Two-dimensional metal-organic coordination networks of Mn-7,7,8,8-tetracyanoquinodimethane assembled on Cu(100): Structural, electronic, and magnetic properties

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A compound two-dimensional (2D) monolayer mixing Mn atoms and 7,7,8,8-tetracyanoquinodimethane (TCNQ) molecules was synthesized by supramolecular assembly on a Cu(100) surface under ultrahigh vacuum conditions. Its structural, electronic, and magnetic properties were analyzed by scanning tunneling microscopy experiment and theory, low-energy electron diffraction, x-ray photoemission spectroscopy, and density-functional theory calculations. The 2D compound has a long-range ordered square planar network structure consisting of fourfold coordinated Mn centers with a Mn:TCNQ ratio of 1:2. The electronic-state analysis revealed a complex charge-transfer scenario indicating strong bonding of TCNQ with both the Mn adsorbates and the Cu surface atoms. The calculations reveal that the Mn centers carry a magnetic moment close to 5\(\mu_B\) with very weak coupling between adjacent Mn centers.

DOI: 10.1103/PhysRevB.80.155458

PACS number(s): 68.43.Hn, 81.16.Fg, 68.37.Ef, 68.43.Bc

I. INTRODUCTION

Organic-based alternatives to conventional magnets, which offer the possibility to form self-organized nanometer-scale structures with novel magnetic properties, have drawn much attention in the past decades.1–5 Famous examples are charge transfer salts of metal-7,7,8,8-tetracyanoquinodimethane (TCNQ), with molecular structure shown in the inset of Fig. 1(b)). TCNQ is chosen to coordinate with transition-metal ions for preparing molecular magnets because local superexchange interactions between the high spin state of metal ions and organic spin carriers are expected.6,7 It has been suggested that the magnetic properties of the metal-TCNQ magnets highly depend on the coordination. Therefore it is important to understand the origin of the magnetism of metal-TCNQ compounds on account of their structural and charge-transfer characters.

Studies of metal coordination at surfaces offer the distinct advantages of high control over surface composition as well as capabilities for structural determination with atomic/molecular resolution, and magnetism measurements. The high degree of composition control in ultrahigh vacuum studies complements well to careful theoretical calculation. A unique way to study metal coordination is through surface-assisted coordination assembly.8–10 On two-dimensional (2D) surfaces, due to surface confinement, the formation of novel 2D coordination systems has been demonstrated.11–14 In this paper we present results of the assembly of two-dimensional Mn(TCNQ)$_2$ coordination networks on a Cu(100) surface. By means of a combined study with scanning tunneling microscopy (STM), low-energy electron diffraction (LEED), x-ray photoelectron spectroscopy (XPS), and density-functional theory (DFT) calculations, we identified that single Mn atoms (ions) form well-defined fourfold square-planar coordination with the cyano groups of TCNQ, resulting in ordered 2D square networks epitaxially grown on the Cu(100) surface. XPS and DFT analyses indicate that a charge transfer to the TCNQ molecules occurs from both the Mn centers and surface Cu atoms. In particular, the magnetic moment of the Mn centers is high (4.9\(\mu_B\)), with weak interactions between Mn centers.

II. EXPERIMENT

The preparation of samples was carried out in two ultrahigh vacuum systems with base pressures of about 3 \(\times 10^{-10}\) mbar. The Cu(100) surface was cleaned by cycles of Ar$^+$-ion sputtering and annealing at 800 K. TCNQ molecules were deposited on a clean Cu(100) surface by organic molecular-beam epitaxy (OMBE) from a Knudsen-cell-type evaporator at a sublimation temperature of 370 K. Mn was subsequently deposited by an electron-beam heating evaporator. The substrate was held at 160 K during and between both deposition steps. After deposition, the sample was annealed for 15 min at a temperature between 380 and 420 K before in situ transfer for STM or XPS characterization, which was done with the sample at room temperature. The home-built Besocke-type STM was operated in the constant current mode with currents of about 0.1–0.5 nA and a sample bias between −0.5 and −1.0 V. X-ray photoemission spectra were measured with an Axis Ultra instrument (Kratos Analytical, Ltd., U.K.), using a monochromatized Al K\(\alpha\) x-ray source of 1486.6 eV and a small spot focusing hemispherical electron-energy analyzer.
The yield of the Mn-TCNQ network phase is quantitative, indicating very efficient diffusion and mixing of each of the two components on the surface. Two differently oriented domains of the network were observed, as marked by the arrows in Fig. 1(a), consistent with mirrored domains and differing by an angle of 16°. These domains have distinct organizational chirality, where the two enantiomeric forms differ in the packing of the TCNQ molecules around the Mn center in a pinwheel formation. Due to the C_s symmetry of the Cu(100) substrate, we also observed the same domains rotated 90°. A high-resolution STM image of the network is shown in Fig. 1(c), which clearly reveals that the network is composed of interconnected TCNQ molecules. The corresponding unit cell is marked by the square in Fig. 1(c), which has a side length of about 1.28 nm. This suggests that the neighboring Mn centers are separated by a distance close to $5a_0$ ($a_0=2.55$ Å, the nearest-neighbor distance of the Cu substrate atoms). The LEED pattern shown in Fig. 1(b) indicates a superposition of two square structures oriented by ±8° with respect to the substrate [010] direction. Both STM and LEED imply that the assembled Mn-TCNQ network periodicity can be expressed as a (4,3/-3,4) superstructure matrix with respect to the substrate Cu(100), equivalent to the Wood notation (5×5)R37°.

A schematic model is overlaid on a STM topograph in Fig. 1(c), where each TCNQ molecule is attached to two Mn atoms and each Mn atom is surrounded by four equivalent TCNQ molecules, giving rise to a Mn(TCNQ)_{12} compound adsorbed on the Cu(100) surface. Based on this model, the optimized structure shown in Figs. 1(d) and 1(e) was obtained by DFT calculations. The hollow site for the Mn atom positions on the Cu(100) is a highly probable geometry, consistent with the experimentally observed structure, and is confirmed as a local energy minimum in the total-energy calculations. The optimum distance between Mn and the four nearest Cu atoms is 2.98 Å, which is slightly longer than the sum of the covalent radii of Cu (1.32 Å) and Mn (1.61 Å for Mn in high spin state). The copper surface shows little relaxation except that the copper atoms which bind to the N atoms of TCNQ move outward by 0.20 Å compared to the average height of the other Cu atoms (the flat part) in that layer. The C atoms of the rings are at an average height of 3.32 Å above the average height of the flat part of the first copper layer, while Mn is 0.89 Å below the average height of the C atoms in the ring. Two of the cyano groups of the TCNQ molecules are strongly bent toward the metal surface with the N atoms binding to the Cu surface atoms with a Cu-N bond of length 2.00 Å. The other two cyano groups bind to two Mn atoms through the N atoms with a Mn-N bond length of 1.61 Å.

We performed XPS characterization to study the charge states of the Mn(TCNQ)_{12} network. XPS signals in the N 1s region are shown in Fig. 2 for (a) TCNQ powder, (b) a TCNQ monolayer, and (c) Mn(155458-2

FIG. 1. (Color online) (a) STM topograph of the Mn-TCNQ assembly on a Cu (100) surface. (b) High-resolution STM topograph of the Mn(155458-2 network. (c) LEED pattern of the Mn(155458-2 network. Inset: chemical structure of the molecule TCNQ. [(d) and (e)] Top and side views of the DFT-optimized structure of the Mn(155458-2 network on the Cu(155458-2 surface color code: Cu red, C gray, N blue, H white, and Mn purple. STM images recorded at (a) 0.4 nA and −0.96 V and (b) 0.3 nA and −0.72 V. LEED image (c) recorded at 16 eV.

III. THEORY

In our DFT calculations, the structures were optimized with the VASP package, which uses a periodic approach with plane waves as basis set. The projected augmented wave method is adopted. The exchange-correlation functional was treated within the generalized gradient approximation of Perdew and Wang with the Vosko-Wilk-Nusair interpolation of the correlation energy. The kinetic-energy cutoff was set to 400 eV. Five-layer slabs, with 10 Å of vacuum between slabs, were used to simulate the surface. The top three copper layers and all the adsorbate atoms were allowed to relax until the largest force component was smaller than 0.04 eV/Å, while the bottom two copper layers were fixed to the experimental bulk Cu geometry. Intersections in the first Brillouin zone were performed using a Monkhorst-Pack grid of (2×2×1) k points. The details of the initial guessed geometries and the supercell employed will be given in the next section.

IV. ANALYSIS

A typical experimental STM topograph of the Mn-TCNQ assembly on Cu(100) is shown in Fig. 1(a). The lower part of the figure shows Mn-free close-packed TCNQ molecules, characterized and modeled in detail elsewhere. Above this region is a periodic network structure of TCNQ with Mn and voids (interspersed with larger vacancies). The molecular phase converts into the network phase as the Mn is vapor deposited to the surface, i.e., with increasing Mn coverage.
soon as they adsorb on the substrate. The N 1s signal of (a) TCNQ powder, (b) TCNQ monolayer grown on Cu(100), and (c) Mn(TCNQ)₂ networks grown on Cu(100).

Through Bader charge analysis of the DFT optimized geometry, we can gain insight into the charge-transfer mechanism by considering the partial systems listed in Table I; we stress that the trends in such charges are more meaningful than the precise numerical values. The atomic coordinates of the three partial systems are the same as in the full 2TCNQ+Mn/Cu(100) system. In the full system, the TCNQ molecule, serving as electron acceptor, carries a negative charge of −1.38|e|, the Cu and the Mn both contribute to the donation. The charge of Mn is +1.33|e|, which approaches the Mn²⁺ state, whereas the Cu(100) surface is charged to +1.43|e|. In the 2TCNQ+Mn system, we found a transfer of −0.73|e| to each TCNQ from Mn and in the Mn-free 2TCNQ/Cu(100) system, a transfer of −1.12|e| occurs from the copper surface. The charge on the TCNQ in the full 2TCNQ+Mn/Cu(100) system, −1.38|e|, exceeds these values: the changes on the N atoms increase by −0.24|e| for N₁ and −0.30|e| for N₂ with respect to the pure (TCNQ)₂ network, where N₁ and N₂ bond to Cu and Mn, respectively. This charge increase is primarily due to donation from Cu to N₁ and from Mn to N₂. For the Mn-free 2TCNQ/Cu(100) system, the redistribution is principally localized in the N-Cu bonds, where each N₁ atom gains −0.24|e|. For the Mn(TCNQ)₂ network in the absence of the Cu substrate, the N₂ atoms gain −0.30|e| relative to N in the pure (TCNQ)₂ network. This analysis reveals that the charge of the N₁ atoms in the Cu-N bonds changes little compared to that of the N₂ atoms in the Mn-N bonds, which is consistent with the XPS data. This is also consistent with a strong electron donation from both Mn and Cu to the TCNQ, especially to the N atoms. This significant charge transfer is a critical aspect of the system, both for structural arrangement and for determining the electronic configuration, as discussed below.

The geometry of the TCNQ also reveals the effect of the charge transfer. While the successive C-C bonds in free TCNQ have alternating long and short distances (cf. the C₀−C₁, C₁−C₂, C₂−C₃, and C₃−C₄ distances listed in Table II), the structure after adsorption (dianion TCNQ⁻² species, cf. Table II), shows a shortened C₀−C₁ distance, an elongated C₁−C₂ distance and nearly equivalent C₂−C₃ and C₃−C₄ distances within the ring.

The optimized Mn(TCNQ)₂/Cu(100) network structure was used to simulate the STM imaging, as shown in Fig. 3(a). The Mn centers dominate the image and are thus too bright compared to the experimental data of Fig. 1(c). After excluding tip effects, we ascribed this discrepancy to the fact that the electronic structure of the Mn(TCNQ)₂ network is not described accurately by the normal DFT approach. In a system such as Mn(TCNQ)₂ the electronic correlation effect due to tightly bound and localized d states on the transition-metal atoms should be taken into account. An improvement that addresses this issue is the LDA+U method. In this approach, a Hubbard-type term is added to the DFT Hamiltonian to account for the strong on-site intra-atomic Coulomb repulsion U. The parameter U describes the energy rise due to placing an extra electron into the d level on a particular site, and an additional parameter J represents the screened exchange energy. While U depends on the spatial extension of the wave functions and on screening, J is an approximation to the Stoner exchange parameter and is almost constant with a value near 1 eV. We applied the DFT+U approach based on the work of Dudarev et al., which uses the single effective parameter U−J (or U_eff=U−J). As a rough approximation, we set the U_eff value for Mn to 4.2 eV. Figure 3(b) shows the STM image produced by the parameter fitting to the projected density of states (PDOS) with the Hubbard U correction. We see that the signal of the Mn centers is significantly reduced relative to that of the TCNQ molecules, which agrees much better with the experimental images than without a Hubbard U correction.

The LDA+U treatment pushes the unoccupied states to higher energy, increasing the energy gap. As shown in Figs. 3(c) and 3(d), with the correction U_eff = 4.2 eV, the gap between the occupied and unoccupied majority-spin states is enlarged from 1.9 to 4.1 eV. Without the Hubbard U correction, there is a large contribution from Mn 3d minority-spin states in the energy range of ±1 eV around the Fermi level.
which results in the Mn atom contributing significantly to the tunneling current and appearing very bright in the simulated STM image. After applying the Hubbard $U$ correction with $U_{\text{eff}}=4.2$ eV, the Mn 3d orbital DOS above/below the Fermi level is shifted up/down in energy in agreement with the $U$ exclusion effect, forming a large energy gap. As shown in Fig. 3(d), the TCNQ levels are not affected by this shift, so that the above-mentioned charge transfers are only negligibly modified. As a result, the PDOS of Mn 3d is smaller than the PDOS of TCNQ at the Fermi level, making the Mn atom in the simulated STM less bright than without Hubbard correction.

We next address the magnetic moment in this structure, adopting the convention, frequently used in the literature, that the above-mentioned charge transfers are only negligibly modified. As a result, the PDOS of Mn 3d is smaller than the PDOS of TCNQ at the Fermi level, making the Mn atom in the simulated STM less bright than without Hubbard correction.

![Fig. 3](image)

**Fig. 3.** (Color online) (a) Simulated constant-current STM image at 0.5 nA and 0.5 V (giving an average tip-to-molecule height of 5 Å), using DFT results for the Mn(TCNQ)$_2$ network structure on Cu(100). (b) Same as (a) but with a Hubbard $U$ correction, $U_{\text{eff}}=4.2$ eV. [(c) and (d)] Spin PDOS for Mn and TCNQ adsorbed on a Cu(100) surface, calculated with VASP without Hubbard $U$ (c), and using $U_{\text{eff}}=4.2$ eV (d). The PDOS of Mn is indicated in blue. The PDOS of the TCNQ molecules is indicated in red solid lines, with positive (negative) values for the majority-spin (minority-spin) PDOS. Square unit cells indicated in (a) and (b) with side length 1.28 nm.

![Fig. 4](image)

**Fig. 4.** (Color online) (a) The PDOS of the top copper layer, of the TCNQ molecule, and of the Mn $d_x$ and $d_y$ orbitals. (b) The PDOS of the Mn $d_z$ orbitals and of the four N atoms bonded to Cu. Note that the PDOS of the $d_x$ and $d_y$ orbitals are equal.

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<th>$C_0N_Cu$</th>
<th>$C_0-C_1$</th>
<th>$C_1-C_2$</th>
<th>$C_2-C_3$</th>
<th>$C_3-C_3$</th>
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<td>1.39(1.37)</td>
<td>1.45(1.45)</td>
<td>1.355(1.35)</td>
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<td>1.475</td>
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<td>1.47</td>
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<td>1.37</td>
<td>1.50</td>
<td>1.40</td>
<td>1.39</td>
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TABLE II. Bond distances (Å) in TCNQ, as neutral monomer, negatively charged dianion dimer, Mn-[TCNQ]$_2$/Cu(100) and TCNQH$_4$ (the theoretical basis set used is specified between parentheses).
the $d_{xz}$ and $d_{yz}$ orbitals are equivalent). The Fermi level is determined by the copper orbitals, which do not contribute to the spin polarization. In Fig. 4(a), three peaks of the TCNQ levels—a sharp peak at $E_{\text{Fermi}}-3.5$ eV (2e), a family of peaks at $E_{\text{Fermi}}-2.5$ eV (4e), and a sharp peak at $E_{\text{Fermi}}-0.5$ eV (2e)—correspond to four Cu-N bonds and do not contribute to the spin. Figure 4(b) displays the PDOS on the four N atoms adjacent to the cyano groups close to Cu which are the dominant contribution. We can therefore interpret the magnetization as resulting from a Mn$^{2+}$ ion interacting with a TCNQ dimer dianion, [TCNQ$^{-2}$]$_2$. Our calculations confirm that the spin is strongly localized on the Mn.

Finally, we searched for a periodic spin arrangement and found that, using the Mn+2TCNQ model with three Cu layers, the ferromagnetic arrangement and the antiferromagnetic arrangement have essentially the same energy (differing by only 5 meV). Therefore, the spin coupling between adjacent Mn ions is small and each Mn ion may be regarded as an independent magnet.

V. CONCLUSIONS

We have performed a combined experimental and theoretical study of a self-assembled Mn-TCNQ network on Cu(100) under ultrahigh vacuum conditions, using STM, LEED, XPS, and DFT calculations. The STM and LEED experiments reveal a highly ordered (5×5)R37° periodic structure. Each unit cell contains two intact flat-lying TCNQ molecules and one Mn atom, which connects four TCNQ molecules together. Theory indicates significant charge transfer in the strong Mn-N bonding within the 2D network, as well as in Cu-N bonds that pin the network to the Cu(100) surface. The calculations reveal that the Mn centers carry a magnetic moment of 4.9μ$_B$, with weak coupling between adjacent Mn magnetic moments. Together, these results characterize a two-dimensional metal-organic system formed by a highly efficient self-assembly process which consists of well-ordered independent magnetic metal centers.

ACKNOWLEDGMENTS

This work was supported in part by Hong Kong RGC under Grant No. CityU 102408, and by the CityU Centre for Applied Computing and Interactive Media. All of the experimental work was performed at MPI Stuttgart and was supported by the European Science Foundation (ESF) EUROCORES-SONS2 program FunSMARTs II. J.I.C. acknowledges financial support from the Spanish MICINN under Contract No. MAT2007-66719-C03-02. C.M. is grateful to the French Consulate General in Hong Kong and Macau for financial support.
We simulated STM images by using the GREEN code (Ref. 34), which is based on a Green’s function representation and employs a linear combination of atomic orbitals. The extended Hückel theory (EHT) was used to compose the semiempirical Hamiltonian. First we fit the EHT parameters to reproduce the DFT DOS of the sample and tip. Then an infinite system composed of two semi-infinite blocks for the tip and the sample was constructed, using the parameterized EHT to simulate the STM image. The W tip has a pyramidal shape with the W(110) plane parallel to the surface, although the exact tip shape and composition used in the experiment are not known.

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