Field-regulated switching of the magnetization of Co-porphyrin on graphene

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Different magnetic coupling mechanisms have been identified for a few monolayers of Co-porphyrin molecules deposited on a graphene-covered Ni(111) single crystal. A relatively strong antiferromagnetic coupling of the first molecular layer via graphene to the Ni crystal in comparison to a weaker intermolecular coupling gives rise to a complex field-dependent response of this hybrid system. By continuously increasing the magnetic field strength, the net magnetization of the molecular system switches from antiparallel to parallel to the field direction at 2.5 T. Utilizing x-ray absorption spectroscopy and x-ray magnetic circular dichroism, the element-specific magnetization and field dependence was probed. The nature of the magnetic couplings is identified by means of density functional theory and orbital-dependent susceptibilities.

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The development of new materials for spintronic applications as an alternative to conventional devices is one of the major challenges of present research. Using the spin of organic materials may be a solution for the limitation of downscaling of current devices [1–3]. Organic molecules, such as metal porphyrins, provide convenient possibilities for a realization of such applications. In recent years, different groups have demonstrated that the magnetization of Mn- and Fe-porphyrins can be controlled by ferromagnetic coupling to ferromagnetic substrates [4–6]. Depending on the magnetization of the film, one can tune the magnetic moments of the molecules to lie either in plane or perpendicular to the surface. With an interlayer of oxygen, the coupling could also be changed, yielding an antiparallel alignment with respect to the magnetic moment of the substrate [7]. Compared to an interlayer of oxygen, graphene has strong advantages when it comes to tuning the exchange interaction between substrate and molecular magnet. The coupling energy via oxygen to the first molecular layer is very high [7,8] compared to using graphene, as presented in this paper. A too strong coupling between the magnetic moments of the molecule and the substrate prevents the possibility to tune the composite system’s net magnetization with, e.g., an external magnetic field. In the present work, we are indeed able to switch the net moment of the Co-octaethylporphyrin (OEP) molecules adsorbed on graphene reversibly between parallel and antiparallel orientations with a magnetic field. The antiferromagnetic coupling on a graphene-covered Ni(111) was calculated for other molecules [9]. Hermanns et al. recently detected an antiferromagnetic coupling of a CoOEP submonolayer to a graphene-covered Ni film [10,11], which followed a study of CoOEP/Ni(100) [12]. Since its discovery, graphene [13] has become one of the most important topics in various fields of material science. Its unique electronic properties make it invaluable for different applications [14]. For instance, by density functional theory (DFT) calculations, graphene was shown to be a reversible spin manipulator for Fe-porphyrin molecules [15].

In this paper, we demonstrate the field dependence of the net magnetization of three layers of CoOEP molecules and a graphene-covered Ni(111) single crystal. Via element-specific x-ray magnetic circular dichroism (XMCD), we can individually determine the magnetization of the Ni crystal and the molecules. The net magnetization of the Co-porphyrin molecules is the sum of two different contributions, i.e., a constant antiparallel magnetization from the first layer and a field-dependent contribution from the second and third ones. This behavior of the system enables us to switch the magnetization by regulating an external magnetic field. We are able to confirm the antiferromagnetic coupling through the graphene layer with DFT calculations and compute the coupling energy as well as spatial- and orbital-dependent susceptibilities. The calculated field-dependent contributions to the magnetism from the two superposing signals fit the experimental data very well.

The samples were prepared in situ at a base pressure of 10−10 mbar. We produced a single-layer graphene on top of the clean Ni(111) single crystal in situ by heating the crystal in a propylene atmosphere, following Ref. [16]. Graphene is known to grow epitaxially on the Ni(111) surface [17]. Subsequently, the CoOEP molecules were thermally evaporated onto the graphene-covered Ni(111) crystal, while the substrate was held at room temperature. The thickness of three layers with an accuracy of ±15% was controlled with a microbalance on top of the evaporator. All molecules are

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FIG. 1. (Color online) Top: Schematic images of the sample: 3 ML CoOEP/graphene/Ni(111). The white and black arrows indicate the directions of the magnetic moments of Ni (blue) and Co (red) in a low magnetic field of 500 mT (left) and in a high magnetic field of 5 T (right). Bottom: XMCD at the Co L2,3 edges of the CoOEP molecules in the low magnetic field (left) and in the high magnetic field (right). Insets show the XMCD at the L2,3 edges of the saturated Ni crystal at 500 mT and at 5 T. All spectra are recorded at $T = 2.8$ K and $\theta = 70^{\circ}$.

oriented and lie parallel to the surface, as shown in Figs. 2 and 3 in the Supplemental Material [18]. The x-ray absorption spectroscopy (XAS) measurements in total electron yield were performed at the undulator beam line DEIMOS at SOLEIL. In the cryomagnet, we could reach a magnetic field up to 7 T and a temperature of 2 K at the sample holder. This results in a temperature of approximately 2.8 K on the sample surface. With the element-specific XMCD technique at the Co and Ni L2,3 edges, we analyzed the magnetic field dependence of the net magnetization of the Ni(111) crystal and the CoOEP molecules.

We have performed first-principles density functional calculations with a full potential plane-wave-based code, VASP [19]. The unit cell used consisted of a slab of three Ni layers in the (111) direction having 108 atoms along with a graphene monolayer with 72 atoms ($6 \times 6$ supercell) on top of the slab. A CoP molecule with 37 atoms was kept on graphene, yielding a total of 217 atoms in the simulation cell. It should be noted that the outermost chains of the CoOEP molecule have been replaced by H atoms in our simulations and, hence, we will denote the molecule as CoP. The size of the supercell was chosen in such a way that the magnetic centers of CoP are at least 15 Å apart from each other in the lateral direction. A $3 \times 3 \times 1$ $k$-point mesh was chosen for geometry optimizations with a force tolerance of 0.01 eV/Å. All of the atomic positions were relaxed except the ones in the lowest Ni layer, which is fixed to have the in-plane lattice constant as that of bulk Ni. A $4 \times 4 \times 2$ $k$-point mesh was considered for calculating total energies and other properties. We have employed a DFT+$U$ approach within the Hubbard model [20] to incorporate strong Coulomb interaction with the Coulomb parameter $U$ and exchange parameter $J$ fixed as 4 and 1 eV, respectively, for Co $d$ orbitals. These parameters have been shown [21] to reproduce certain experimental results for 3$d$ transition-metal-centered porphyrin molecules. All of the calculations were performed including an empirical form of dispersion correction given by Grimme [22].

Figure 1 shows the XMCD spectra at the L2,3 edges for the CoOEP molecules as well as for the Ni single crystal, along with the schematic illustrations of the corresponding samples. The black arrows illustrate the magnetic moments of each Co ion, while the white arrow represents the macroscopic magnetization of the Ni(111) crystal. The spectra were recorded at a temperature of 2.8 K and at grazing x-ray incidence ($\theta = 70^{\circ}$ with respect to the surface normal), with the external magnetic field parallel to the beam. The XMCD signal at the Ni L2,3 is plotted as the inset in both Co spectra, showing the same XMCD signal for both magnetic fields, i.e., 500 mT and 5 T, because the magnetization of the crystal is saturated for $B > 200$ mT. In the spectrum at the high magnetic field of $B = 5$ T, the XMCD intensity at the Co L3 edge is negative, just like the L3 edge of the Ni XMCD. However, in a lower magnetic field of 500 mT, the XMCD signal at the Co L3 edge changes its sign, whereas the shape of the XMCD signal is unchanged. Further details about the field-dependent XMCD are presented in Fig. 4 in the Supplemental Material [18]. At a temperature of 2.8 K, the spin fluctuation of the weakly coupled, paramagnetic Co-porphyrin molecules is too strong to determine a net alignment of the Co moments in a small magnetic field of 500 mT. Hence, the XMCD signal for low magnetic fields is dominated by the first CoOEP layer which is coupled antiferromagnetically to the ferromagnetic substrate. All of the magnetic moments of the Co ions in the first layer are oriented antiparallel to the magnetization of the Ni film and therefore antiparallel to the applied field. The second and third layers are only weakly coupled (see
details below) and are only slightly affected by the small field, as is expected for a paramagnetic system. These Co magnetic moments point in various directions and add up to a negligibly small magnetization in the field direction. In Fig. 2, this is demonstrated in a clear way where the magnetization curves at the Co L3 and Ni L3 edges are presented. In this is demonstrated in a clear way where the magnetization curves at the Co L3 and Ni L3 edges are presented. In this is demonstrated in a clear way where the magnetization curves at the Co L3 and Ni L3 edges are presented. In both cases, the measurements were performed at constant photon energies of 775.4 eV for Co and 850.9 eV for Ni, at which the XMCD intensity is maximum, and normalized to pre-edge energies of 772.0 and 845.0 eV, respectively, while the magnetic field was varied. The magnetization of the Ni single crystal increases strongly in a small magnetic field and saturates at about 200 mT. Simultaneously, at this field strength, the Co magnetization rises to its maximum value, antiparallel to the field, demonstrating the antiferromagnetic coupling. In higher fields, the Ni magnetization remains constant, while the absolute value of the Co XMCD signal decreases and reaches zero at a field of nearly 2.5 T. At this point, the magnetization of the molecule in the second and third layers annihilates the magnetization of the first one. In higher fields, the net magnetization switches its sign because the amount of parallel aligned magnetic moments is higher than the antiparallel magnetic moments of the first layer. Close to zero magnetic field, the magnetization of the Ni crystal vanishes due to spontaneous formation of magnetic domains and so does the magnetization of the Co ions. The hysteresis curve for Co, as shown in Fig. 2, has been fitted with the superposition of two contributions, 

\[
M(B) = M_1 \tanh\left(\frac{\mu_B (B - B_{c1})}{k_B T}\right) + M_{2,3} \tanh\left(\frac{\mu_B B}{k_B T}\right),
\]

with \(S = \frac{1}{2}\) and \(B_{c1} = 72\) T, i.e., the exchange field between the Co moment of the first molecular layer and the Ni surface, obtained from our DFT+U calculations. The fitting parameters \(M_1\) and \(M_{2,3}\) represent the saturation magnetizations of the first layer, and the second and third layers, respectively. Since the exchange field is much larger than the external magnetic field, \(\tanh\left(\frac{B - B_{c1}}{k_B T}\right)\) is approximately \(-1\), leading to a field-independent antiparallel contribution \(-M_1\) originating from the first layer. We fit the second and third layers’ contribution paramagnetically because of the weakness of the intermolecular interaction. The fit yields \(M_{2,3} = 1.8 |M_1|\), indicating that the film thickness is approximately 2.8 ML, which is in good agreement with our assumption of 3 ML.

Our DFT+U calculations also reveal an antiferromagnetic coupling between the Co moment in CoP and the Ni surface. However, the coupling strength is much smaller compared to chemically adsorbed molecules on magnetic surfaces [23]. Due to the presence of the graphene layer, CoP is physisorbed which significantly weakens the magnetic coupling. The chemical properties of graphene are also modified due to strong hybridization with the Ni(111) layer. A perfect lattice matching helps one of the C \((C_A)\) sublattice atoms to chemically bind to Ni atoms, while the other C sublattice \((C_B)\) atoms remain sp^2 bonded. This imbalance induces different magnetizations on different C sublattices. \(C_A\) develops a moment of 0.018 \(\mu_B\) and aligns antiparallel to the Ni moment, while \(C_B\) gets magnetized parallel to the Ni moment with a moment of 0.03 \(\mu_B\), yielding a ferrimagnetic graphene layer. We have, in the calculations, considered CoP adsorption sites on top of \(C_A\), \(C_B\), and in the middle of the graphene hexagon, which we refer to in the rest of this paper as Top-A, Top-B, and Hexagon, respectively. We find from our calculations that the physisorbed CoP finds itself spatially 3.07, 3.04, and 3.11 Å away from the graphene layer for the Top-A, Top-B, and Hexagon sites, respectively. The Top-A site is found to be the most stable one, while the Top-B and Hexagon sites are 14.6 and 23.5 meV higher in energy.

From the total-energy calculations, we find that the central Co atom in CoP couples antiferromagnetically with the underlying Ni surface for all adsorption sites, in agreement with our experiments. In contrast to a previous study [10] for a similar system, we find that the exchange-coupling energies \((E_{FM} - E_{AFM})\), where \(E_{FM}\) and \(E_{AFM}\) represent the total energies in the ferromagnetic and the antiferromagnetic orientation, respectively) are small (4.2, 9.9, and 3.1 meV/cell for Top-A, Top-B, and Hexagonal adsorption sites, respectively). While oxygen as a buffer layer has a saturation coverage at half a monolayer, e.g., on Co/Cu(100) [7,8], leading to well separated and localized O \(p_z\) orbitals, the C \(\pi\) orbitals build a delocalized \(\pi\) band. Due to this delocalization, the molecules couple antiparallel, independent of their absorption site. We want to comment that if CoP is replaced by FeP, studied in Ref. [24], Fe couples ferromagnetically to Ni for the Top-A site due to the involvement of different types of orbitals in mediating the exchange interaction.

To understand the exchange interactions, we present, in Fig. 3, the density of states of the out-of-plane orbitals for CoP-Co, (CoP)-C, (CoP)-N, (Graphene)-C, and surface layer Ni for the most stable configuration, Top-A. The exchange splittings of the Co and Ni states demonstrate the antiferromagnetic configuration, which is further vividly represented in the magnetization density plot (inset). The magnetization density (blue) that appears on the Co atom is solely from a singly occupied \(d_{z^2}\) orbital and couples antiferromagnetically with Ni moments (red). It may be observed that the Ni 3d and graphene \(p_z\) states hybridize and should therefore have a stronger chemical interaction than between graphene and...
CoP. Also, there is no hybridization visible between Ni and Co 3d orbitals, and which excludes a direct exchange mechanism between CoP and Ni substrate. Instead, exchange is indirect and is mediated via the graphene states.

To shed more light on the exchange coupling between surface Ni and Co atoms, we have analyzed the orbital dependence of the susceptibility, which is proportional to the interatomic pair-exchange interaction (considering energy-independent local potentials), expressed as \( \chi_{ij}^{\text{\text{exchange}}} (E_F) \propto \text{Im} \int dE \sum_{m,m'} G_{m,m'}^{ij} (E) G_{m,m'}^{ji} (E) \), with \( m, m' \) being magnetic quantum numbers (\( m = \pm 1 \)) for d orbitals, while \( G_{m,m'}^{ij} \) denotes the Green function between \( i \) (Co) and \( j \) (Ni atom beneath the C atom of the graphene layer in the Top-A position; see Fig. 3) atoms for the spin channel \( \sigma (\sigma = \uparrow, \downarrow) \). The orbital-projected Green functions were generated by the projection of Kohn-Sham orbitals onto spherical harmonics in a sphere around each atom. The \( m \)-dependent susceptibility as a function of Fermi energy \( E_F \) is shown in Fig. 4. In these calculations, first \( E_F \) was determined self-consistently (denoted as zero in Fig. 4). Then the value of \( E_F \) was varied using a rigid band model by fixing the self-consistent electronic structure and calculating the orbital-dependent \( \chi^{ij} \) for each \( E_F \). It can be observed from Fig. 4 that the in-plane orbital contributions are almost zero throughout, while the out-of-plane (\( d_{xz} = d_{dz^2} + d_{dz^2} \)) contribution is significantly below the Fermi energy. At \( E_F = 0 \), the major contribution comes from the \( d_{z^2} \) orbital with a relatively small contribution from \( d_{2z} \) orbitals. Note also that the contributions for \( d_{z^2} \) and \( d_{2z} \) orbitals are of different sign, meaning that the total contribution (summed over orbitals) will be slightly reduced.

Now we discuss the exchange interactions between the molecular layers. As argued before, based on our experiments, the top and middle layers behave like paramagnets, while the lowest layer is exchange coupled to the Ni substrate via graphene. We have found from our calculations for a bilayer of CoP that the high-symmetric, perfectly parallel CoP layers are antiferromagnetically coupled to each other. The optimized geometry has the two Co atoms in two consecutive layers shifted laterally, which is also discussed in the literature for CoPc [25]. The spatial separation between the parallel layers reduces the exchange strength considerably due to lowering of orbital overlap, while randomness in the orientation has a higher impact in reducing interlayer exchange coupling. In Fig. 4 (inset), the sharp fall of exchange energy due to increasing separation between two CoP layers and angular deviations between them is shown. Two CoP molecules making an angle to each other not only increases the effective distance between Co atoms, but also causes the type of orbitals participating in the exchange to be modified. This results in a drastic decoupling of the exchange interaction between the layers. As mentioned before, this randomness in orientation can occur for dense molecular layers on surface due to the outermost ligands in CoOEP. The applied magnetic field thus can orient these loosely coupled layers and turn the whole system ferromagnetic, which confirms the argument of field-dependent switching of the magnetization. It should be noted that the out-of-plane magnetic anisotropy energy is high (0.19 meV) compared to the extremely small in-plane anisotropy (easy-plane system) as given by our DFT + U calculations for CoP in the gas phase, in which we included spin-orbit coupling. This indicates that the field-dependent switching most probably takes place by in-plane rotation of Co moments.

In summary, we have demonstrated the switching of magnetization of three layers of CoP deposited on 1 ML graphene-covered Ni(111) substrates by external magnetic fields. From our analysis, with the aid of density functional calculations, we find that the CoP layer adjacent to the graphene layer is strongly exchange coupled antiferromagnetically to the Ni.
substrate, whereas the top and the middle CoP layers are coupled via very weak, short-ranged interactions and, hence, can easily be switched by external fields. This achievement is a result of an effective decoupling of the second and third molecular layers, generated by the graphene interlayer, which is the key to the field-regulated switching. The realization of a field-regulated switching of molecular magnets is an important step ahead towards future molecular spintronic concepts, as the spin injection into the organic layer can be regulated by a magnetic field.

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