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## Tunable Quantum Dot Arrays Formed from Self-Assembled Metal-Organic Networks

F. Klappenberger, <sup>1,\*</sup> D. Kühne, <sup>1</sup> W. Krenner, <sup>1</sup> I. Silanes, <sup>2,3</sup> A. Arnau, <sup>2,4,5</sup> F. J. García de Abajo, <sup>6</sup> S. Klyatskaya, <sup>7,1</sup> M. Ruben, <sup>7,8</sup> and J. V. Barth <sup>1</sup>

<sup>1</sup>Physik Department E20, TU München, 85748 Garching, Germany

<sup>2</sup>Donostia International Physics Center (DIPC), 20018 San Sebastian, Spain

<sup>3</sup>Instituto de Hidráulica Ambiental de Cantabria (IH), 39005 Santander, Spain

<sup>4</sup>Centro de Fisica de Materiales CSIC-UPV/EHU, Materials Physics Center MPC, 20080 San Sebastian, Spain

<sup>5</sup>Depto. Fisica de Materiales UPV/EHU, Facultad de Quimica, 20080 San Sebastian, Spain

<sup>6</sup>Instituto de Óptica—CSIC, Serrano 121, 28006 Madrid, Spain

<sup>7</sup>Institut für Nanotechnologie, Karlsruher Institut für Technologie (KIT), 76344 Eggenstein-Leopoldshafen, Germany

<sup>8</sup>IPCMS-CNRS, Université de Strasbourg, F-67034 Strasbourg, France

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The confinement of Ag(111) surface-state electrons by self-assembled, nanoporous metal-organic networks is studied using low-temperature scanning tunneling microscopy and spectroscopy as well as electronic structure calculations. The honeycomb networks of Co metal centers and dicarbonitrile-oligophenyl linkers induce surface resonance states confined in the cavities with a tunable energy level alignment. We find that electron scattering is repulsive on the molecules and weakly attractive on Co. The tailored networks represent periodic arrays of uniform and coupled quantum dots.

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The controlled adjustment of materials properties by modification on the atomic scale is a major goal of nanoscale science. On the close-packed faces of noble metals the surface-state electrons are well suited to monitor and influence the properties of the surface [1]. In 1993 Crommie et al. demonstrated the engineering of quantum well states by the manipulation of individual adatoms [2]. Since then the quasiparticle excitations of the surface-state band have been studied with great interest both experimentally [3–8] and theoretically [9–11], especially because their lifetime is connected to elementary scattering processes [10,11]. The high level of control over local electronic properties achievable by atom manipulation was demonstrated in the quantum mirage effect [12], in the controlled modification of the electronic structure of an adsorbate [13], and in quantum holographic encoding [14].

However, for the design of a targeted electronic spectrum of an *entire* surface it is impractical to use atom manipulation due to the slow serial production process. Since not only metal adatoms but also organic molecules scatter the surface-state electron waves [15], the extended regular structures that can be produced with molecular self-assembly [16] allow the tuning of electronic properties not only locally but surface wide, for example, in between linear molecular lines [17]. An elaborate electronic structure has been bestowed to Ag(111) by a chiral kagomé network providing a 2D periodic array of quantum dots [18]. With the reflectivity for the electron waves being finite, the leakage-induced electronic overlap between neighboring quantum dots results in dispersive bands [19].

Here, we use supra-molecular design to construct metalorganic networks providing regular hexagonal cavities on the Ag(111) metal surface [20,21]. The obtained roomtemperature stable, nanoporous arrays create a novel periodic structure to the surface-state electrons, in which the energy positions of the different resonances depend on the lateral dimensions of the confinement region. These dimensions are easily tunable by the length of the linking molecule. The electronic characteristics of the confined states have been reproduced both in energy and real space by Green's functions based electronic structure calculations using a boundary element method. Our analysis indicates that the molecules produce a repulsive scattering potential whereas the Co potential is slightly attractive. Thus the coordination bond markedly alters the scattering properties of the Co atoms. The crystal-like quality of our networks provides a perfect template for the lateral engineering of surface states [22].

The experiments were carried out in a vacuum apparatus, where a clean Ag(111) surface was prepared by a standard procedure. Metal-organic networks of Co and dicarbonitrile-quaterphenyl (NC-Ph<sub>4</sub>-CN), respectively, dicarbonitrile-sexiphenyl (NC-Ph<sub>6</sub>-CN) molecules were produced as described earlier [20,21]. Posteriorly, the sample was transferred into our homemade [23] beetle-type low-temperature scanning tunneling microscope (STM) where data were recorded at  $\approx 8$  K. Scanning tunneling spectroscopy (STS) was carried out by open-feedback loop dI/dV point spectra with bias modulation at ~1400 Hz, an amplitude of 5 mV rms, and a lock-in time constant of 20 to 50 ms. The dI/dV maps were extracted from a set of 84  $\times$ 84 normalized point spectra. The spectroscopic maps are displayed as measured without convolution or high pass filtering. The automated procedure for taking a set of spectra takes 10 to 20 h, therefore maps are slightly distorted due to drift. More experimental details can be found in the supporting information (SI) [24].

We reported previously [21] that after the evaporation of 3 parts NC-Ph<sub>6</sub>-CN and 2 parts Co [Fig. 1(a)] large domains of RT stable metal-organic networks self-assemble as exemplified in Fig. 1(b). These supramolecular structures provide regular arrays of uniform hexagonal pores. The spectroscopic dI/dV map obtained at a bias value  $V_B = 101 \text{ mV}$  [Fig. 1(c)] demonstrates a standing electron wave pattern indicating confinement [2,3,8,18,25] of the electrons of the surface state described by the energy dispersion  $E(k) = E_0 + \frac{\hbar^2}{2m^*}k^2$ , where  $E_0 = -65$  meV is the onset energy,  $m^* = 0.42m_e$  the effective mass [25,26],  $\hbar$  the reduced Planck constant, and k the electron wave vector. Thus each pore represents a quantum dot. Earlier work showed that for this network type the size of the unit cell is controlled by the length of the organic linker [20]. Consequenently, the confinement imposed by the networks can be tuned accordingly. As an example we chose NC-Ph<sub>4</sub>-CN [Fig. 1(d)] to construct quantum dots with a reduced confinement area [Fig. 1(e)]. Now, the same standing-wave pattern is obtained at a higher  $V_R$  = 204 mV. In comparison to confinement in previous, sputtered or grown, nanostructures [4,7,8,27] the intensity distributions of the network-confined resonances exhibit more regular shapes directly reflecting the hexagonal symmetry of the networks. Furthermore, the identical spectroscopic features in the different pores [Figs. 1(c) and 1(f)] reveal the crystal-like quality that can be achieved by using

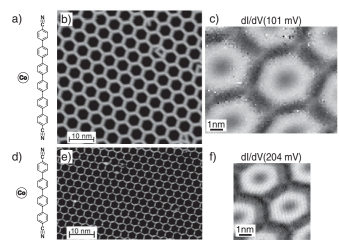


FIG. 1. (a) The self-assembly of NC-Ph<sub>6</sub>-CN and Co produces (b) a crystal-quality metal-organic 2D network with honeycomb-shaped pores (STM image,  $V_B=1~\rm V,~I_T=0.1~\rm nA).$  (c) The experimental dI/dV map (STS,  $V_B=101~\rm mV)$  shows a uniform electron standing-wave pattern in all pores qualifying them as quantum dots. The lateral confinement dimensions are tunable by choice of the length of the molecules. (d) The shorter linking species creates (e) a network of the same symmetry, but with a smaller unit cell (STM image,  $V_B=0.8~\rm V, I_T=0.1~\rm nA)$  resulting (f) in the same electron wave pattern as in (c), but at an increased bias voltage STS,  $v_B=204~\rm mV.$ 

self-assembly protocols for the production of such arrays of quantum dots.

For a more quantitative investigation of the confinement properties we focus on the NC-Ph<sub>6</sub>-CN case in the following before discussing the tunability in more detail later. A set of representative points of the hexagonal unit cell [Fig. 2(a)] was chosen to conduct dI/dV point spectroscopy [Fig. 2(b)]. In contrast to the reference spectrum (gray dashed line), recorded over a pristine surface with the same tip that was used to obtain all the spectra of the set, a strong position dependent variation is present in the spectra distributed over the quantum dot. The c spectrum (black, plus symbols), i.e., the spectrum at the center of the hexagon, shows two clear maxima at  $V_B = -6$  and 205 mV, with the second maximum being enclosed by a shoulder on the low as well as on the high energy side ( $V_B \approx 140$  and 280 mV). The "halfway" spectrum (red, crosses), obtained at half of the distance between the center of the cavity and the center of a molecule, displays its most prominent peak at 68 mV. The "molecule" spectrum (blue squares), on top of the center of a molecule, indicates the smallest local density of states (LDOS) of all positions and appears without sharp features. The Co spectrum (green diamonds), on top of a threefold coordinated Co center, shows more intensity than the molecule spectrum for  $V_B$  < 150 mV, and very similar intensity for higher bias values. Therefore, the difference between the Co and the molecule spectrum is a first indication that, with respect to the electron scattering close to the Fermi level, the two positions behave differently.

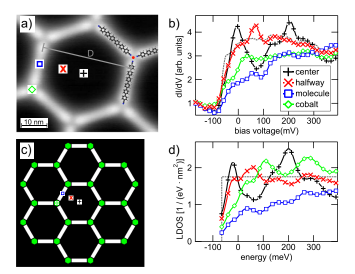


FIG. 2 (color online). (a) Topograph of a single quantum dot. The symbols depict the positions at which the dI/dV spectra of (b) were taken. D is the distance between parallel sides. (b) The experimental dI/dV spectra (symbols) demonstrate a strong position dependent modulation of the nearly constant DOS of the pristine surface (dashed line). (c) The 2D potential model with black, white and green areas corresponding to potential values of  $V_0 = 0$  meV,  $V_{\rm mol} = 500$  meV, and  $V_{\rm Co} = -50$  meV. (d) The spectra calculated with the BEM.

In the following we will analyze the confinement in a first approximation by comparing it to a quantum particle in a 2D hexagonal box [27]. The lowest eigenstates  $\Psi_n$  of this system that exhibit intensity in the center of the hexagon are  $\Psi_1$  and  $\Psi_4$ . They are reflected in the two maxima of the c spectrum in Fig. 2(b), whereas the maximum of the halfway spectrum is connected to  $\Psi_2$ . The numerically obtained eigenvalues  $\lambda_n$  for the hexagonal box [27] scale with the inverse of the area of the hexagon A = $\cos(\frac{\pi}{6})D^2$  and thus the eigenstate energies  $E_n$  can be calculated by  $E_n = E_0 + \frac{\lambda_n}{m^*A}$ . For the  $20 \times 20$  supercell obtained in Ref. [21] the distance between parallel hexagon sides amounts to D = 57.78 Å. With this D the energy values  $E_n$  deviate from the experimental ones ([24], Table I). However, by using an effective diameter  $D_{\rm eff} =$ 1.05D good agreement is obtained. This indicates a substantial penetration of the wave functions into the confining potential, i.e., an appreciable overlap between neighboring quantum dots. The origin of this overlap will be further discussed after the presentation of a more comprehensive theoretical analysis.

Next, we investigate the lateral intensity distributions of the confined resonances. The  $\Psi_1$  state [Fig. 3(a)] exhibits a domelike structure inside the cavity. The molecules appear dark, the Co atoms at the coordination sites show medium brightness. A central depression surrounded by a bright ring leads to a donut shape for  $\Psi_2$  [Fig. 3(b)]. In this case also the corners exhibit a higher electron density than the molecules. At 200 mV [Fig. 3(c)] the map is characterized by a protrusion in the center encompassed with a dark inner and a brighter outer ring. At the energy of the 5th eigenstate [Fig. 3(d)] the central maximum is very shallow and the outer ring is divided into six bright spots near to the centers of the molecules. The comparison with the theoretical state densities of Ref. [4] allows extracting two conclusions. First, the intensity in the outer ring of the  $\Psi_4$  maps is too high compared to the central peak, but can well be explained if a mixing with  $\Psi_3$  is taken into account. Thus the map at 200 mV is not an eigenmode, but a mixture with the 3rd state that should appear at  $V_B = 174 \text{ mV}$  (see

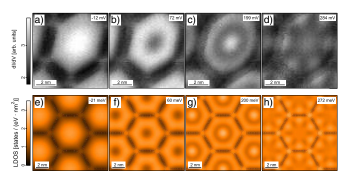


FIG. 3 (color online). (a)–(d) Measured dI/dV maps of a hexagonal quantum dot at indicated bias. (e)–(h) The electron wave patterns obtained with the BEM for the model potential of Fig. 2(c) reproduce well the characteristics of the experimental maps.

SI [24]), but is not resolved because of the width of the resonances. Second, for n = 2 to 5, all eigenstates are expected to display maxima inside the cavity near the corners. Since the experimental maps do not show these maxima, the scattering potential of the Co atoms must be less reflective than that of the molecules.

We have gained further insight into the confinement properties of the network through simulations using a scalar version of the electromagnetic boundary element method (BEM) [18,22,28]. The Schrödinger equation is solved for a 2D potential consisting of regions of constant potential with abrupt boundaries. The solution is expressed by boundary sources, which are propagated in each region via the electron Green's function. The sources are determined from the continuity of both the wave function and its gradient, thus defining a system of integral equations [24].

Using the already mentioned values for  $E_0$  and  $m^*$  for the surface-state electron band, we have defined the effective scattering potential landscape as depicted in Fig. 2(c), where the black regions have zero potential and in which the molecules are described by rectangles (white) with the same length (2.96 nm), widths (0.25 nm, C-C distance perpendicular to the long molecular axis), and effective scattering potential value ( $V_{\rm mol} = 500$  meV) that were successfully employed earlier [18]. The value  $V_{Co} = -50$  meV for the hexagonal Co regions (green) and the energy-independent phenomenological broadening (25 meV) were adjusted to obtain the best agreement with experiment.

Using the BEM, we calculated the energy dependent LDOS at selected (x, y) positions and compared them to the measured dI/dV spectra. Despite the simplicity of the model, the simulated spectra [Fig. 2(d)] agree rather well with the measured dI/dV point spectra [Fig. 2(b)] regarding peak positions, intensities, and width. The only exception is the Co spectrum, which deviates from the experiment for  $E \gtrsim 100$  meV.

Furthermore, conductance maps [LDOS(x, y)] of the resonances  $\Psi_i$  [Figs. 3(e)–3(h)] were calculated at the energies  $E_i$  indicated by the maxima in the dI/dV spectra shown in Fig. 2(d). In this model description, the surfacestate electrons are restricted to a 2D plane and, therefore, there is not any z coordinate representing the height. Thus, we compare with experimental spectroscopic data taken under open-feedback-loop conditions, corresponding to constant height conditions. Note that all simulated maps are displayed using an identical color scale, and that the same is true for the experimental maps. Close examination of the simulated standing-wave patterns reveals a reproduction of the experimental data in great detail within the cavities. For the Co sites, the agreement is very good for energies E < 100 meV, but too much intensity is present at higher energies in the simulated maps, as anticipated from the dI/dV point spectra. The approximation of describing the actual three-dimensional scattering of surface-state electrons by an energy-independent 2D effective scattering

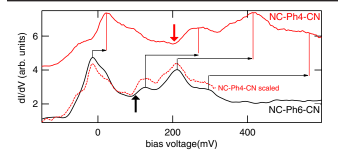


FIG. 4 (color online). The c spectrum for cavities made with NC-Ph<sub>6</sub>-CN linkers (black) and with NC-Ph<sub>4</sub>-CN linkers (red). The four local maxima are present in both spectra and are upshifted for the shorter molecules as clearly demonstrated by the scaled version (red, dashed) of the NC-Ph<sub>4</sub>-CN spectrum, for which the bias values were scaled by the ratio of the confining areas  $R_{\rm Exp}$  (for details see [24]). The bold arrows mark the energies of the maps presented in Figs. 1(c) and 1(f).

potential has some limitations and, therefore, one cannot expect a fully correct description of the real system.

The overall agreement between the measured and the simulated shapes of the resonances substantiates that the difference between the eigenstates of a purely hexagonal quantum box and our system originates from the attractive scattering potential at the Co atoms. In contrast, an isolated Co adatom on the Ag(111) surface is characterized by an effective repulsive scattering potential for electrons at the Fermi level [29]. Thus, our study indicates that the metalligand interactions induce a significant change of the embedded Co centers and their interplay with the surface-state electrons. Even though in an extended system it is not trivial to assign a specific charge to an individual entity, we interpret this result using the following picture. The electron charge transfer from the Co centers to the CN-ligands changes the character of the electron-Co interaction from repulsive (isolated Co adatom) to attractive (Co bond to CNligands) for electron scattering close to the Fermi level. However, the HOMO-LUMO gap of the molecules represents a repulsive potential for these electrons, even after the formation of the metal-organic coordination.

Now we discuss the tunability of the energetic positions of the quantum dot states in more detail. In Fig. 4 we compare the c spectra of the networks constructed from NC-Ph<sub>6</sub>-CN and NC-Ph<sub>4</sub>-CN. The characteristic features (highlighted by vertical lines) shift to higher energies for the shorter molecules. As shown in [24], we obtain a constant scaling ratio  $R_{\rm exp} = 1.74$  for all four features marked in the Figure proving that the energy levels are controlled by the quantum dot dimensions. A version of the NC-Ph<sub>4</sub>-CN spectrum with the bias values scaled by  $R_{\rm exp}$ highlights that the shape of the spectrum remains nearly unchanged when reducing the length of the linkers. The ratio of the quantum dot areas for the two networks is R =1.83, thus compared to confinement in a hexagonal box the expected scaling factor is somewhat larger than  $R_{\rm exp}$ . This difference is consistent with the enlargement of the confining area produced by the attractive Co potential that does not scale with the molecular length.

In conclusion, we have self-assembled RT stable metalorganic networks behaving as arrays of coupled quantum dots. The preservation of the metal-organic binding motif upon a variation of the organic linker allows to control the confinement geometry and accordingly the electronic level alignment. A further functionalization of the molecules can provide an additional parameter to tailor the system. Thus, we introduced a versatile method to tune the interface electronic properties by employing an easy-to-control supra-molecular engineering protocol.

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- \*florian.klappenberger@ph.tum.de
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