Fe Spin Reorientation across the Metamagnetic Transition in Strained FeRh Thin Films


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A spin reorientation accompanying the temperature-induced antiferromagnetic (AFM) to ferromagnetic (FM) phase transition is reported in strained epitaxial FeRh thin films. $^{57}$Fe conversion electron Mössbauer spectroscopy showed that the Fe moments have different orientations in FeRh grown on thick single-crystalline MgO and in FeRh grown on ion-beam-assist-deposited (IBAD) MgO. It was also observed, in both samples, that the Fe moments switch orientations at the AFM to FM phase transition. Perpendicular anisotropy was evidenced in the AFM phase of the film grown on IBAD MgO and in the FM phase of that grown on regular MgO. Density-functional theory calculations enabled this spin-reorientation transition to be accurately reproduced for both FeRh films across the AFM-FM phase transition and show that these results are due to differences in strain.

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Equiatomic FeRh has raised growing interest in the last decade because of the fundamental properties of its first order phase transition [1–6] as well as its potential applications in thermally assisted magnetic recording (TAMR) [7,8]. The occurrence of a temperature- or field-induced metamagnetic transition from antiferromagnetic order (AFM) to ferromagnetic order (FM) $\sim 350$ K makes it a very interesting material for storage media applications [7].

The existence of a magnetic phase transition in equiatomic FeRh was seen by Fallot in 1938 [9], but it was only in 1961 that the low-temperature phase was identified as AFM, as reported by de Bergevin and Muldawer [10,11], and later confirmed by Kouvel and Hartelius [12] and Shirane et al. [13]. Note that the magnetic phase transition is accompanied by a lattice expansion of $\sim 1\%$ from the AFM phase to the FM phase. However, the origin of the transition is still debated. The different origins suggested in the literature range from an increase in the electronic density of states (DOS) [14,15] to spin-wave excitations [3] and instability of the Rh magnetic moment [2,6]. An increased DOS in the FM phase was experimentally seen [16] but it is insufficient to account for the large entropy change at the transition. In addition, an excess specific heat was observed in the FM phase [17], consistent with magnetic contributions to the entropy. It appears likely that the transition is caused by a combination of these phenomena. Time-resolved magneto-optical Kerr effect measurements showed that the lattice expansion occurs later than the onset of ferromagnetism in the photon-induced AFM-FM phase transition [18,19]. However, recent time-resolved x-ray circular dichroism [20] and a combination of time-resolved x-ray diffraction and magneto-optical Kerr effect experiments [21] did not confirm this result and suggest that magnetic and structural changes through the transition occur simultaneously.

In this Letter, we report on the study of the AFM-FM transition in epitaxially grown FeRh films by $^{57}$Fe conversion electron Mössbauer spectroscopy (CEMS). Such a microscopic technique is a very sensitive tool for the investigation of the magnetic properties of Fe-based thin films via nuclear hyperfine interactions [22–25]. In particular, this technique enables us to determine the direction of the magnetic moment of the Fe atoms with respect to the incident $\gamma$ radiation in both AFM and FM phases. CEMS allowed a spin-reorientation transition to be observed through the AFM-FM transition; i.e., the preferred direction of the Fe magnetic moments changes between the AFM and FM phases. Using density-functional theory (DFT), we show that the spin-reorientation transition is well understood as a result of an electronic structure modification under tetragonal lattice distortion, leading to a different magnetocrystalline anisotropy (MCA) for the FM and AFM states of these thin film samples.

$\sim 150$ nm thick Fe$_{0.98}$Rh$_{0.02}$ films were grown epitaxially at 573 K by magnetron sputtering from an equiatomic FeRh target onto both a thick single-crystal (001) MgO substrate and an a-SiO$_2$/Si substrate coated with ion-beam-assist-deposited (001) MgO (IBAD MgO) [26]. This IBAD growth produces a biaxially-textured MgO film, whose lattice parameters are given in [26], onto which an epitaxial body-centered tetragonal FeRh film can be grown. The samples were post-annealed for 2 h at 873 K.
TABLE I. Room temperature lattice constants (measured for the AFM phase and extrapolated for the FM phase using the experimental linear thermal expansion coefficient of $1.13 \times 10^{-5}$ K$^{-1}$, as determined from the thermal variation of the out-of-plane lattice constant), corresponding $c/a$ ratio, and calculated spin magnetic moments (bold) of FeRh thin films in the AFM and FM phases.

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<th>AFM</th>
<th>FM</th>
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<tr>
<td></td>
<td>$a=b$ (Å)</td>
<td>$c$ (Å)</td>
</tr>
<tr>
<td>FeRh//MgO</td>
<td>2.980</td>
<td>3.004</td>
</tr>
<tr>
<td>FeRh//IBAD MgO</td>
<td>3.005</td>
<td>2.959</td>
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The composition of the films was determined via Rutherford back-scattering (RBS) measurements, and both films were shown to be epitaxial by four-circle x-ray diffraction (XRD) which showed four in-plane (110) peaks. Temperature-controlled XRD data in the temperature range 300–500 K showed a tetragonal distortion of the films, particularly in the AFM phase. Because films typically relax strain by nucleating defects, a higher strain is likely in thinner films. The lattice constant, corresponding to a broad distribution of spin orientation and indicating that for this sample, the Fe spin orientation in the FM phase no longer lies in the film plane.

The hysteretic behavior of the hyperfine field as a function of temperature is seen, and the transition temperature upon heating is found to be $390 \pm 10$ K for Fe$_{0.98}$Rh$_{1.02}$/MgO heated up to 450 K (FM phase), $x \sim 2$ [Fig. 1(a)], corresponding to a broad distribution of spin orientation and indicating that the 2nd and 5th Mössbauer lines are not completely absent, as it should be for a truly perpendicular orientation of the Fe magnetic moments, is attributed to the imperfect collimation of the $\gamma$ ray beam. For Fe$_{0.98}$Rh$_{1.02}$/MgO measured upon cooling at 340 K (FM phase), the increase of the line intensity $x$ up to $\sim 4$ on the Mössbauer spectrum indicates a spin-reorientation transition from out-of-plane in the AFM phase to in-plane in the FM phase [Fig. 1(d)].

The hyperfine field plotted as a function of temperature [Figs. 2(a) and 2(b)] shows that the local magnetic moment carried by Fe is higher in the FM phase than in the AFM phase for both samples, in agreement with results obtained on bulk samples [13] and polycrystalline thin films [29]. The hysteretic behavior of the hyperfine field as a function of temperature is seen, and the transition temperature upon heating is found to be $390 \pm 10$ K for Fe$_{0.98}$Rh$_{1.02}$/MgO [Fig. 2(a)] and $350 \pm 10$ K for Fe$_{0.98}$Rh$_{1.02}$/IBAD MgO [Fig. 2(b)], in good agreement with the macroscopic magnetization data [Figs. 2(c) and 2(d)]. SQUID magnetometry measurements show that the transition temperature

![FIG. 1 (color online). CEMS spectra obtained at different temperatures on (a) Fe$_{0.98}$Rh$_{1.02}$/MgO and (b), (d) Fe$_{0.98}$Rh$_{1.02}$/IBAD MgO. The room temperature spectra (a),b) correspond to the AFM phase and the high-temperature spectra (c),(d) to the FM phase. Note that (c) was obtained upon heating and (d) upon cooling.]

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magnetization above the transition is grown on IBAD MgO [Fig. 2(d)], and in both cases the $k_{31}$. To integrate over the 3D Brillouin Zone (BZ) a method in multiple scattering formulation was applied Kohn-Rostoker (KKR) Green’s function band structure the local spin density approximation (LSDA). To calculate framework of density-functional theory (DFT), using served experimentally, our CEMS measurements were ac-

therefore lowers the transition temperature. FeRh FM phase, which naturally stabilizes this phase and MgO in this present work provides a closer match to the

Experimental results in [30] indicated that the spin magnetic moment in the AFM state is equal to zero, which is in qualitative agreement with the experimental increase of the hyperfine field by $\sim 6\%$.

The MCA energy of the FM and AFM states is plotted in Fig. 3 as a function of the $c/a$ ratio, exhibiting a nearly-linear dependence for each magnetic state. The tetragonal distortion breaking the cubic symmetry of the system results in a modification of the electronic structure, which leads to an energy difference between the states with out-of-plane and in-plane magnetization directions. By varying the spin-orbit coupling (SOC) of each component away from its actual value, we find a pronounced decrease of the MCA when the SOC for Rh approaches zero, both in FM and AFM states, which indicates the important role of the Rh atoms for the MCA in FeRh films. While in the FM state the hybridization is different for spin-up and spin-down states, resulting in a nonzero magnetic moment, in the AFM state the hybridization is similar for both spin states, which indicates the important role of the Rh atoms for the MCA in FeRh films. While in the FM state the hybridization is similar for both spin states, which indicates the important role of the Rh atoms for the MCA in FeRh films.
magnetization data (at 400 K) is the shape anisotropy, whose value deduced from the experimental in-plane anisotropy, as found experimentally, and the FM phase. When \( c/a > 1 \) (corresponding to FeRh///IBAD-MgO), the calculations yield the out-of-plane (z direction) easy magnetization direction for the AFM state \((c/a = 0.985)\) and in-plane easy direction for the FM state \((c/a = 0.993)\), in full agreement with the experiment. When \( c/a < 1 \) (corresponding to FeRh///MgO), the AFM state \((c/a = 1.008)\) has an in-plane anisotropy, as found experimentally, and the FM state \((c/a = 1.016)\) an out-of-plane MCA direction, while experimentally a broad distribution of Fe moment orientation was seen for the FM phase. Note, however, that the results presented in Fig. 3 do not account for the shape anisotropy, whose value deduced from the magnetization data (at 400 K) is \( \sim 7.6 \times 10^5 \) J/m\(^3\) (1.28 \times 10^{-4} \) eV/f.u.). Calculation of the MCA for \( c/a = 1.016 \) gives 0.45 \times 10^{-4} \) eV/f.u., a factor of 3 smaller than the shape anisotropy. The existence of out-of-plane moments could result from an underestimate of the MCA by the DFT calculation [34], or from a low anisotropy magnetic domain configuration in which magnetic moments point out-of-the plane, even though the MCA energy is lower than the demagnetizing energy [35]. Since the latter favors an in-plane configuration of the magnetic moments, the fact that the direction of the Fe magnetic moments was experimentally found to be distributed over a broad range of orientations in the FM phase of FeRh///MgO indicates the onset of a spin-reorientation towards the normal to the film plane, and suggests the possibility of stabilizing a perpendicular FM phase for a higher \( c/a \) ratio.

It should be noted that perpendicular anisotropy in FM FeRh (001)///MgO (001) had already been suggested in [36] based on magnetization measurements showing coercivity along the hard axis; that is however weak evidence and no theoretical calculations were performed to understand the origin of this observation. It is important to mention that while Cao et al. [36] attributed this spin reorientation to magnetoelastic anisotropy originating from the lattice expansion across the magnetic phase transition (meaning it would exist only in the FM phase), our work shows that it actually results from the magnetocrystalline anisotropy and exists in both magnetic phases.

In summary, we have demonstrated that strained FeRh undergoes a spin-reorientation through the AFM-FM metamagnetic phase transition, which is very well confirmed by first-principles calculations accounting for epitaxial strain. The excellent agreement between theory and experiment indicates that the spin reorientation is driven by strain, and is tuned by the use of different substrates. It is important to note that bulk samples do not exhibit such a spin reorientation since the magnetocrystalline anisotropy is zero for \( c/a = 1 \) in both phases. This effect is therefore specific to thin films and is likely to be even larger in thinner films such as those contemplated for TAMR. Strained thin films could be of considerable interest for future applications based on AFM spintronics, and the possibility of getting a FM FeRh thin film with perpendicular anisotropy deserves attention.

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