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Site-specific Assembly of Fullerene Nanorings Guided by Two-dimensional Au Clusters

Yang-chun Xie¹, Mahroo Rokni Fard¹, Dogan Kaya¹, Deliang Bao^{2,3}, Richard. E. Palmer¹, Shixuan Du^{*2}, Quanmin Guo^{*1}

ABSTRACT

Molecules can self-assemble rather easily into straight one-dimensional wires via covalent or non-covalent bonding. To organize molecules into cyclic structures such as rings or loops where the number of molecules forming each structure is accurately controlled is a much more challenging task. Here we demonstrate the construction of Fullerene nanorings on the (111) plane of gold using single-atomic-layer-high gold islands as the seeding agent. C₆₀ molecules are trapped by the step edges of the seeding Au island leading to the formation of molecular rings. The smallest ring consists of six C₆₀ molecules encircling 19 Au atoms. Scanning tunneling microscope imaging reveals that the rings are formed at specific locations on the reconstructed Au(111) surface with the diameter of the ring controlled by the size of the gold island. Molecular mechanics modeling provides a detailed understanding of the relationship between the number of molecules in the ring and the number of gold atoms within the Au island.

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INTRODUCTION

The first manmade nanoring on a solid surface was created by manipulation of iron atoms on a copper surface using the scanning tunnelling microscope¹. The electron charge density oscillation within such a ring follows closely the prediction of quantum mechanics and for this reason this type of ring became known as a quantum corral. This pioneering work has inspired continued efforts to find new and more versatile routes to creating quantum corrals by means of supramolecular chemistry² or by directed surface assembly³. The selfassembly process such as that used by Klappenberger et al³ is a much more efficient approach than the original atom manipulation method¹ and is hence more promising for practical mass production. Selfassembly works well with a range of different materials such as colloids⁴, and metal nanoparticles⁵ as well as molecules⁶. The assembly process can be controlled rather accurately by altering the forces among the elemental blocks in the system⁷⁻¹⁴, or as demonstrated recently by changing the geometric shape of the building blocks¹⁵. Nanostructures achieved by molecular self-assembly on surfaces are usually in the form of large extended periodic structures of a repeating unit. To make isolated nanorings, one can choose a template to confine the growth of interesting structures to specific locations. Here we report the formation of Fullerene nanorings on gold surfaces where twodimensional Au clusters are used as the nucleation centres. The Fullerene nanorings resemble those formed naturally in photosynthetic complexes by Rsp. Photometricum cells¹⁶. Since Fullerenes are active ingredient in organic solar cells, the formation of rings may prove to be a significant step in producing solar cells with a structure mimic that of photosynthetic bacteria.

METHODS

Experimental

Experiments were performed in an ultra-high vacuum (UHV) system equipped for variable temperature scanning tunneling microscopy¹⁷⁻²⁰. The gold sample was a thin film of ~ 300 nm thick prepared by physical vapour deposition onto freshly cleaved highly oriented pyrolitic graphite (HOPG). The Au film was cleaned in the UHV chamber by many cycles of Ar+ ion sputtering and thermal annealing to 1000 K. C₆₀ molecules were deposited onto the sample using a homemade evaporation source. Au atom deposition is performed using a Createc high temperature effusion cell. An Omicron VT-STM was used for microscopic imaging of all structures. Electrochemically etched tungsten tips were used after initial heat treatment. Image processing was conducted using Image-SXM and Image J.

Computational

The molecule mechanics (MM) method is performed to simulate the Au-C₆₀ complex configurations. The COMPASS force field is employed to investigate the interactions between C₆₀ molecules and Au islands. The force field type for Au and C element is Au_m and c3a, respectively. As for six and seven C₆₀ molecules, the model is built with a 16×17 Au(111) surface containing three atomic layers, with the adsorbing Au island and C₆₀ molecules on one side of the slab. For more than seven C₆₀ molecules, the substrate expands to a 20×20 unit cell. The atoms in Au clusters are located at fcc region on the Au(111) surface. The models are not periodic. In relaxation, the slabs are all fixed, while all C atoms in C₆₀ molecules and Au atoms in clusters are fully relaxed until the system energy gradient is lower than 1×10⁻⁴ kcal/mol.

The density functional theory (DFT) calculation is also performed with Perdew-Burke-Ernzerhof (PBE) flavour of the generalized gradient approximation (GGA) in all-electron plane-wave formulation in the VASP code. An energy cut-off of 400 eV is used. The C₆₀-Au edge system is modelled by a 6×10 Au(111) surface containing three atomic layers of Au and a C₆₀ molecule adsorbed on one side of the slab. Some Au atoms in the layer closed to C₆₀ molecule is taken off to simulate an edge on the surface. Au atoms in the bottom two layers are fixed, while the edge layer and C₆₀ molecule are relaxed until the atomic forces are lower than 0.02 eV/Å. Van der Waals (vdW) interaction is considered with DFT-D2 method of Grimme in our calculation.

RESULTS AND DISCUSSION

Figure 1 illustrates the basic procedure implemented for the assembly of C₆₀ nanorings. First, Au atoms are deposited onto the Au(111) substrate which is kept at temperatures between 110 K and 150 K, giving rise to small single-atomic-layer-high Au islands. C₆₀ molecules are subsequently deposited onto the same surface while the sample remains at < 150 K. The step edges of the preformed Au islands thus become decorated by the C₆₀ molecules. The temperature of the sample is then increased gradually to room temperature to allow selfregulated growth and ripening of stable structures.

The Au(111) surface has a well-known herringbone-type of reconstruction which is characterized by a two-dimensional network of dislocations^{21,22}. The dislocation, analogous to the edge dislocation found in three-dimensional materials, is formed by the termination of an atomic row²³⁻²⁴. Fig. 2a shows an STM image of the clean Au(111) surface revealing the pattern of the herringbone reconstruction and

the presence of two types of elbow sites. Each elbow is associated with a defect. One side of the defect, where the extra row of Au atoms is located, is under compressive stress, and the opposite side is under tensile stress. The Au island nucleates at the defect and expands into the area under tensile stress. For the bulged elbow site, the Au island is thus found in the fcc domain, whereas the Au island is found in the hcp domain at the pinched elbow site as shown in Fig. 2b. The amount of gold atoms deposited in our study is low enough such that the Au island at the bulged elbow site does not spread into the neighbouring hcp region, and the Au island at the pinched elbow site does not spread into the fcc region. When C_{60} molecules are deposited onto the surface at 110 K-150 K, they diffuse until they are captured by the preformed Au islands resulting in the formation of irregular C60-Au complexes as shown in Fig. 2c. It can be seen that Au clusters are fully or partially surrounded by C_{60} molecules. The molecules appearing "bright" in the image are those sitting directly above the Au island. The location of the C₆₀-Au complex follows that of the pre-formed Au island: the C_{60} -Au complex is confined inside the fcc domain at the bulged elbow and inside the hcp domain at the pinched elbow, respectively.



Figure 1. Schematic of self-assembly of C_{60} -Au complex structures. Green circles represent C_{60} molecules, the large orange circle represents a two-dimensional Au island. The Au island is preformed by deposition of Au atoms onto Au(111) in the temperature range of 110 K-150 K. Subsequent deposition of C_{60} molecules conducted between 110 K and 150 K gives rise to C_{60} -decorated Au islands. By warming up to room temperature, two types of well-defined structures are formed: i) a C_{60} nanoring around the Au island; and ii) a compact island of C_{60} with some molecules sitting above the Au island.



Figure 2. Formation of C₆₀-Au complex structures at low temperatures. (a) STM image of the clean herringbone-reconstructed Au(111) surface. (b) Small Au islands formed at the elbow sites after deposition of 0.03 ML of Au at 120 K. Triangles drawn in the inset show the locations of the Au islands. (c) Structures formed after adding 0.015 ML of C₆₀ at 150 K onto a surface having pre-formed Au islands. Red circles highlight a few bare Au islands with no C₆₀ molecules attached.

The C₆₀-Au complex formed at 150 K as shown in Fig. 2c is far from uniform in terms of both size and shape. The number of C₆₀ molecules linked to a Au island ranges from 1 to about 15 with an average value of ~6. With just 0.015 ML of C₆₀ deposited, there is also a very small number of Au islands with no C₆₀ molecules attached. When the temperature of the sample is raised, not much change is observed below 240 K. Above 240 K, major structural transformations take place involving atom/molecule transportation across the surface. Bare Au islands with no C₆₀ attached are the least stable and the first to disappear. Au islands with only a small fraction of its step edges

covered by C_{60} molecules are the next to disintegrate. The Au atoms released from the fragmentation of these Au islands are incorporated into other islands. Similarly, there are also C_{60} molecules diffusing on the surface and they are ready to join nearby C_{60} -Au complexes.

In between 240 K and 290 K, rapid exchange of Au atoms and C_{60} molecules among the C60-Au complexes leads to the formation of two types of stable structures at room temperature, Fig. 3. One type of stable structures takes the form of compact islands with an exact number of C₆₀ molecules, known as magic number clusters; these are analyzed and reported in an earlier publication²⁵. Here we will concentrate on the alternative type of C_{60} -Au complex: the C_{60} nanoring formed around step edges of the Au island. In Fig. 3a, one can see full rings of C_{60} as well as partial rings. The full rings are mostly located in the fcc domains next to pinched elbow sites, with a small number also found inside the fcc domains far away from the elbow sites. There are Au islands straddling the discommensuration lines such that part of the island is inside the fcc domain and the rest inside the hcp domain. For this type of Au islands, the step edges inside the fcc domain are decorated by C₆₀ molecules, whilst the step edges within the hcp domain are exposed. Hence partial molecular rings are formed around these Au islands. Fig. 3b shows an image from a larger area demonstrating the spatial distribution of the two types of structures.

The structural transformation taking place between 240 K and 290 K can be summarized in the following way. At the bulged elbow sites, small compact C₆₀-Au hybrid clusters are formed²⁵ in the fcc domain. The fcc domain next to the bulged elbow site is relatively narrow, ~ 1.2 nm, and this puts an upper limit on the size of the Au island. At the pinched elbow site, Au atoms and C₆₀ molecules are able to move out of the hcp region as temperature is increased and they re-assemble at the adjacent fcc region, giving rise to either a compact C₆₀-Au cluster or a C₆₀ nanoring. The fcc region next to the pinched elbow is relatively wide, ~ 3.8 nm, hence it can accommodate larger Au islands. In Fig. 3b, the nanorings highlighted by white circles, consisting of about 15 molecules, are formed around Au islands that have grown to the maximum size allowed by the width of the fcc region next to the pinched elbow. The rings inside the yellow circles in Fig. 3 have not yet reached the maximum allowed size.



Figure 3. Formation of C_{60} -Au complex structures at room temperature. (a) STM image showing the presence of C_{60} nanorings at room temperature. (b) STM image from a larger area. Rings inside white circles consisting of ~ 15 molecules, are formed around Au islands already reached their maximum size allowed by the width of the fcc domain. Rings inside yellow circles are formed around Au islands yet to reach their maximum size. Structures inside white squares are partial rings formed around Au islands that straddle the hcp and fcc domains.

Figure 4 shows examples of nanorings observed at room temperature. Due to the limited number of rings observed, it is yet not possible to produce a reliable size distribution for the rings. The majority of rings have 6 to 15 molecules. The shape of the ring does not simply depend on the number of C_{60} molecules ($N_{C_{60}}$), for example, with eleven C_{60} molecules, five different shaped rings are shown in Fig. 4. It is likely that the shape of the ring also depends on the number of Au atoms (N_{Au}) enclosed. We performed molecular mechanics (MM) simulation to find out how the shape of the ring with a fixed number of molecules is influenced by the number of enclosed Au atoms.





The smallest ring consists of 6 molecules. The largest ring included in this set consists of 23 molecules (this is not the largest possible size). A ring with a given number of molecules, 11 for example, can have different shapes. The yellow arrow for the ring with 15 molecules points at a C_{60} molecule sitting further away from the step edge of the Au island than other molecules. The yellow arrow for the ring with 23 molecules points at a gap in the ring.

We first examined the ring with six C_{60} molecules. Figs. 5a and b show that for such a ring, the optimized configuration is that each molecule sits next to one side of a hexagonal Au island consisting of 19 Au atoms. The distance between adjacent molecules is about 1 nm. For the carbon atoms that are in direct contact with the Au surface, the C-Au distance is about 0.25 nm. Both values are consistent with those reported previously for C_{60} molecules on the same Au(111) surface²⁶. Figs. 5c and d show that for eleven C_{60} molecules, different shaped rings can be formed. With 75 Au atoms in the gold island, the molecular ring takes a slightly distorted hexagonal shape, Fig. 5c. With 64 Au atoms in the gold island, the same number, eleven, of C_{60} molecules form an elongated ring as shown in Fig. 5d. Therefore, it can be stated that for a fixed number of C_{60} molecules the shape of the ring is determined by the number of Au atoms enclosed by the molecules. Under certain conditions, the Au island happens to be the "wrong size" such that the perimeter of the Au island cannot support an integral number of C_{60} molecules while keeping the C_{60} - C_{60} distance at 1 nm. In such cases, molecules line up along the step edges of the gold island maintaining the 1nm distance between nearest neighbour molecules. In Fig. 5e, the perimeter of the Au island with 27 atoms can support 7 molecules but is too short for 8 molecules. When the eighth molecule is added to the existing structure, it stays at a larger distance from the step edge of the gold island. This kind of ring with a peripheral molecule has been observed in experiments as shown in Fig. 4 (ring with 15 molecules for example). The peripheral molecule is less strongly bound, and if this molecule moves away, a gap will occur in the molecular ring. Figs. 5g and h show 8-molecule rings with a gap. The Au island in Fig. 5h is 14 atoms larger than that in Fig. 5g and consequently the gap is larger. In Fig. 4, the STM image from the ring with 23 molecules shows such a gap which is not large enough to accommodate another molecule.



Figure 5. MM simulation of C₆₀ nano-ring formation. (a) Top view of a ring with six C₆₀ molecules and 19 Au atoms. (b) Side view of the 6-C₆₀ ring. (c) Ring with eleven molecules and 75 Au atoms. (d) Ring with eleven molecules but a reduced number, 64, of Au atoms. The changing number of Au atoms from 75 to 64 leads to a change in the geometry of the molecular ring. (e)-(h) The evolution of a nano-ring configuration by changing the number of Au atoms in an 8-C₆₀ ring. With 27 Au atoms, the perimeter of the Au island is not long enough to accommodate 8 molecules, hence one of the molecules in (e) is kept at a longer distance from the step edge. With 33 Au atoms, the perimeter of the Au island causes the formation of a gap in the molecular ring. The number in black at the top-right corner of each figure represents $N_{C_{60}}$, while the number in red represents N_{Au} in the gold island.

By assuming that the Au island always takes a hexagonal shape, which is a reasonable assumption for atomic islands on the (111) plane of fcc metals, we can determine N_{Au} rather precisely for a ring containing N- C_{60} molecules. The detailed method for calculating N_{Au} can be found in the Supplementary information. From experimental findings we know that the shape of the rings ranges from regular hexagons to elongated ones. Figure 6 shows the relationship between N_{AU} and $N_{C_{eq}}$. The red curve corresponds to the result calculated for regular hexagonal rings, while the green dashed line corresponds to that of the elongated hexagonal island. The degree of elongation is limited by the stability of the metal island which needs to minimize its total step energy. The shaded area between the two curves represents the possible values of N_{AU} and $N_{C_{EQ}}$ predicted by the model. For a number of experimentally observed rings consisting from 6 to 16 C_{60} molecules, the number of Au atoms in each ring has been evaluated using the relaxed model in Fig. 5 and Fig. S3. The outcome is shown in Fig. 6 with the filled squares. Therefore, the modeling provides a rather accurate evaluation of the number of gold atoms enclosed by the experimentally observed C_{60} rings. This overcomes the difficulty in direct counting the number atoms by STM.



Figure 6. The relationship between the number of C_{60} molecules ($N_{C_{60}}$) and the number of Au atoms enclosed (N_{Au}). The red curve is based on the model with a regular hexagon; whereas the green curve is based on an elongated hexagon. The formulae producing the curves are displayed in the figure (see details in the Supplementary Information).

The width of the fcc region next to a pinched elbow site is 3.8 nm. Experimental observations show that the rings are mostly confined within this fcc region. This suggests an upper limit for the number of Au atoms in the Au island of about 160. Consequently, the C_{60} ring associated with such a Au island in the fcc region consists of no more

than 15 molecules. Some rings are located far away from the elbow site, in regions where the discommensuration lines are straight. The width of the fcc domain in such regions is 2.7 nm. The Au island in the fcc domain far away from the elbows hence contains no more than ~110 Au atoms and 13 molecules are expected to decorate the step edges of the gold island. As can be seen in Fig. 3b, there exist local areas where fcc domains are larger than 3.8 nm due to the irregular bending of the discommensuration lines. This offers the opportunity to form a small number of molecular rings with more than 15 molecules.

Although the discommensuration lines tend to confine the Au island, they do not completely prevent the Au islands from growing to very large sizes. At RT and above, the Au island can expand by pushing further apart the discomensuration lines. Fig. 7 shows an STM image of a Au island which is 13 nm across. Au island of this size contains more than 1500 Au atoms and is clearly faceted. The usual herringbone reconstruction is observed within the island as revealed by the discommensuration lines^{27,28}, Fig. 7b. Because the amount of Au and C_{60} deposited on the surface is small, nearly all rings formed upon warming up the sample to room temperature are confined between two pairs of discommensuartion lines. Each Au island is protected by a ring of molecules. However, when the temperature of the sample is further increased to 400 K, some rings are observed to fragment. The atoms and molecules released from the fragmentation are captured by other stable rings resulting in the formation of the extra large rings such as that shown in Fig. 7.



Figure 7. STM image of a Au island 13 nm across. (a) The island is formed as a result of increasing the sample temperature to ~ 400 K. The island is faceted and its step edges consist of close-packed atoms. (b) Herringbone reconstruction takes place within the island. The image in (b) is the same as in (a) shown with enhanced contrast.

The nanorings are stable at room temperature as shown by sequential imaging of the same area over an extended period of time. Fig. 8 shows STM images of six nanorings observed during a 5 minutes interval. The rings keep their form rather well even for rings with irregular shapes. One of the rings, Figs. 8c and d, has a gap which is seen not to move with time. There might be edge diffusion of Au atoms occurring at the atomic level, but the overall morphology of the Au island is locked by the capping C_{60} molecules. The structure of the nanoring can be changed under certain conditions by the STM tip, for example, by using a larger bias voltage. Fig. 8i shows an image of a ring with 15 molecules. Fig. 8j shows an image of the same ring after scanning with an increased voltage of -2.46 V, which causes a clear change of the ring due to the re-organization of C_{60} molecules and Au atoms. The ring has also captured an extra molecule.



Figure 8. Stability of the nanorings at room temperature observed with the STM. (a), (c), (e) and (g) show the morphologies of the initial rings. (b), (d), (f) and (h) shows the same rings after 5 minutes at room temperature. The non-circular shape of the rings allows any change to be easily identifiable. (i) and (j) are STM images demonstrating the change of the ring due to the perturbation by the STM tip.

In all the experiments performed so far, the nanorings were always found to co-exist with compact magic number C₆₀-Au clusters. The fraction of the nanorings can be changed by altering the parameters used for deposition. For example, when more Au atoms are deposited to form relatively large bare gold clusters, the number of rings increases significantly. Fig. 9a shows an STM image after 0.09 ML of Au and 0.17 ML of C₆₀ are deposited at 150 K and warmed up the sample to room temperature. More than 50% of the features observed afterwards are rings. This is much higher than ~5% of rings found in Fig. 3 when smaller amounts of Au atoms and C₆₀ molecules are deposited. Compact clusters can also be transformed into rings by adding more Au atoms as demonstrated by the STM images in Figs. 9b and c. In Fig. 9b, the two clusters inside the white circles are identical clusters, each contains 14 C₆₀ molecules. Four of the fourteen molecules sit on the Au island. After the image in Fig. 9b was collected, 0.003 ML of Au is deposited onto the sample at RT. Thus, some Au atoms join the 14-molecule clusters. The four bright molecules are found to jump down each of the Au islands and become part of the molecular ring surrounding the now enlarged Au islands, Fig. 9c. During the structural transformation into rings, the cluster near the top-left corner mass-migrates collectively to a new location where the ring is formed. This collective migration suggests a cohesive interaction that keeps the atoms and molecules in a group as they travel on the surface. It is not likely that the whole cluster would move as an intact unit. Instead, the net displacement is accomplished probably by the movements of individual atoms/molecules in a loosely bound group.



Figure 9. Scheme to increase the proportion of nanorings. (a) The fraction of nanorings is increased significantly when the amounts of Au and C₆₀ deposited at 150 K are increased. The numbers next to the rings are the number of C₆₀ molecules in the ring. Transformation from compact clusters to nanorings is demonstrated in (b) - (e). In (b) the two compact clusters within white circles are identical clusters each consisting of fourteen C₆₀ molecules. Four of the fourteen molecules sit directly above the Au island. These two clusters capture more Au atoms at room temperature and change into rings (c). The ring on the left is found at a different location from that of the initial compact cluster, indicating a collective migration of molecules and atoms during the formation of the ring. The yellow arrow in (d) shows the direction of displacement of the C₆₀-Au complex. The images in (b) and (c) are contrast enhanced and displaced in (d) and (e), respectively, to show the locations of the C₆₀-Au complexes relative to the locations of the discommensuration lines.

SUMMARY

Molecular rings of C_{60} , resembling the structures formed naturally in photosynthetic complexes by *Rsp. Photometricum* cells, have been produced on the Au(111) surface. Both the size and location of the rings can be controlled by choosing suitable preparation procedures. Such molecular rings, encapsulating a metal or semiconductor core have the potential to be developed into building blocks for new types of solar cells based on efficient charge transfer between the metal and the molecule²⁹ or high-density magnetic recording medium if the metal core is formed by magnetic materials.

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SUPPORTING INFORMATION

Supplementary Information on theoretical modeling accompanies this paper.

AUTHOR CONTRIBUTIONS

Y-C Xie, M Rokni Fard, D Kaya and D-L Bao made equal contributions to the manuscript.

REFERENCES

- 1. Crommie, M. F.; Lutz, C. P.; Eigler, D. M. Confinement of electrons to quantum corrals on a metal surface. *Science* **1993**, 262, 218-220.
- Kondratuk, D. V.; Perdigao, L. A.; Esmail, A. M. S.; Oshea. J. N.; Beton, P. H.; Anderson, H. L. Supramolecular nesting of cyclic polymers. *Nature Chemistry* 2015, 7, 317-322.
- Klappenberger, F.; Kuhne, D.; Krenner, W.; Silanes, I.; Arnau, A.; Garcia de Abajo, F. J.; Klyatskaya, S.; Ruben, M.; Barth, J. V. Dichotomous array of chiral quantum corrals by self-assembled nanoporous Kagome network. Nano Lett. 2009, 9, 3509-3514.
- 4. Kim, P. Y.; Oh, J. W.; Nam, J. M. Controlled co-assembly of nanoparticles and polymer into ultra long and continuous onedimensional nanochains. J. Am. Chem. Soc. **2015**, 137, 8030-8033.
- 5. Gorgoll, R. M.; Tsubota, T.; Harano, K.; Nakamura, E. Cooperative self-assembly of gold nanoparticles on the hydrophobic surface of vesicles in water. J. Am. Chem. Soc. **2015**, 137, 7568-7571.
- Heim, D.; Ecija, D.; Seutert, K.; Auwarter, W.; Aurisicchio, C.; Fabbro, C.; Bonifazi, D.; Barth, J. V. Self-assembly of flexible onedimensional coordination polymers on metal surfaces. J. Am. Chem. Soc. 2010, 132, 6783-6790.
- 7. Barth, J. V. Molecular architectonic on metal surfaces. Annu. Rev. Phys. Chem. 2007, 58, 375-407.
- Rosei, F.; Schnack, M.; Naitosh, Y.; Jiang, P.; Gourdo, A., Lagsgaard, I.; Stensgaard, I.; Jochim, C.; Besenbacher, F. Properties of large organic molecules on metal surfaces. *Prog. Surf. Sci.* 2003, 71, 95-146.
- Theobald, J. A.; Oxtoby, N. S.; Philips, M. A.; Champness, N. R.; Beton, P. H. Controlling molecular deposition and layer structure with supramolecular surface assemblies. *Nature* (London) 2003, 424, 1029-1031.

- 10. Pawin, G.; Wong, K-L.; Kwon, K-Y.; Bartels, L. A homomolecular porous network at a Cu(111) surface. *Science* **2006**, *313*, 961-962.
- 11. Pivetta, M.; Pacchioni, G. E.; Schlickum, U.; Barth, J. V.; Brune, H. Formation of Fe cluster superlattice in a metal-organic quantumbox network. *Phys. Rev. Lett.* **2013**,110, 086102.
- Hanke, F.; Haq, S.; Raval, R.; Persson, M. Heat-to-connect: Surface commensurability directs organometallic onedimensional self-assembly. ACS Nano 2011, 5, 9093-9103.
- 13. Ecija, D.; Urgel, J. I.; Papageorgiu, A. C.; Joshi, S.; Auwarter, W.; Seitsonen, A. P.; Klyatskaya, S.; Ruben, M.; Barth, J. V. Five-vertex Archimedean surface tessellation by lanthanide-directed molecular self-assembly. *Proc. Natl. Acad. Sci. U.S.A.* **2013**,110, 6678-6681.
- Fernandez-Torrente, I.; Monturet, S.; Franke, K. J.; Fraxedas, J.; Lorrente, N.; Pascual, J. I. Long-range repulsive interaction between molecules on a metal surface induced by charge transfer. *Phys. Rev. Lett.* 2007, 99, 176103.
- 15.Lu, F.; Yager, K. G.; Zhang, Y-G.; Xin, H-L.; Gang, O. Superlattices assembled through shape-induced directional binding. *Nature Comm.* **2015**, 6, 6912.
- 16. Scheuring, S.; Sturgis, J. N. Chromatic adaptation of photosynthetic membranes. *Science* **2005**, 309, 484-487.
- 17. Tang, L.; Xie, Y-C.; Guo, Q. Complex orientational ordering of C₆₀ molecules on Au(111). J. Chem. Phys. **2011**, 135, 114702.
- Xie, Y-C.; Tang, L.; Guo, Q. Underlayer growth of a nano-porous single atomic layer of gold. J. Phys. Chem. C. 2012, 116, 5103-5109.
- 19.Li, F. S.; Tang, L.; Zhou, W. C.; Guo, Q. Resolving the Au-adatomalkanethiolate bonding site on Au(111) with domain boundary imaging using high-resolution scanning tunneling microscopy. J. Am. Chem. Soc. **2010**, 132, 13059-13063.
- 20. Zhang, X.; Tang, L.; Guo, Q. Low temperature adsorption and growth of C₆₀ monolayers on Au(111). J. Phys. Chem. C. **2010**, 114, 6433-6439.
- 21. Scheuring, S.; Sturgis, J. N. Chromatic adaptation of photosynthetic membranes. *Science* **2005**, 309, 484-487.
- 22. Barth, J. V.; Brune, H.; Ertl, G.; Behm, R. Scanning tunneling microscopy observations on the reconstructed Au(111) surface. *Phys. Rev. B.* **1990**, *42*, 9307-9318.
- 23. Harten, U.,; Lahee, A.; Toennis, J.; Woll, C. Observation of a soliton reconstruction of Au(111) by high-resolution helium-atom scattering. *Phys. Rev. Lett.* **1985**, *54*, 2619-2622.
- 24. Chambliss, D. D.; Wilson, R. J.,; Chiang, S. Nucleation of ordered Ni island arrays on Au(111) by surface-lattice dislocations. *Phys. Rev. Lett.* **1991**, 66, 1721-1724.
- 25. Xie, Y-C.; Tang, L.; Guo, Q. Co-operative assembly of magic number C₆₀-Au complexes. *Phys. Rev. Lett.* **2013**, 111, 186101.

- 26. Wang, L-L.; Cheng, H-P. Density functional study of the adsorption of a C₆₀ monolayer on Ag(111) and Au(111) surfaces. *Phys. Rev. B* **2004**, 69, 165417.
- 27.Sun, J-T.; Gao, L.,; He, X-B.; Cheng, Z-H.; Deng, Z-T.; Lin, X.; Hu, H.; Du, S-X.; Liu, F.; Gao, H-J. Surface reconstruction transition of metals induced by molecular adsorption. *Phys. Rev. B* 2011, 83, 115419.
- 28. El-Batanouny, M.; Burdick, S.; Martini, K.; Stancioff, P. Double sine-Gordon solitons- A model for misfit dislocations on the Au(111) reconstructed surface. *Phys. Rev. Lett.* **1987**, *58*, 2762-2765.
- 29. Hunt, M. R. C.; Modesti, S.; Rudolf, P.; Palmer. R. E. Chargetransfer and structure in C₆₀ adsorption on metal surfaces. *Phys. Rev. B* **1995**, *51*, 10039-10047.

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