Magnetism of FePt Surface Alloys

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The complex correlation of structure and magnetism in highly coercive monoatomic FePt surface alloys is studied using scanning tunneling microscopy, x-ray magnetic circular dichroism, and *ab initio* theory. Depending on the specific lateral atomic coordination of Fe either hard magnetic properties comparable to that of bulk FePt or complex noncollinear magnetism due to Dzyaloshinski-Moriya interactions are observed. Our calculations confirm the subtle dependence of the magnetic anisotropy and spin alignment on the local coordination and suggest that 3D stacking of Fe and Pt layers in bulk $L1_0$ magnets is not essential to achieve high-anisotropy values.

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Several exciting experiments on Fe atoms in nanostructures have shown recently that the magnetism of lowcoordinated Fe is very different from bulk, rich in features, with a strong tendency towards nonparallel alignment of neighboring spins. Examples include antiferromagnetism in Fe monolayers on tungsten surfaces [1], and the complex noncollinear spin alignment in Fe monolayers on Ir(111) [2]. Common to these results is the presence of lowcoordinated Fe atoms and bond lengths different from bulk Fe, as well as the direct contact between Fe atoms and a 5d element substrate. The complexity of the magnetism in such systems is a result of the quasi-2D interplay between exchange interaction, magnetic anisotropy, induced moments in 5d atoms, and the emergence of Dzyaloshinski-Moriya (DM) [3] interactions due to the broken inversion symmetry. In this work, we succeeded in embedding Fe atoms directly in the surface layer of a Pt(997) substrate and created a new class of lowdimensional materials in which mechanisms that lead to complex magnetic behavior can be studied in more detail. STM images with atomically resolved chemical contrast are presented and the system's magnetic properties are analyzed by x-ray magnetic circular dichroism (XMCD). We show that depending on the specific lateral coordination of Fe the magnetism of these surface alloys (SA) can exhibit large out-of-plane magnetic anisotropy, comparable to that of high-anisotropy bulklike FePt alloy films (10–100 nm thickness) [4–6], and noncollinear magnetism due to DM interactions. While in bulk FePt large anisotropies are due to noncubic 3D stacking of Fe and Pt layers [7.8], we find a substantially different mechanism for SAs.

FePt SAs are grown on Pt(997) substrates with a highly regular step periodicity of 2 nm [9]. FePt alloy layers

confined to the topmost layer of the Pt(997) surface form during molecular-beam epitaxy (MBE) deposition of submonolayer coverages of Fe at temperatures of 500 to 550 K. The substrate and alloy preparation and structural characterization is described in Ref. [10]. STM images shown in Fig. 1 resolve SA structures that form during



FIG. 1 (color). STM images of Fe_xPt_{1-x} SAs formed during deposition of Fe atoms onto Pt(997) at 525 K. (a) $Fe_{50}Pt_{50}$: Mixture of Fe and Pt atoms in the surface layer consistent with the expected stoichiometry. Fe 2 × 2 and 2 × 1 superstructures on a local scale are clearly visible (right side). (b) $Fe_{25}Pt_{75}$: Partial $Fe_{50}Pt_{50}$ SA with preferential 2 × 1 superstructure confined close to the step edges.

the deposition of Fe at 525 K. The SA grows from the step edges into the terraces; offering less Fe leads to an alloy which is confined to the area around the step edges, as shown in Fig. 1(b) for Fe₂₅Pt₇₅. Figures 1(a) and 1(b) show equiatomic (Fe₅₀Pt₅₀) and iron-poor (Fe₂₅Pt₇₅) coverages, respectively. The enlarged areas of individual terraces in Fig. 1(a) show typical SA configurations, especially 2×1 and 2×2 structures, which are also illustrated in the cartoons of Fig. 1. We observe that the degree of disorder depends on the iron concentration; the 2×1 and—especially— 2×2 structures are more predominant for low Fe coverages. In equiatomic SAs (Fe₅₀Pt₅₀), shown in Fig. 1(a), the 2×1 and 2×2 structures are confined to very limited areas. We find the lateral atomic environments of the Fe and Pt atoms in Fig. 1 to be of crucial importance for the understanding of anisotropy and coercivity.

The *in situ* magnetic characterization of the FePt SAs was carried out using XMCD measurements [11] at a fixed temperature of T = 12 K and magnetic fields up to 2.5 T. From the difference between absorption spectra (XAS) for opposite sample magnetization \tilde{M} the magnetic moments projected on the vector of the incoming photon spin can be determined [12]. In order to probe the systems magnetic anisotropies we measured the magnetic behavior in different directions parallel and perpendicular to the substrate steps, in the $(11\overline{2})$ and $(\overline{1}10)$ planes, respectively. The polar angle θ is the angle of incidence of the x rays with respect to the surface normal, as shown in Fig. 2(a). The upper part of Fig. 2(b) displays typical XAS spectra at the Fe $L_{3,2}$ and Pt $N_{7.6}$ absorption edges of the Fe₅₀Pt₅₀ surface alloys. The difference in the XAS for opposite magnetic fields (plotted in the lower part as the XMCD signal) reflects a magnetization at both Fe and Pt absorption edges. The Pt exhibits a dichroic signal as a result of an induced magnetic moment



FIG. 2 (color). (a) Geometry: XMCD was measured along different Pt(997) directions parallel and perpendicular to the substrate steps (within the planes in blue and red, respectively). θ is the polar angle with respect to the surface normal. (b) XAS of the 0.5 ML FePt SA at the Pt $N_{7,6}$ (left) and Fe $L_{3,2}$ edges (right). The dashed two-step function accounts for the contribution of the 2p-4s transitions to the Fe XAS. The difference between the signals for opposite magnetization (blue and black traces) yields the XMCD spectra (red traces). The photon energies at which M(B) for Fe and Pt is measured [Fig. 3(a)] are indicated by vertical lines.

due to a strong hybridization between Fe 3*d* and Pt 5*d* states. The complicated Fano line shape at the Pt $N_{7,6}$ edges has been analyzed in detail in an earlier study of bulk CoPt₃ [13]. We used the dichroism at the Fe L_3 and the Pt N_7 absorption edges at 72.0 and 708.3 eV, respectively, to obtain element-specific magnetization curves. Figure 3(a) shows typical hysteresis loops for Fe and Pt along $\theta = 0^{\circ}$ and 70°. We observe a nearly square-shaped hysteresis loop along the surface normal ($\theta = 0^{\circ}$), corresponding to the polar magnetic easy axis (perpendicular anisotropy). The coercive field $B_c = 0.71$ T along this direction suggests the presence of a large magnetic anisotropy energy (MAE). With increasing polar angle θ , the loops loose their squareness [blue and red curves in Fig. 3(a)].

The $\theta = 70^{\circ}$ hysteresis loops exhibit an additional inplane anisotropy, since the two in-plane directions parallel and perpendicular to the step edges yield slightly different coercivities [Fig. 3(a), right]. This symmetry breaking is attributed to the coexistence of ordered 2×2 and 2×1 superstructures. Both surface structures support uniaxial MAE $\sim K_u \sin^2 \bar{\theta}$, and our XMCD experiments reveal easyaxis anisotropy ($K_u > 0$) ($\bar{\theta}$ is the angle between *M* and the surface normal). However, only the 2×1 , but not the $2 \times$ 2 surface, exhibits a K'_1 -type anisotropy [see Eq. (3.2) in Ref. [14]], which favors in-plane spin alignment parallel or perpendicular to the monatomic wires, depending on the sign of K'_1 . A similar effect is also expected from the residual influence of the steps.

We find congruent Pt and Fe hysteresis loops in Fig. 3(a), suggesting a collinear alignment of Fe and Pt moments. To distinguish between a ferromagnetic (FM) and anti-FM ordering, we observe that the sign of the XMCD signal at the Fe $L_{3,2}$ and Pt $N_{7,6}$ edges in Fig. 2(b) exhibits the same relation as found by Shishidou *et al.* for the Co $M_{3,2}$ and Pt N_7 edges of the bulk ferromagnet CoPt₃ [13]. By analogy, we conclude that in the FePt SA the Pt and



FIG. 3 (color). (a) Upper panels: Hysteresis of the Fe₅₀Pt₅₀ SA taken at the Fe L_3 edge for directions $\theta = 0^\circ$, 70° . The data are normalized to the nondichroic XAS signal. The measured directions are color coded with orientation labels explained in Fig. 2(a) and in the text. Lower panels: Corresponding measurements at the Pt N_7 edge. (b) Hysteresis of Fe₅₀Pt₅₀ for $\theta = 0^\circ$, 35° and 70° together with a Stoner rotation fit using $K_u = 0.35 \text{ meV/f.u.}$ and $m_{\text{f.u.}} = 3.53 \mu_B/\text{f.u.}$

Fe moments are FM coupled. Moreover, the relative size of the XMCD signal at the Pt N_7 edge suggests that the induced moment of Pt within the alloy layer is comparable to the one found in CoPt₃. For the latter, a Pt moment of $0.26\mu_B/\text{atom}$ was determined by neutron scattering [15,16].

Our experimental results are supported by fully relativistic ab initio density functional theory (DFT) simulations. The electronic structure of the surface alloys on Pt(111)was calculated by the KKR multiple scattering formalism, and the local LSDA was used for the exchange and correlation effects. The anisotropy was extracted from calculations of the magnetic torque, and in the investigation of random SAs, the disorder was treated in the coherentpotential approximation. The calculations confirm a parallel alignment of Fe and Pt spins, and predict induced Pt moments of $0.27 \mu_B$ per Pt atom in a FePt layer with 2×1 structure on Pt(111) and $0.21\mu_B$ per Pt atom for the disordered FePt surface alloy. For the first subsurface layer we obtain Pt moments of about $0.2\mu_B$ and $0.18\mu_B$, respectively. The magnetism of the surface alloy depends on the Fe and the Pt sublattice due to interatomic hybridization, hence it has to be treated as one system. It is best defined as the top Fe₅₀Pt₅₀ alloy layer plus the Pt layer underneath, giving an overall chemical composition, or formula unit (f.u.), of FePt₃.

Using DFT, we systematically calculated various FePt model structures on Pt(111), as observed in our STM experiments. Pt(997) steps were simulated by a finite monolayer high cluster of Pt on Pt(111). For the Pt-rich 2×2 structure on Pt(111), we found an AFM coupling between Fe nearest neighbors, leading inevitably to magnetic frustration due to the hexagonal symmetry of the SA [Fig. 1(a)]. By contrast, the simulation of ideal 2×1 SA structures yields a strong FM exchange within each Fe chain of 60 meV between neighbor atoms. The coupling between adjacent Fe chains, however, is much smaller, about 1 meV per atom. It will be shown below that this weak interchain coupling facilitates the formation of complex spin configurations due to DM interactions. The simulation of random Fe₅₀Pt₅₀ SAs yields strong FM exchange of about 48 meV between Fe nearest neighbors.

The magnetic hysteresis of nanoscale systems with rigid FM coupling is well-described by the coherent-rotation or Stoner-Wohlfarth model. The calculation of M(B) and the extraction of the anisotropy constant K_u requires the knowledge of the magnetic moment per Fe atom, which includes both Fe and the induced Pt moments. The Fe moment was estimated from the XMCD spectra using the sum rules for 3d metals [17,18]. From the saturated spectra along the magnetic easy axis we derive an effective spin moment per hole, $m_S^{\text{eff}} = m_S + 7m_T$, of $0.62\mu_B/\text{hole}$, where m_T is the magnetic dipole term accounting for the anisotropy of the spin density. Because of the influence of m_T the actual spin moment m_S per hole in the polar direction is expected to be approximately 15%–20% larger

than the experimentally determined m_S^{eff} . Under the assumption of 3.7 *d*-band holes per atom [19], the actual spin moment per Fe atom is $2.75\mu_B$, which is fairly close to the DFT prediction ($m_S = 3.2\mu_B$). Taking the experimental Fe moment and adding three Pt neighbors with $\mu_{\text{Pt}} = 0.26\mu_B$ each, we obtain a total moment of $3.53\mu_B$ per FePt₃ formula unit (or Fe atom).

Figure 3(b) compares our M(B) data at $\theta = 0^{\circ}$, 35° and 70° with the Stoner-Wohlfarth model. Best agreement is achieved for an effective magnetic anisotropy constant $K_{\mu} = 0.35$ meV per Fe atom. This anisotropy is comparable to that of highly anisotropic $L1_0$ FePt bulk alloys, $K_u = 5 \text{ MJ/m}^3 (5 \times 10^7 \text{ erg/cm}^3)$ [20], or 0.8 meV per Fe atom (In terms of anisotropy fields, $1 \text{ meV}/\mu_B =$ 34.6 T). The strong spin-orbit coupling of Pt, $\xi_{Pt} =$ 0.6 eV, compared to Fe, $\xi_{Fe} = 0.08$ eV, is known to significantly contribute to the anisotropy. Figure 3 shows that the Stoner-Wohlfarth model works well in the coherentrotation regime, that is, at larger fields, and adds evidence to the existence of a magnetically stable monatomic Fe₅₀Pt₅₀ alloy layer. The much less perfect agreement in small field, especially for $\theta = 0^\circ$, is a common phenomenon caused by real-structure imperfections [14] and, in the present case, by the neglect of magnetostatic interactions. Both effects reduce the coercivity.

The magnetization loops exhibit a delicate dependence on the details of the Fe-Pt coordination. Figure 4 compares the polar magnetization curve of the Fe₅₀Pt₅₀ SA in (a) with loops of a SA layer formed from only 0.35 ML Fe at 525 K (b), and with nonalloyed Fe stripes of 4 atoms' width formed from 0.5 ML Fe at 350 K (c). Figure 4(b) shows that confining the alloy to a region close to the step edges drastically alters the loop shape. Compared to the square loop of the Fe₅₀Pt₅₀ alloy, the loop in (b) is *S* shaped, and remanence and coercivity are reduced by 78% and 68%, respectively. The overall shape of the loop (b) is very similar to that of the nonalloyed Fe stripes of 0.5 ML Fe (c). Note that the available field of 2.5 T is insufficient



FIG. 4 (color). M(B) perpendicular to the surface at the Fe L_3 edge of (a) the FePt SA prepared with 0.5 ML Fe, (b) a SA prepared with 30% less Fe, and (c) a nonalloyed Fe-stripe with 0.5 ML Fe. All three samples can be shown to exhibit a perpendicular easy magnetization axis.

to saturate the magnetization, however, full saturation has been achieved at B = 6 T on nonalloyed 0.5 ML Fe on Pt(997) in separate experiments at a different beam line. What makes $Fe_{50}Pt_{50}$ surface alloys such hard magnets? High coercivity and slow saturation require high magnetocrystalline anisotropy and strong exchange coupling. In $L1_0$ -ordered bulk FePt, the strong Fe-Fe intralayer exchange provides the basis for magnetic order and ferromagnetism. Our experiments and the *ab initio* theoretical results show that the situation in two-dimensional FePt is more complex. Here, strong out-of-plane anisotropy is found for equiatomic FePt coverages (0.5–0.6 ML Fe) where the area of the ordered regions is substantially decreased. The DFT simulation of the statistically disordered Fe₅₀Pt₅₀ SA yields a weak in-plane anisotropy of 0.09 meV per Fe atom, while a full Fe monolayer has a weak out-of-plane anisotropy of 0.03 meV per Fe atom. By contrast, for a perfect 2×1 structure, our calculations yield a very large in-plane MAE of 1.06 meV/f.u.. Intriguingly for this system DM [3] interactions play an important role. The calculated DM coupling vector \vec{D}_{ij} between two closest magnetic moments of Fe in different chains is of the same order of magnitude as the ordinary exchange coupling $(|\vec{D}_{ij}| = 4.6 \text{ meV}; J = 8.8 \text{ meV})$. This leads to the creation of a noncollinear magnetic structure with coupled in-plane and out-of-plane components of the total magnetic moment. In this case the discussion of magnetic anisotropy on the basis of M(B) becomes difficult: for instance, when in- and out-of-plane anisotropies for individual sites have similar magnitudes it will result in similar magnetization curves that cannot be distinguished in a simple way. This effect is particularly pronounced for the system with 0.35 ML Fe coverage [Figs. 1(b) and 4(b)] as it consists of almost perfect stripes of the Fe-Pt 2×1 alloy and it offers a convincing explanation of the out-of plane magnetization component and of the lower magnetization at 2.5 T compared to Fe₅₀Pt₅₀.

As the Fe coverage is increased to about 0.5–0.6 ML [Figs. 1(a) and 4(a)], 2×1 stripes are destroyed by the substitution of Fe into the Pt chains. This results in the formation of small Fe-rich surface regions as those sketched in Fig. 4(a) with lateral Fe-Pt interfaces. Our calculations show that these islands of strongly exchange coupled Fe atoms exhibit a pronounced out-of-plane anisotropy of 0.3–0.5 meV per Fe atom. Without the Fe-rich areas, that is, for a Fe concentration just below 50% of a full monolayer, the anisotropy is small, as discussed earlier. A slight increase of the Fe concentration beyond 50%-60% results also in a significant decrease of the anisotropy, and for a Pt-rich 2×2 structure (Fe₆₅Pt₃₅), the calculated anisotropy is only 0.15 meV per Fe atom. Thus, FePt SAs containing 0.5-0.6 ML of iron seem to hit the "sweet spot" for strongest out-of plane anisotropy.

In conclusion, the key result of this paper is in the identification of structural and magnetic phases in FePt surface alloys, depending on the Fe concentration, and the complex mechanism that leads to a maximum in the magnetic anisotropy for a Fe concentration of 50%–60%. The FePt 2 × 1 stripe phase exhibits an intriguing interplay between exchange and DM interaction, which results in noncollinear spin structures that appear in the experiments in the form of reduced anisotropy and total moment. Our work, thus, extends recent studies on noncollinear spin-structures of full Fe monolayers on other 5*d* transition metal surfaces [1,2] to surface alloys. Strong out-of-plane anisotropy comparable to bulk $L1_0$ alloys are achieved for some surface alloy structures. Both bulk $L1_0$ and surface alloy exploit the strong spin-orbit coupling of Pt, but the anisotropy is realized differently in the two structures, and the surface anisotropy exhibits a more complex physics.

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