (c) Same as in panel a, but for an asymmetric trimer 5. The high resolution image evidences that one of the blades is reversed with respect to 4, which is the cause of the asymmetry on the trimer. Scale bars are 1 nm ($V_s=5$ mV, $I_t=1$ nA, scale bar: 1.2 nm).

Imaged darker than the others, whereas the peripheral benzene rings appear brighter. Ring-specific current contrast appears to be linked to the Clar sextet structure of graphenoid systems [36]. The structure 4 accommodates nine Clar sextets in each blade, but only the six outer rings appear brighter in the images, which are the ones with probably less distortion. As we show in the Supporting Note 4, the different contrast can be well compared with the electrostatic potential landscape of the nanostructure [37, 38], appearing brighter in the images those rings with larger electron accumulation. This supports the association of ring contrast in constant-height
current images at $V \sim E_F/e$ to variations of electron density due to conjugation (as e.g. in Clar sextets) although other factors may also affect the tunneling transmission, such as vertical displacements and local density of states around the Fermi level.

The appearance of azulene moieties plays a key role in the planarization of the molecule. Compound 3 is heavily distorted in gas phase due to the overlap of five hydrogen atoms at each conjoined cove region, three of them lying directly over the same site. The steric repulsion induces an out-of-plane twist in the free compound 3 ($\sim 25$ degrees, see results of Force-Field simulations in Supporting Note 2), which hinders the planar adsorption on the surface. Compound 4 is on the contrary rather planar, and the azulene fragments simply endow the three blades some more flexibility than pure graphenoid structures [23, 24]. On a metal surface, the twisted structures 3 are forced to planarize by the combined effect of van der Waals forces and intramolecular restoring forces, both acting against steric hindrance between H atoms and weakening the C-H bonds. The catalytical role of the metal substrate at high temperatures results in the hydrogen removal [16].

Contrary to dehydrogenation reactions of fjord [16] and cove [19] cavities, the CDH of conjoined cove regions does not directly lead to cyclization, but additional molecular twists are required to create new C-C bonds. The deprotonated species is flat, flexible, thermally excited, and active to capture gold adatoms [39], which could facilitate the additional C-C coupling for azulene formation [40]. The energy gain of the reaction is clear: the incorporation of azulene moieties in the free molecule reduces the steric energy mostly due to the removal of the strong twist of compound 3 (Supporting Note 2). On the surface, the energy gain is considerably larger as the more planar structure 4 maximizes the contact with the gold surface.

We investigated the electronic structure of the produced trimers by means of differential conductance ($dI/dV_s$) imaging, exploring the extension of frontier orbitals. Point spectra acquired over the molecules (shown in the Supporting Note 3) found $dI/dV_s$ spectral features at -0.8 V and 1.2 V associated with HOMO and LUMO derived molecular states. Constant height maps of $dI/dV_s$ measured at these bias voltage values are shown in Figure 4a and 4b, respectively. The $dI/dV_s$ images show a characteristic intramolecular pattern extending all around the carbon platform rather than localized in some parts or edges, supporting the conjugated character of these states. To interpret these maps, we compared them with DFT simulations of the density of states (DOS) of the HOMO (Figure 4d) and LUMO (Figure 4e) states for a fully relaxed, C3-symmetric trimer 4 in the gas phase (see Supporting Note 4). The simulations confirm that the $dI/dV$ modulations in both cases are correlated with the lobes of the orbital wavefunction, which run as axial-wavefronts for the LUMO and tilted-waves for the HOMO.

Remarkably, a highly localized conductance feature emerges at the center of the [18]annulene core when the sample bias is tuned to e.g. 2.4 V (Figure 4c). The state appears with a peculiar three fold shape and with very little DOS weight in the rest of the carbon platform. Pore-localized states in non-covalent molecular architectures on surfaces have been detected due to the confinement of surface electrons [41–44]. The [18]annulene core is, on the contrary, a rigid structure arising from covalently bonded precursors. Our DFT calculations on the free species nicely reproduce the shape of this state (Figure 4f), including three smaller maxima at each of the three cavities at the joints between the blades. The pore state is thus an orbital of the trimer structure and can probably be depicted as a symmetry-adapted overlap of three specific blade orbitals [22, 45]. In fact, a precursor state can be already imaged for dimer structures appearing occasionally over the sample (see Supporting Note 3).

Interestingly, we find that the pore state belongs to the family of the so-called Super-Atom Molecular Orbitals (SAMOs [46]), representing the set of high-lying unoccupied orbitals with principal quantum number n ≥ 3 [47]. SAMOs have very little weight around the carbon cores, and extend well outside the molecular backbone, occupying the empty space in cavities and pores [47, 48]. Due to their large extension, SAMO-states are able to overlap with neighbours to form electronic bands in molecular crystals. In fact, the pore state shown in Fig. 4c is the result of the in-phase combination of three n=3 orbitals of the propeller blade, resulting in a tubular state crossing through the [18]annulene core (see Supporting Note 4).

In conclusion, we have shown the on-surface synthesis of a unique propeller-shape nanographene molecule with a [18]annulene pore and the remarkable formation of six azulene moieties at its three blades. The creation of azulene moieties follows a novel cyclodehydrogenation pattern in conjoined cove regions, which leads to two new C-C bonds and relaxation of the twisted regions into a flat-lying molecule on the surface. Our work thus highlights an additional reaction pathways allowed by the on-surface synthesis that can be exploited by clever precursor design, opening new possibilities in the growth of graphene-like structures. For example, the discovered reaction pattern suggests a control strategy to introduce Stone-Wales defects in graphene flakes or ribbons, as well as the synthesis of Haeckelites, which are proposed to endow metal character to flat $sp^2$ carbon flakes [23]. Furthermore, we found peculiar electronic states residing at the central pore of the trimers at high energies, which could be identified as SAMO-like resonances of the nanographene with the support of DFT simulations. These image-like states are expected to en-
FIG. 4: STS results and pore state of trimer 4: Constant height $dl/dVs$ maps picturing the shape of (a) the HOMO and (b) LUMO levels of an propeller shape C3-symmetric trimer 4. (c) Constant height $dl/dVs$ maps at higher energy, picturing a molecular state with large weight over the [18]annulene core (see also Supporting Note 4). (d,e,f) Isosurfaces of constant wave-function amplitude and phase (colour) obtained from DFT simulations of the free trimer 4. The orientation and number of modes of HOMO and LUMO orbitals (dashed black lines), as well as the shape of the pore state, shows remarkable agreement with the experimental maps (a:$I_t$=0.1 nA; b:$I_t$=0.07 nA; c:$I_t$=0.07 nA).

dow porous graphene with novel properties such as new electronic bands or enhance the photochemical reactivity of captured molecular guests.

**Organic synthesis:** of the organic synthesis of dibromo polycyclic aromatic compound 1 are given in the Supporting Note1.

**STM measurements:** The experiments were conducted in custom-made Low-Temperature (4.8 K) STM, under Ultra-High Vacuum conditions. We used a Au(111) single crystal as substrate, prepared by repetitive cycles of Ar$^+$ sputtering and annealing to 500°C, until the surface is atomically clean and flat. High resolution STM imaging was done employed CO-functionalized tips. For this purpose, small amounts of CO were dosed after the last reaction step on a sample. Analysis of STM and STS data was performed with the WsXm [49] and SpectraFox [50] (http://www.spectrafox.com) software packages.

**Computational details:** The electronic structure and geometry of compound 4, in its free standing form, were calculated using density functional theory, as implemented in the SIESTA code [51]. All the atoms of the molecule were fully relaxed until forces were $< 0.01$ eV/Å, and the dispersion interactions were taken into account by the non-local optB88-vdW functional [52]. The basis set consisted of doublezeta plus polarization (DZP) orbitals and diffuse 3s and 3p orbitals for C atoms, and DZP orbitals for H atoms. A cutoff of 300 Ry was used for the real-space grid integrations, and the -point approximation for sampling the three-dimensional Brillouin zone.

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Note 1: Organic synthesis of the molecular precursor

General Methods:

All reactions were carried out under argon using oven-dried glassware. TLC was performed on Merck silica gel 60 F254; chromatograms were visualized with UV light (254 and 360 nm). Flash column chromatography was performed on Merck silica gel 60 (ASTM 230-400 mesh). $^1$H and $^{13}$C NMR spectra were recorded at 300 and 75 MHz (Varian Mercury 300 instrument), respectively. Low-resolution electron impact mass spectra were determined at 70 eV on a HP-5988A instrument. High-resolution mass spectra (HRMS) were obtained on a Micromass Autospec spectrometer.

Molecules 6, 7 and 8 were prepared following published procedures [6]. Commercial reagents and anhydrous solvents were purchased from ABCR GmbH, Aldrich Chemical Co. or Strem Chemicals Inc., and were used without further purification. $n$-BuLi was used in solution in hexane (2.4 M).

Supporting Figure 1: Chemical structure of intermediate molecules 6, 7 and 8.
Synthesis of 3,6-dibromophenanthrene-9,10-dione (6):

To a solution of 9-10-phenantroquinone (9, 10.2 g, 49 mmol) and benzoyl peroxide (0.97 g, 4.00 mmol) in nitrobenzene (75 mL) Br₂ (5.1 mL, 100 mmol) was added over 30 min. at room temperature. The resulting mixture was heated at 110°C for 16 h. After cooling, the reaction crude was filtered and the residue was washed several times with hexane affording compound 6 (11.7 g, 66%) as a yellow solid. ¹H NMR (298 K, 300 MHz, CDCl₃) δ: 8.10 (d, J = 1.6 Hz, 2H), 8.05 (d, J = 8.3 Hz, 2H), 7.65 (dd, J = 8.3, 1.5 Hz, 2H) ppm.

![Supporting Figure 2: Synthesis of compound 6.](image)

Synthesis of 6,9-dibromo-1,3-diphenyl-2H-cyclopenta[l]phenanthren-2-one (7):

A mixture of compound 6 (1.00 g, 2.75 mmol) and 1,3-diphenylpropan-2-one (625 mg, 2.91 mmol) in MeOH (40 mL) was heated at 80 °C while a solution of KOH (160 mg, 2.85 mmol) in MeOH (4 mL) was slowly added. After addition, the mixture was refluxed for 30 min. and then cooled in an ice bath. The precipitate formed was filtered and washed with MeOH (4 x 20 mL) to yield compound 7 (1.13 g, 76%) as a brownish solid. ¹H NMR (298 K, 300 MHz, CD₂Cl₂) δ: 7.93 (d, J = 2.0 Hz, 2 H); 7.46-7.32 (m, 12 H); 7.11 (dd, J = 1.9 Hz, J = 8.6 Hz, 2 H) ppm.

Synthesis of 3,6-dibromo-9,14-diphenyl-12-(trimethylsilyl)benzo[f]tetraphen-11-yl trifluoro-methanesulfonate (10):

To a mixture of 7 (200 mg, 0.38 mmol) and 8 (220 mg, 0.42 mmol) in MeCN/CH₂Cl₂ (1:1, 28 mL) anhydrous CsF (84 mg, 0.55 mmol) was added. The resulting mixture was heated at 50°C for 16 h. Solvents were removed under vacuum and the residue was purified
Supporting Figure 3: Synthesis of compound 7.

by column chromatography (SiO$_2$; hexane/CH$_2$Cl$_2$ 4:1) to afford 10 (116 mg, 37%) as a greenish solid. $^1$H-NMR (300 MHz, CDCl$_3$) $\delta$: 8.34 (s, 1H), 8.33 (s, 1H), 8.14 (s, 1H), 7.86 (s, 1H), 7.62 7.50 (m, 10H), 7.44 (dd, $J = 9.0$, 1.9 Hz, 2H), 7.14 (d, $J = 8.9$ Hz, 2H), 0.35 (s, 9H). ppm. $^{13}$C-NMR (75 MHz, CDCl$_3$) $\delta$: 152.9 (C), 140.4 (C), 140.0 (C), 136.9 (CH), 135.9 (C), 135.7 (C), 133.0 (C), 132.7 (C), 132.6 (C), 132.2 (2CH), 132.03 (2CH), 131.97 (CH), 131.4 (C), 130.1 (C), 130.0 (C), 129.9 (C), 129.6 (2CH), 129.5 (2CH), 129.4 (2CH), 129.3 (2CH), 128.7 (C), 128.4 (CH), 128.3 (CH), 126.2 (2CH), 122.0 (C), 121.8 (C), 118.44 (q, $J = 320$ Hz,CF$_3$), 115.8 (C), -0.88 (3CH$_3$) ppm. MS (El) $m/z$ (%): 808 (M$^+$, 80), 675 (28), 660 (16), 596 (17), 424 (14), 243 (11), 73 (100). HRMS (El) for C$_{38}$H$_{27}$O$_3$F$_3$SiSBr$_2$, calculated: 805.9769, found: 805.9739.

Supporting Figure 4: Synthesis of compound 10.

Synthesis of 3,6-dibromo-9,11,12,13,14,16-hexaphenyldibenzo[a,c]tetracene (1).

To a mixture of 10 (75 mg, 0.09 mmol) and tetraphenylcyclopentadienone (30 mg, 0.08 mmol) in MeCN/CH$_2$Cl$_2$ (1:1, 4 mL), anhydrous CsF (42 mg, 0.28 mmol) was added. The resulting mixture was heated at 50°C for 16 h. Solvents were removed under vacuum and
the residue was purified by column chromatography (SiO₂; hexane/CH₂Cl₂ 3:2) to afford 1 (45 mg, 61%) as a yellow solid. 

\[ ^{1}H\text{-NMR (300 MHz, CDCl}_3\delta: 8.21 (s, 2H), 8.20 (s, 2H), 7.37 - 7.29 (m, 12H), 7.17 - 7.10 (m, 10H), 7.04 (dd, } J = 9.1, 1.5 \text{ Hz, 2H), 6.90 - 6.81 (m, 10H) ppm. } ^{13}\text{C-NMR (75 MHz, CDCl}_3\delta: 140.8 (2C), 140.5 (2C), 139.2 (2C), 138.8 (2C), 138.5 (2C), 136.0 (2C), 132.9 (2C), 132.4 (2CH), 132.1 (4CH), 131.3 (4CH), 131.1 (4CH), 131.0 (2C), 130.96 (2C), 130.6 (2C), 129.4 (2CH), 129.0 (4CH), 127.5 (2CH), 127.4 (4CH), 127.2 (2C), 126.6 (4CH), 126.2 (2CH), 125.8 (2CH), 125.4 (2CH), 121.4 (2C) ppm. MS (EI) } m/z (%): 941 (M⁺, 100), 863 (20), 569 (8), 414 (40). HRMS (EI) for C₆₂H₃₈Br₂, calculated: 940.1340, found: 940.1348.

**Supporting Figure 5:** Synthesis of compound 1, the molecular precursor used in this study.
Spectroscopic data

Supporting Figure 6: $^1$H and $^{13}$C NMR spectra of compound 10
Note 2: Molecular structure and steric energy

Figure 8 shows the minimum energy structure of the free compounds 3 and 4 obtained from molecular mechanics simulations using the MM2 Force Field implementation. This method provides the steric energy for each relaxed structure, as a measure of intramolecular

Supporting Figure 7: $^1$H and $^{13}$C NMR spectra of precursor molecule 1
distortion from an ideal $sp^2$ force field configuration. We found that both forms have similar steric energy (70 Kcal/mol in 3, 61 Kcal/mol in 4). However, while in 3 most of the energy comes from dihedral torsions, in 4 it comes from C-C bond stretch and bends. On a metal surface, planarization forces are expected to reduce the twist angle of compound 3, forcing the hydrogen atoms to approach against their mutual steric hindrance, until dehydrogenation reaction occurs at sufficiently elevated temperatures. The deprotonated radical compound is planar and with negligible steric energy. The formation of compound 4 from this point requires in-plane distortions, probably are mediated by gold adatoms, which are prone to bond to active sites in molecular species [1, 2].

Supporting Figure 8: (a,b) Force Field (MM2) relaxed structures of compounds 3 and 4, respectively. While 3 shows a strongly twisted shape, 4 appears flat after MM2 energy minimization. Yellow atoms in (a) point to the five H atoms suffering steric repulsion in the CC region. Blue atoms in (b) correspond to the new azulene moiety created after removal of four H atoms from each CC region.
Note 3: Scanning Tunneling Spectroscopy results:

dI/dV point spectra on trimer (4):

Scanning tunneling spectroscopy was performed over the trimer 4 to study their electronic configuration. We measured the differential conductance dI/dV using a lock-in amplifier. To detect frontier orbitals, we measured constant height spectra on the trimer blades (Fig. 9). The alignment of Highest Occupied and Lowest Unoccupied Molecular Orbitals (HOMO and LUMO) are detected as sudden increase in dI/dV signal at the corresponding bias alignment: -0.7 V and +1.2 V. The constant height dI/dV maps in Figure 4a and 4b of the main manuscript, were performed around the energy position of these two resonances.

Supporting Figure 9: HOMO and LUMO orbitals of trimer 4: (a) dI/dV spectrum taken at the center of a lobe of a trimer (black curve), showing a strong feature at +1.2 V associated to the LUMO orbital of the molecule (V_s=-1.0 V, I_t=0.3 nA). The spectrum of the bare Au(111) substrate is shown as a reference (green curve). Inset: constant current spectrum (I_t=0.2 nA) showing a resonance peak attributed to the HOMO orbitals of the trimer.
To explore high-lying molecular states (i.e. at large sample bias) we use constant-current dI/dV spectra (Figure 10a). The spectra show several features at high bias, specially when acquired over the [18]annulene core of the molecule, which are attributed to molecular states localized at the pore cavity, supported by constant-height dI/dV maps at the indicated bias values (Figure 10(b-e)).

**Supporting Figure 10:** STS constant current ($I_t=0.2$ nA) point spectra and corresponding dI/dV maps showing the evolution of the pore state of a symmetric trimer 4: (a) constant current dI/dV spectra taken at the pore (orange), at the covalent link (red). The spectrum of the Au(111) surface is shown as a reference (green); (b-e) constant height dI/dV maps taken with a metallic tip at the characteristic energies observed in a.
**dI/dV maps of dimer structures:**

A small amount of dimers were also found over the sample. Fig. 11 shows high resolution STM images and constant height dI/dV spectral maps of one of such structures. We note that at high bias, precursor states of pore resonances appear localized at the semi-cavities at the connecting point. This constitutes a proof that the pore states are indeed molecular states instead of confined surface states.

![Supporting Figure 11](image_url)

**Supporting Figure 11:** STM image (\(V_s=5\) mV, \(I_t=1\) nA) and corresponding dI/dV map of the dimer structure, taken with a CO functionalized tip. The conductance maps were taken in the constant height mode.
Note 4: Density Functional Simulations of free trimer structures

To interpret the electronic properties of the trimers we performed first-principle density functional theory (DFT) simulations of the atomic structure and electronic configuration of free-standing compound 4. High-resolution experimental observations allow us to conclude that the trimers adopt a planar conformation when adsorbed on the Au(111) surface. For this reason, despite not being the energetically most favorable conformation in gas phase, we have considered a planar trimer for our calculations, shown in Figure 12a.

Electrostatic Potential Energy Landscape:

The experimental high resolution images show a clear contrast, with some of the peripheral rings appearing brighter and the 7-membered rings darker (Fig. 3a of the manuscript). It has been proposed that such intramolecular contrast could be related to an inhomogeneous electron density in the aromatic platform, which eventually could be associated to the stabilization of a Clar sextet structure in the graphenoid system [3]. In turn, the inhomogeneous charge distribution affects the electrostatic potential landscape, which is known to have a significant influence on the features measured by high-resolution STM with a CO tip [4].

Supporting Figure 12: (a) Relaxed structure of free-standing compound 4, (b) charge density isosurface at 0.03 e/Å³ with a color range of [-272,272] meV, (c) constant height density of states at ~3.0Å, above the molecular plane.

In order to access to electrostatic potential energy landscapes, we simulated in Fig. 12b charge density isosurfaces of the free trimer 4. Clearly, a larger electron accumulation (more blue color) and, accordingly, a larger (more repulsive) electrostatic potential is observed at
the peripherial rings. In contrast, over the 7-membered rings we find less charge density (more red color). Such intramolecular differences can also be detected in constant height maps of the total density of states (Fig. 12c). Thus, our DFT results agree with the interpretation of the "electronic" origin of the contrast in the high-resolution STM images of the trimer.

**Pore super-atom states:**

DFT finds at high energy above the Fermi level a set of states with larger weight in the [18]annulene core and in the three semipores between the trimer blades. Figure 13a shows two of these states appearing at 3.33 eV (left) and 3.57 eV above E_F (right). In both cases, the wave function is even with respect to the carbon plane (Figure 13b). This proofs that the pore states found in the experiment are Super Atom Molecular Orbitals as defined by Hu and coworkers [5].

![Supporting Figure 13](image-url)

**Supporting Figure 13:** (a) Wavefunction amplitude isosurfaces (0.02 electrons/bohr^3) of two states, with large weight at the [18]annulene pore. The color indicates the relative phase. (b) Wavefunction amplitude isosurface (0.03 electrons/bohr^3) of the state at 3.57 eV showing the even parity of this state with respect to the carbon plane.
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