Ferromagnetic Coupling of Mononuclear Fe Centers in a Self-Assembled Metal-Organic Network on Au(111)


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The magnetic state and magnetic coupling of individual atoms in nanoscale structures relies on a delicate balance between different interactions with the atomic-scale surroundings. Using scanning tunneling microscopy, we resolve the self-assembled formation of highly ordered bilayer structures of Fe atoms and organic linker molecules (T4PT) when deposited on a Au(111) surface. The Fe atoms are encaged in a three-dimensional coordination motif by three T4PT molecules in the surface plane and an additional T4PT unit on top. Within this crystal field, the Fe atoms retain a magnetic ground state with easy-axis anisotropy, as evidenced by x-ray absorption spectroscopy and x-ray magnetic circular dichroism. The magnetization curves reveal the existence of ferromagnetic coupling between the Fe centers.

As the size of a magnetic material decreases, the stability of the collective magnetic ground state becomes weaker, leading to paramagnetic behavior at atomic-scale dimensions [1]. The identification and study of mechanisms for stabilizing magnetically oriented ground states aims to provide routes for the design and construction of nanomaterials (clusters and thin films) that retain some magnetic order at temperatures as high as possible. At the limit of individual atoms deposited on a metal surface, magnetic anisotropy and interatomic coupling are the key ingredients to tune the magnetism of a nanostructure [2]. The anisotropy of the atomic environment may lower the magnetic moment but, in turn, defines easy directions for magnetization. Exchange interactions coupling the magnetic moment of neighbor atoms drive the formation of magnetically ordered states, but this effect quickly vanishes with increasing interatomic distance. Only the indirect exchange coupling through valence electrons of the metal support (the Ruderman-Kittel-Kasuya-Yosida (RKKY) interaction [3–7]) may be operative. It has been shown that RKKY interactions lead to collective ferromagnetic or antiferromagnetic ground states, but the exchange energy is small ($J \sim 50 \mu$eV [1,7,8].

An interesting approach to enhance the magnetic anisotropy of atoms in a thin film is to augment the ligand field splitting of the $d$ levels by coordination to specifically selected organic ligands [2]. The formation of a coordination bond to molecular end groups induces a charge redistribution at the metal sites and casts the ligand field by adopting high-symmetry metal-organic structures and periodic networks [9–11]. In bulk metal-organic materials, the organic linkers connecting magnetic metal centers also mediate the magnetic coupling between them by means of electronic states extended along the organic backbone [12,13]. The presence of such a superexchange mechanism for magnetic coupling in metal-organic thin films could optimize the stabilization of collective magnetic ground states, but its detection still remains a challenge.

Here, we show the existence of ferromagnetic coupling between Fe ions within a self-assembled metal-organic film grown on a Au(111) surface. Single Fe atoms lie in three-dimensional coordination sites and are connected to neighboring Fe atoms by 2,4,6-tris(4-pyridyl)-1,3,5-triazine (T4PT) molecules [see Fig. 1(b)]. Analysis of x-ray absorption spectroscopy and x-ray magnetic circular dichroism, the magnetization curves reveal the existence of ferromagnetic coupling between the Fe centers.
absorption spectroscopy (XAS) and x-ray magnetic circular dichroism (XMCD) measurements reveals a high spin $S = 2$ ground state with an out-of-plane magnetic anisotropy. Magnetization curves of the ultrathin layer exhibit deviations from the case of noninteracting paramagnetic moments, which are interpreted as a finite ferromagnetic interaction between the mononuclear centers on the basis of a mean-field model.

Fe-T4PT networks were grown on an atomically clean Au(111) single crystal. The gold surface was first precov- ered at room temperature by a submonolayer amount of T4PT molecules, sublimated from a Knudsen cell, and by Fe atoms. Subsequently, the substrate was annealed to 400 K to activate the formation of coordination bonds. The structure of the resulting surface was characterized in situ, in our low temperature (4.8 K) scanning tunneling microscope (STM). XAS and XMCD measurements in total electron yield were performed at the beam line ID08 at the European Synchrotron Radiation Facility (ESRF). The beam size on the sample was about 1.4 mm (FWHM) in the horizontal and 50 $\mu$m in the vertical direction. The sample was kept at a temperature of 8 K in a variable external magnetic field up to 5 T and exposed to x-rays under different incidence angles. Here, STM measurements (Fig. 1) were carried out in situ at several macroscopically separated areas, indicating that the sample was covered by a homogeneous metal-organic network and that there are no uncoordinated Fe clusters.

Large-scale STM images show the tendency of Fe atoms and T4PT molecules to coordinate in networks with a high degree of order [Fig. 1(a)]. The presence of the Au(111) herringbone reconstruction underneath the metal-organic layer indicates a weak interaction with the substrate. STM images recorded at low bias voltage ($V_S < 0.5$ V) resolve the skeleton of the T4PT molecules [Fig. 1(c)]. Each pyridyl group participates in a threefold node, a configuration that can only be stable if the natural repulsion between electrophilic end groups is overcome by insertion of a bonding center prone to coordinate with the $N$ lone-pair electrons [9,11]. Hence, we anticipate the inclusion of Fe atoms at these sites [14].

The structure of the Fe-T4PT network shows a peculiar dependence on the bias voltage applied during its inspection [Figs. 2(a) and 2(b)]. Starting from the low bias voltage structure in Fig. 2(a), an increase of the sample bias voltage above 0.6 V results in the observation of a new set of T4PT molecules centered on the Fe sites and rotated by $\varphi = 29^\circ$ [Fig. 2(b)]. These new features can not be associated to tunneling through different orbitals of the same molecules due to their different orientation. Furthermore, we observe domains with mirror symmetry around dislocation lines at high bias voltage that are absent when inspected at low bias voltage (see Supplemental Material S1 [15]). On the basis of these facts, we conclude that the film consists of a bilayer of T4PT molecules: Fe atoms coordinate to the pyridyl nitrogen atoms [Figs. 2(c) and 2(d)]. The presence of the top T4PT molecule centered directly on top [second layer in Fig. 2(d)] results in the Fe-T4PT network. Fe atoms are threefold coordinated by three T4PT molecules to coordinate in networks with a high degree of order [Fig. 1(a)]. The presence of the Au(111) herringbone reconstruction underneath the metal-organic layer indicates a weak interaction with the substrate. STM images recorded at low bias voltage ($V_S < 0.5$ V) resolve the skeleton of the T4PT molecules [Fig. 1(c)]. Each pyridyl group participates in a threefold node, a configuration that can only be stable if the natural repulsion between electrophilic end groups is overcome by insertion of a bonding center prone to coordinate with the $N$ lone-pair electrons [9,11]. Hence, we anticipate the inclusion of Fe atoms at these sites [14].

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A +2 oxidation state is typical in bulk mononuclear iron complexes [22], thus supporting the three-dimensional coordination motif described in Fig. 2(d). Multiplet calculations of the XAS and XMCD difference spectra indicate an $S = 2$ high-spin state (see Supplemental Material S2 [15]).

The presence of a magnetic moment is further corroborated by XMCD spectra shown in Fig. 3(b). The XMCD difference signal of the $L_3$ peak exhibits a characteristic dip-peak structure at the low-energy side. While the line shapes are similar for different angles of incidence, there is a large change in the XMCD peak height: the signal at normal incidence is more than a factor of 2 higher than that at grazing incidence ($20^\circ$). The higher XMCD signal in normal incidence points towards an easy-axis magnetic anisotropy along the direction normal to the surface, which is a result of the spin-orbit-induced mixing of nearly degenerate Fe 3$d$ levels in the surrounding ligand field.

Figure 4 shows the evolution of the Fe $L_3$ XMCD signal with applied external magnetic field for both perpendicular and grazing incidence. The curvature of these plots allows us to quantify the anisotropy of the system. We start first by simulating the magnetization curves assuming noninteracting paramagnetic moments under the presence of magnetic anisotropy by means of a spin Hamiltonian approach:

$$\mathcal{H} = D S_z^2 - \mu_B g \mathbf{B} \cdot \mathbf{S}.$$  \hspace{1cm} (1)

The first term describes the anisotropy energy through the zero-field-splitting parameter $D$ and $S_z$, the spin component normal to the molecular plane (a negative $D$ stands for an easy axis along $z$). The second term accounts for the Zeeman energy ($S$ is the spin vector, $\mathbf{B}$ is the vector of the external field, $\mu_B$ is Bohr’s magneton, and we consider a $g$ factor of 2). Solving Eq. (1) for the two directions of measurement and calculating the magnetization from the thermal population of the resulting levels allows us to simulate magnetization plots as in Fig. 4 for several conditions. The curvature of the magnetization plots is an expression of the anisotropy of the system and can be...
tuned by adjusting the parameter $D$. The larger curvature in normal incidence corroborates that this direction is the easy axis for magnetization ($D < 0$). However, the experimental curvature cannot be properly reproduced, no matter what value of $D$ is used. The dashed line in the inset of Fig. 4 shows the mean squared deviation between experiment and simulations evaluating Eq. (1) as a function of $D$. We obtain large standard deviations for all values of $D$, even when the curvature is simulated using unrealistically high $D$ values. We thus conclude that the experimental data are not compatible with noninteracting anisotropic paramagnetic moments but bear instead evidence for a magnetic coupling between neighboring Fe moments [29].

To include magnetic interactions between nearest neighbors, we resort to a mean-field description to simulate the magnetization curves. In this model, the magnetic interaction of a spin with all other spins in the neighborhood is replaced by an effective mean magnetic field added to the external applied field and proportional to the magnetization of the ensemble with proportionality factor $T_C = \sum J_i /(3k_B)$. Here, $J_i$ is a Heisenberg-type interaction of the spin with a spin at site $i$ and $k_B$ is the Boltzmann constant. The sum goes over all sites with nonvanishing $J_i$. $T_C$ represents the Curie temperature in the limit of vanishing anisotropy.

The solid lines in Fig. 4 are the result of such a simulation with $D = -0.35$ meV and $T_C = 1.8$ K. These values correspond to a global minimum (see solid line in the inset of Fig. 4 and Supplemental Material S4 [15]), reproducing correctly the experimental magnetization curves. While the model may not capture the details of the coupling mechanism, it clearly evidences the necessity to include ferromagnetic coupling of the Fe atoms to explain the experimental curves. If we assume a coupling to only the six nearest neighbor Fe atoms, the coupling constant $J_i = k_B T_C / 2$ has to be on the order of ~80 $\mu$eV to reproduce the experimentally observed magnetization curves [31]. Since the measurements are carried out well above $T_C$, the effect of correlated fluctuations, which would lead to an underestimation of the derived coupling strength in the mean-field description, is small.

Due to the relatively large distance between the Fe centers ($r = 1.3$ nm), a direct exchange coupling is very unlikely to be the origin of the observed ferromagnetic coupling. An indirect coupling mechanism via the polarization of the surface and bulk conduction electrons has been shown to exist on metal surfaces [6,7,32,33]. The RKKY interaction exhibits an oscillating ferromagnetic-antiferromagnetic coupling with the distance $r$ of the magnetic moments. It can be described as $J_{\text{RKKY}} \propto [\cos(2k_F r)]/(2k_F r)^d$ [7,34], where $k_F$ is the Fermi wave vector and $d$ defines the dimensionality of the conduction electron system. The Fermi wave vector of the Rashba-split surface state amounts to $k_{F,1} \approx 1.7$ nm [35]. Hence, a pairwise interaction of the Fe atoms at a distance of 1.3 nm via RKKY interactions would result in a weakly coupled antiferromagnetic ground state. Furthermore, we expect that the sixfold ligated Fe atoms are lifted from the surface [19,20] and thus only interact weakly with the substrate electrons. Despite small effects that may change the scattering phase shift of the conduction electrons, we may tentatively exclude a significant contribution of an RKKY-mediated coupling.

The molecular ligands coordinated to two neighbor Fe atoms can also play an important role in mediating magnetic interactions through a superexchange mechanism. Ferromagnetic and antiferromagnetic exchange interactions in metal-organic frameworks are well-known in bulk materials [12] for nitrogen-based organic linkers and transition-metal atoms. The superexchange mechanism is enhanced when conjugated molecular states bridge the magnetic centers [36], as is the case for Fe-T4PT networks. The spin polarization qualitatively follows a simple alternation rule: an even number of linker atoms in the pathway leads to antiferromagnetic coupling, whereas an odd number results in ferromagnetic coupling. In the case of the Fe-T4PT network, the alternation rule results in a ferromagnetic coupling between neighboring Fe centers. This is consistent with the positive (ferromagnetic) Heisenberg-type interaction $J_i$ deduced from the magnetization curves.

Our results clearly sustain the presence of ferromagnetic coupling of Fe atoms connected by organic molecules in a highly ordered metal-organic network. The peculiarity of the observed network is that it consists of a bilayer structure, in which the Fe is coordinated by three pyridyl groups and one triazine unit. The central triazine ligand may further be responsible for a lifting of the Fe atoms from the metallic surface due to the trans effect [19,20]. This configuration eases the formation of a +2 oxidation state with a net spin. The resulting magnetic moments of the Fe ions are coupled ferromagnetically with each other, probably via a superexchange mechanism mediated by the organic linkers. These results further demonstrate that magnetism in two-dimensional metal-organic networks on a metal surface is feasible without the use of ferromagnetic substrates and suggests a route to engineer its magnetic character by enclosing the magnetic centers in self-assembled three-dimensional coordination cavities. To increase the stability of such magnetic networks for potential applications, a larger anisotropy and significantly stronger magnetic coupling strengths are required. Chemical design strategies toward this goal may comprise the use of radical ligands, known from bulk metal-organic ferromagnetic materials [37,38].

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Clusters of a few atoms size already show a significant broadening in XAS due to hybridization with the neighbor atoms (see e.g., Refs. [2,24,25]).

We rule out that the magnetization is due to Fe clusters, as we do not observe their presence in any STM images taken at various sample locations of the very same preparation of the Fe-T4PT network. Uncoordinated Fe clusters can be further excluded by the line shape [23] of the XAS and XMCD signals and by the magnetization behavior. The magnetization of potential residual Fe clusters would have an in-plane anisotropy [30], contrary to our observed easy-axis anisotropy.

Growth of T4PT films without deposition of Fe atoms resulted in a completely different layer structure, where direct bonds between N atoms of pyridyl moieties are fully absent.