On-surface synthesis and characterization of nitrogen-substituted undecacenes

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Heteroatom substitution in acenes allows tailoring of their remarkable electronic properties, expected to include spin-polarization and magnetism for larger members of the acene family. Here, we present a strategy for the on-surface synthesis of three undecacene analogs substituted with four nitrogen atoms on an Au(111) substrate, by employing specifically designed diethano-bridged precursors. A similarly designed precursor is used to synthesize the pristine undecacene molecule. By comparing experimental features of scanning probe microscopy with ab initio simulations, we demonstrate that the ground state of the synthesized tetraazaundecacene has considerable open-shell character on Au(111). Additionally, we demonstrate that the electronegative nitrogen atoms induce a considerable shift in energy level alignment compared to the pristine undecacene, and that the introduction of hydro-aza groups causes local anti-aromaticity in the synthesized compounds. Our work provides access to the precise fabrication of nitrogen-substituted acenes and their analogs, potential building-blocks of organic electronics and spintronics, and a rich playground to explore π-electron correlation.
Acenes, polycyclic aromatic hydrocarbons (PAHs) consisting of linearly fused benzene rings, have attracted a great deal of interest for their remarkable electronic properties. As the length of the acene is increased, the energy gap between its highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) reduces rapidly, with already anthracene and tetracene (three and four linearly fused benzene rings, respectively) possessing semiconducting properties. The small electronic gap together with high-electron mobility renders acenes attractive materials in organic electronics, for applications such as organic field-effect transistors (OFETs), organic light-emitting diodes (OLEDs), organic photovoltaic (OPV) devices, and spintronic devices. Acenes have been shown to exhibit singlet fission, which would considerably increase OPV photoconversion efficiencies. In addition to the many potential applications, acenes, and especially higher acenes, are very good candidates to be open-shell polyradicals, while a recent quantum Monte Carlo study predicted a closed-shell ground state.

Heteroatom substitution is an effective method to tune the electronic, magnetic, and physico-chemical properties of graphene-based materials. When applied to acenes, replacing CH groups with heteroatoms (B, P, O, S, N) produces oligoheteroacenes, whose properties depend on the type, number, and position of the heteroatoms. For example, boron and nitrogen substitution has been shown to offer precise control over the radical character and chemical stability of acenes. If all the heteroatoms are nitrogen, the compounds are called azacenes (or N-heteroacenes). Azacenes inherit the remarkable properties of acenes due to the nitrogen atom leaving the topology of the π-system equivalent with respect to the substituted CH group. The electronegative nitrogen atoms allow for finely tunable frontier molecular orbital energy alignment. Azacenes have been shown to exhibit n-type semiconducting behavior and have been used in thin-film transistors, OLEDs, and in OPV devices.

Synthesis and characterization of larger acenes (from pentacene onwards) has been a challenge for solution chemistry due to their low solubility and high reactivity. Nevertheless, successful solution synthesis protocols are available for higher acenes by functionalizing them with stabilizing and protecting groups. Furthermore, the Strating–Zwanenburg reaction has been successfully used for the synthesis of hexacene to undecacene (except for decacene), although such compounds need to be stabilized in a matrix in order to keep them from decomposing after a few hours. In addition, solution chemistry has also been applied to successfully synthesize azapentacenes, azahexacenes, and azahexatetraacenes by utilizing various bulky protecting groups.

On-surface chemistry has emerged as an alternative, powerful method that allows synthesizing new organic materials by one- or two-dimensional spatial confinement on single-crystal substrates under ultrahigh vacuum (UHV) conditions. The molecular structure and electronic properties of the targeted nanomaterials can be identified via advanced scanning probe techniques, such as scanning tunneling microscopy/spectroscopy (STM/STS) and non-contact atomic force microscopy (nc-AFM). In the last decade, the research field of on-surface synthesis has experienced a considerable expansion and provided an appealing playground for the design and investigation of organic materials that have remained elusive due to synthetic limitations in solution chemistry. For instance, the on-surface synthesis of graphene nanoribbons (GNRs) has revealed the discovery of novel chemical reactions, or the synthesis of carbon-based nanographenes with intriguing electronic properties have significantly toward the development of the on-surface chemistry toolbox.

Particular attention has recently been paid to the synthesis and characterization of higher acenes on surfaces, with the largest synthesized members being decacene, undecacene, and dodecacene. Herein, the use of a-diketone-epoxy- and hydrogen-protected acene precursors, which provide them with the required solubility and stability, followed by thermal or light-induced activation once deposited on the metal substrate, have typically been the synthesis strategies employed for the successful formation of pristine higher acenes. Interestingly, the experimental frontier orbital gap for the series of acenes of increasing length is monotonically decreasing to around 1.0 eV for undecacene and unexpectedly increasing to around 1.4 eV for dodecacene, which the authors attribute either to a charge transfer or an electronic correlation effect.

Despite these successful cases exist for the synthesis of higher acenes, and despite the existence of on-surface approaches to synthesize heteroatom-substituted graphene-derived systems, the investigation of higher heteroacenes is so far unexplored.

Here, we introduce the on-surface formation of hydrogated tetraazaundecacene (N,N′N″,N‴-tetrahydro-6,12,19,25-tetraazaundecacene) 1, its analog with two edge-fused five-membered rings 2, and the 6,12,19,25-tetraazaundecacene 3 via thermal annealing and STM tip manipulation of a novel-design precursor molecule on a coinage metal surface under UHV conditions (Fig. 1). We also demonstrate that a similarly designed precursor can be used to synthesize the pristine undecacene 4. For this purpose, diethano- and diethano-bridged precursors (5 and 6, respectively), which are thermally stable upon sublimation, were synthesized in solution (see Supplementary Methods: Precursor Synthesis and Characterization).

In addition to the detailed chemical and structural properties unveiled via STM and nc-AFM measurements, we report on STS experiments to characterize the electronic structure of the synthesized compounds. In combination with theoretical calculations, this allows us to determine whether the obtained compounds present an open or closed-shell electronic ground state on the substrate. Additionally, ab initio computational methods are used to characterize and compare the electronic properties, such as the radical character, ionization potential, and aromaticity, of the synthesized compounds in detail.

Results and discussion

Deposition of precursor 5 on Au(111) and thermally induced transformations. In order to study the on-surface reactions described in Fig. 1, a submonolayer coverage of 5 was sublimed onto an Au(111) surface held at room temperature. Figure 2a shows a representative large-scale STM image revealing the presence of several rod-like self-assembled molecules mainly located at the face-centered cubic (fcc) regions of the Au(111) surface. High-resolution STM images (Fig. 2b) allow us to discern the intramolecular features of 5, where two bright protrusions per molecule of the similar apparent height of 2.3 Å (measured at a sample bias of −0.5 V) are observed. According to structural models, the self-assembled molecules are stabilized by N–H interactions between adjacent molecules with an experimental projected average distance of 3.2 ± 0.3 Å. Such experimental features, observed in Fig. 2b, are in good agreement with the DFT-optimized geometry and the corresponding STM simulation of 5 on the Au(111) surface (Fig. 2c, d), which indicate that 5 adopts a syn conformation with the two out-of-plane ethano bridges pointing upwards and an adsorption height of 2.8 Å with respect to the underlying surface.
Following the proven strategy for the formation of higher acenes on metallic surfaces, we have annealed the substrate at 280 °C, which induces the planarization of most of the molecular species, as judged from the absence of the bright protrusions of higher apparent height (Fig. 3a). This observation is attributed to the cleavage of the ethano protecting groups via retro-Diels–Alder reaction, observed for ~90% of the molecules, which coexist with species where one or two bright protrusion per molecule are still present. Notably, a statistical analysis of the sample (out of >200 molecules) shows that 25% of the new species appear as smooth rod-like structures, while the remaining molecules present one (33%) or two (42%) lateral protrusions. To characterize the chemical structure of the observed species, nc-AFM measurements using a CO-functionalized tip were performed.

![Diagram](image)

**Fig. 1 On-surface synthetic strategy.** Schematic representation of the synthesis of N,N',N'',N'''-tetrahydro-6,12,19,25-tetraazaundecacene 1, its analog with two edge-fused five-membered rings 2, 6,12,19,25-tetraazaundecacene 3, and the pristine undecacene 4 from the corresponding specifically designed precursors 5 and 6.

![Image](image)

**Fig. 2 Identification of the precursor 5 after deposition on Au(111).** a Large-scale STM topography image of the surface after room temperature deposition of 5. V_b = −1 V, I_t = 100 pA, scale bar = 5 nm. Δz denotes the relative height w.r.t. clean Au(111) surface. b Magnified view of one of the self-assembled structures shown in panel a containing three molecules. V_b = −0.5 V, I_t = 50 pA, scale bar = 1 nm. c Top and side views of the DFT equilibrium geometry of a cluster of three molecules 5 on Au(111). d DFT-simulated constant-current STM image of the three self-assembled molecules shown in panel a. V_b = −0.5 V, scale bar: 1 nm.
simulated nc-AFM image depicted in Fig. 3b, c confirms the successful formation of the hydrogenated tetraazaundecacene species 1. The corresponding DFT equilibrium geometry, shown in Fig. 3d, indicates physisorption with an adsorption height of 3.1 Å. Furthermore, Fig. 3e, f allow us to elucidate the chemical structure of the lateral protrusions (highlighted with a green arrow in Fig. 3a), which are attributed to five-membered rings (see Supplementary Fig. 1 for a zoom-in nc-AFM image of 2, where the newly formed five-membered rings are clearly discerned). They hypothetically derive from the bond cleavage at the bridgehead position of bicyclo[2.2.2]octadieno unit, with the terminal part of the resulting partial structure re-bonding to the neighboring nitrogen atom. These steps lead to the removal of two protons of the ethano group by dehydrogenative aromatization. Again, for product 2, the DFT equilibrium geometry shows physisorption with an adsorption height of 3.1 Å (Fig. 3g).

The observed hydrogenation of the edge nitrogen atom(s) might appear surprising at first sight, but it must be noted that the deprotection reaction takes place on a surface exhibiting a significant density of mobile hydrogen adatoms. Furthermore, under UHV conditions, hydrogen is the predominant compound in the rest gas of the vacuum chamber, and hydrogen atoms may also be produced by surface-promoted dissociation after cleavage of the ethano protecting groups. Similar passivation of reactive molecules or GNRs on surfaces has been reported before.

Next, we have probed the electronic structure of the obtained species 1 via scanning tunneling spectroscopy (STS) measurements. Figure 3h shows the voltage-dependent differential conductance spectrum (dIdV vs V) with peaks in the density of states at −0.45 V and +1.10 V, which we assigned to the positive and negative ion resonances (PIR and NIR, respectively). The spectrum has spurious peaks between −0.35 and −0.10 V, but these do not show features in the dIdV maps of the molecule and therefore are attributed to tip states (Supplementary Fig. 2). Using a CO-functionalized tip in constant-height mode, we mapped the dIdV spectra acquired on 1 at the energetic positions of the NIR, PIR, and two subsequent ion resonances detected at −0.80 V and −1.20 V, shown in Fig. 3i, left column, respectively from top to bottom. These dIdV maps show an excellent agreement with the DFT-calculated local density of states (LDOS) maps of the frontier orbitals (Fig. 3i, middle and right column) and correspond to the positive ion resonance peaks at −0.80 V and −1.20 V, respectively. The LDOS maps for this system are remarkably sensitive to height variation and therefore the simulation height matching best to the experimental data was chosen. For the height variation and comparison to experimental constant-current dIdV maps, see Supplementary Fig. 3. The assignment of the ion resonances to the molecular orbitals confirms the chemical structure of 1, and more specifically, the possibility of these species to be assigned to tetraazaundecacene 3 is ruled out at this stage by the presence of the HOMO−1/HOMO−2 resonance at −0.80 V (as this resonance is absent for 3, see Supplementary Fig. 4). The assignment of PIR and NIR to HOMO and LUMO allows us to infer an experimental HOMO-LUMO gap of 1.20 V.
Thus, tip-induced release of the protecting group diketone moieties equipped in heptacene and nonacene feedback loop off, as previously reported for the cleaving of by applying a voltage ramp (Fig. 4).

LUMO gap of 1.55 eV of 1 on Au(111), which is in good agreement with the theoretically predicted gap of 1.77 eV, calculated using many-body perturbation theory within the GW approximation, including screening effects from the underlying gold substrate (for details, see the Calculation methods section and Supplementary Methods: Computational Details).

**Synthesis of tetraazaundecacene by STM tip-induced deprotection.** Hydrogen passivation of the edge nitrogen atoms can be prevented if the annealing step is avoided and alternatively a tip-induced cleavage of the ethano bridges is applied. Figure 4a depicts the sequential tip-induced cleavage of the protecting groups from eight self-assembled molecules 1, which is achieved by applying a voltage ramp (−1.0 to −3.0 V) to the STM tip with feedback loop off, as previously reported for the cleaving of diketone moieties equipped in heptacene and nonacene precursors. Thus, tip-induced release of the protecting group can be employed to produce the targeted tetraazaundecacene species 3 (see Supplementary Fig. 5 for a comparison between experimental and simulated nc-AFM images between species 1 and 3). The resulting tetraazaundecacene molecules 3 remain self-assembled, stabilized by N···H interactions with identical average intermolecular distances as those described for the precursor 5 in Fig. 2c. The existence of N···H hydrogen bonds for 3, that enables the self-assembly, is also supported by analysis of the electrostatic potential (Supplementary Fig. 6), while the hydrogenated species 1 and 2, and the pristine undecacene 4 lack this interaction.

Figure 4b shows the DFT-optimized geometry of the five self-assembled tetraazaundecacene molecules 3 highlighted by a gray dashed rhombus in Fig. 4a, with the N···H interactions indicated by red ellipses. The DFT calculated adsorption height of 3 within this self-assembled cluster structure is 3.1 Å. Constant-height dI/dV maps acquired on a similar cluster of five self-assembled 3 (Fig. 4c) reveal the NIR at 0.90 V, PIR at −0.45 V and an additional resonance at −1.20 V. The experimental ion resonance maps are in good agreement with the DFT calculated LDOS maps (Fig. 4d, e), further confirming the successful synthesis of 3, and allow to assign the NIR and PIR to frontier orbitals and the resonance at −1.20 V to HOMO−1 (see Supplementary Fig. 4 for additional maps and calculated height dependence). The only qualitative difference between the LDOS maps calculated for the broken-symmetry open-shell singlet solution (Fig. 4d) and for the closed-shell solution (Fig. 4e) is the existence of the nodal plane along the center of the molecule in the HOMO map compared to the SOMO map. The corresponding experimental PIR map does not exhibit a reduction in the signal along the center, indicating that the experimental ground state has considerable open-shell character. The experimental frontier orbital gap, derived from the difference between PIR and NIR, is 1.35 eV. A GW calculation with image charge corrections for the open-shell ground state gives a gap value of 0.97 eV.
which matches better with the experiment than the closed-shell gap of 0.74 eV, thus giving further evidence for a significant open-shell character in the ground state of tetraazaundecacene on Au(111).

**Synthesis of pristine undecacene.** In addition to the diethano-bridged precursor 5 affording the tetraazaundecacenes discussed above, we have expanded our on-surface synthesis protocol toward the formation of pristine undecacene 4, taking advance of precursors containing diethano protecting groups (compound 6). Annealing of 6 on Au(111) to 220 °C results in a variety of products, including the targeted undecacene 4 (Supplementary Fig. 7). Experimental dI/dV maps acquired on 4 reveal PIR and NIR at −0.25 and 0.70 V, respectively, which can be assigned to frontier orbitals by comparing to DFT-simulated LDOS maps (Supplementary Fig. 7d–f). The experimental maps of the frontier orbitals show features of both, closed and open-shell calculated maps, which is again an indication of considerable open-shell character, similar to the case of tetraazaundecacene 3. This is in agreement with the demonstration of radial contributions in the ground state of higher acenes in literature and computational studies predicting that undecacene tilts more to the open-shell by DFT (Supplementary Table 1) with similar ionization potential predicted by state frontier gaps are 2.2 and 2.3 eV, respectively. Also, the gap for undecacene (difference between PIR and NIR) is 0.95 eV, in line with values reported in literature. This experimental gap matches very well with the GW (plus image charge screening correction) calculation in 1.03 eV, while the equivalent calculation for the closed-shell solution gives a gap of 0.70 eV, thus giving further evidence for the open-shell character of undecacene adsorbed on Au(111).

**Theoretical characterization of the synthesized compounds.** To compare the electronic structure of the synthesized compounds in further detail, we performed gas-phase DFT geometry optimization (Supplementary Fig. 8) and calculations of the ground spin state, ionization potential, and radical character (Supplementary Table 1). As expected, due to the equivalent topology of the π-system and electron filling in the undecacene 4 and the tetraazaundecacene 3, their electronic properties are practically equivalent except for the ionization potential and thus the energy alignment of the orbitals. For both systems, DFT with the B3LYP functional predicts an open-shell singlet ground state with the triplet state being 0.2 eV and the closed-shell state 0.5 eV higher in energy. Additionally, the ground state frontier gaps are ~1.7 eV, and unrestricted Hartree–Fock with spin-projection predicts a biradical character of 0.95 and a total of 3.7 unpaired electrons in both cases. The equivalence of the π-systems is also demonstrated in Supplementary Fig. 9, which shows that a wide range of frontier orbitals for the closed-shell solution are equivalent for 4 and 3. To quantify the orbital energy alignment, we performed ΔSCF calculations that predict the ionization potentials of undecacene 4 and tetraazaundecacene 3 to be 5.63 and 6.12 eV, respectively, showing that the electron-negative nitrogen atoms of tetraazaundecacene cause the electronic orbitals to be aligned 0.5 eV lower with respect to the vacancy level than those of pristine undecacene.

The hydrogenated species 1 and 2 are both predicted to be closed-shell by DFT (Supplementary Table 1) with similar electronic properties: triplet state energy is respectively +0.94 and +1.08 eV higher than the closed-shell state, and the ground state frontier gaps are 2.2 and 2.3 eV, respectively. Also, the ionization potential predicted by ΔSCF is similar for both: 5.47 and 5.42 eV for 1 and 2, respectively. Interestingly, the geometry of 1 and 2 is fully planar (Supplementary Fig. 8), which indicates that the nitrogen atoms do not break the π-conjugation and each of them provides 2 electrons to the π-system of the molecule. In the case of 1, this is also confirmed by comparing the DFT-predicted electronic orbitals to those of 4 and 3 (Supplementary Fig. 9). The orbitals of 1 exhibit the same nodal structure as those of 4 and 3, with 2 extra orbitals occupied by the 4 extra electrons provided by the NH groups. The lower-lying orbitals show the same energy-ordering for the three systems, but the extra occupations in 1 cause the orbitals from HOMO–2 (number 151) onwards to reorder in energy with respect to 4 and 3. Interestingly, the reordering happens such that the same orbital stays HOMO in all three cases.

In order to further characterize the π-system and the impact of the nitrogen substitution, we calculated the aromaticity pattern of the systems using the nucleus independent chemical shift (NICS) method. More specifically, we used the NICSzz(i) variant, which considers the out-of-plane component of the magnetic shielding tensor due to the π orbitals at 1 Å above the center of each ring. To characterize electron delocalization in the π-system and the aromatic ring currents, we performed anisotropy of the induced current density (ACID) calculations on the π orbitals of the systems (π-ACID). The results are shown in Fig. 5. The aromaticity patterns for the undecacene 4 and the tetraazaundecacene 3, calculated for the closed-shell solution, show a similar behavior: the central region of the acene is strongly aromatic and a strong diatropic current flows around the whole perimeter of the molecule. The open-shell solution for both cases, however, exhibits a positive NICSzz(i) value in the central region (Supplementary Fig. 10). This behavior has been reported in literature, although for shorter acenes and therefore showing a less significant effect. The positive NICSzz(i) value is an artificial effect induced by the contamination of the open-shell singlet ground state with the triplet state (Supplementary Table 1), which exhibits Baird’s anti-aromaticity. The π-ACID analysis for the open-shell singlet shows a diatropic current, although weaker than in the closed-shell solution, but still indicating aromaticity in the system. The hydrogenated tetraazaundecacene 1 and its analog with edge-fused five-membered rings 2 show almost equivalent NICSzz(i) pattern on the acene backbone (Fig. 5c, d, Supplementary Fig. 10 and 11), in line with the similarity of the electronic properties previously discussed, showing that the five-membered rings, which are aromatic themselves, have negligible effect on the aromatic pattern of the remaining system. Interestingly, for both systems, the cycles containing the nitrogen show sharp antiaromaticity. This further confirms the picture that the nitrogen atoms provide 2 electrons each to the π-system, making the nitrogen-containing cycles have 8 π electrons, which are formally anti-aromatic according to Hückel’s rule. The π-ACID analysis confirms this nicely: the rings containing the two nitrogen atoms show a clear paratropic contribution in the delocalized π-system, while the remaining parts show a diatropic current around their peripheries.

In conclusion, the on-surface synthesis of tetraazaundecacene, hydrogenated tetraazaundecacene and its analog with edge-fused five-membered rings reported in this article represents a fundamental step toward the precise engineering and tailoring of the unique electronic properties of the (hetero)acene family. Nc-AFM, STM, and STS differential conductance map unambiguously demonstrate the successful formation of the three targeted structures on an Au(111) surface. By comparing the experimental differential conductance maps and the frontier orbital gap with density functional theory LDOS maps and GW-calculated gap, we are able to assign an experimental ground state with considerable open-shell character for the synthesized tetraazaundecacene on Au(111), while the other two compounds are found to be closed-shell. Calculations based on DFT
Fig. 5 Computational characterization of the aromaticity of the synthesized undecacene and tetraazaundecacenes. **a** NICS\(_{(1)}\) patterns (each cycle is represented by the NICS\(_{(1)}\) evaluated at the center) and π-ACID plot (magnetic field is applied perpendicular to the plane of the system, pointing toward the reader; red arrows indicate aromatic diatropic and blue arrows anti-aromatic paratropic ring current) for the closed-shell solution of the undecacene 4. **b** NICS\(_{(1)}\) pattern and π-ACID plot for closed-shell solution of the tetraazaundecacene 3. **c** NICS\(_{(1)}\) pattern and π-ACID plot for the hydrogenated tetraazaundecacene 1. **d** NICS\(_{(1)}\) pattern and π-ACID plot for the hydrogenated tetraazaundecacene with edge-fused five-membered rings 2. All π-ACID plots are showing the isosurface at isovalue 0.05 a.u.

demonstrate that the tetraazaundecacene retains the remarkable electronic properties of the pristine undecacene with the exception of a considerable downshift in the orbital energies introduced by the electronegative nitrogen atoms. Additionally, characterization of the aromaticity shows that the hydrogenated tetraazaundecacene and its analog with edge-fused five-membered rings exhibit unique sharp anti-aromaticity in the rings containing the nitrogen atoms. Finally, the presented procedure offers a new perspective to investigate the complex electronic correlation effects in acenes by chemical substitution and opens up a new path toward precisely tailored building blocks of organic electronics and spintronics.

**Methods**

**Sample preparation and STM/nc-AFM measurements.** Experiments were performed under UHV conditions (base pressure below 4 × 10\(^{-10}\) mbar) with a commercial low-temperature STM/AFM from ScienT Instruments.

The Au(111) substrate was prepared by repeated cycles of Ar\(^{+}\) sputtering (E = 1 keV) and subsequent annealing to −450 °C for 10 min. All STM images shown were taken in constant current mode, unless otherwise noted, with electrochemically etched tungsten tips at a sample temperature of 4.8 K. Scanning parameters are specified in each figure caption. Molecular precursors (5 and 6) were thermally deposited onto a clean Au(111) surface held at room temperature with a typical deposition rate of 0.5 Å min\(^{-1}\) and 0.8 Å min\(^{-1}\) (sublimation temperatures −400 and 410 °C, respectively).

Nc-AFM measurements were performed with a tungsten tip attached to a Qplus tuning fork sensor. The tip was a posteriori functionalized by the controlled adsorption of a single CO molecule at the tip apex from the previously CO-dosed surface. The functionalized tip enables the imaging of the intramolecular structure of organic molecules. The sensor was driven at its resonance frequency (27.220 Hz) with a constant amplitude of ~70 pm. The shift in the resonance frequency of the tuning fork (with the attached CO-functionalized tip) was recorded in constant-height mode (Omicron Matrix electronics and HF2Li PLL by Zurich Instruments). The STM and nc-AFM images were analyzed using WSxM.

**Calculation methods.** The CP2K software package was used to calculate the equilibrium adsorption geometries and to perform the many-body perturbation theory calculations in the GW approximation. We used PBE functional with DFT-D3 dispersion corrections. The GW quasiparticle energy levels were renormalized according to the image charge model by Neaton et al. to account for the screening by the metal substrate. The AiiDAlab platform was used to perform the calculations for the molecules adsorbed on Au(111).

The gas-phase electronic structure, LDOS maps, and aromaticity calculations were performed with the Gaussian software package. The B3LYP functional was used in the spin-restricted and unrestricted formalism (respectively for the closed and open-shell states). The 6-311G** basis set was used for geometry optimizations, while single point and aromatic properties were calculated using the 6-311+G** basis set. The NICS\(_{(1)}\) was calculated with the GIAO-B3LYP method by applying the canonical molecular orbital (CMO) natural chemical shielding analysis (NCS) within the NBO program suite. The ACID calculations were performed with the CSGT method.

**Data availability**

The data that supports the results of this study are available on the Materials Cloud platform (https://doi.org/10.24435/materialscloud:mc-kj).

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Author contributions
J.I.U., H.H., R.F. conceived the experiments and C.A.P. and K.E. conceived the theoretical investigation. H.H., S.S., S.G., Y.S.C., N.A., and H.Y. synthesized the precursors. J.I.U. and M.D.G. performed the STM and nc-AFM experiments. K.E. performed the theoretical calculations. K.E., J.I.U., and C.A.P. analyzed the data. K.E. and J.I.U. co-wrote the manuscript. All authors discussed the results and commented on the manuscript.

Competing interests
The authors declare no competing interests.

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