Electronic and magnetic structure of Ga$_x$Fe$_{1-x}$ thin films

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The electronic as well as magnetic properties of Ga$_x$Fe$_{1-x}$ thin films were studied by soft x-ray measurements. Using x-ray magnetic circular dichroism the Fe majority-spin band was found to be completely filled for $x \approx 0.3$. With further enhanced Ga content, the Fe moment as well as the angular dependence of the x-ray magnetic linear dichroism decrease strongly, which we attribute to the formation of D0$_3$ precipitates. Moreover, the magnetocrystalline anisotropy drops significantly.

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In a magnetic material, the spin-orbit interaction, a purely relativistic effect between the electron spin and its orbital moment, couples the magnetic moments to the lattice. It is at the electronic origin of magnetocrystalline anisotropy, i.e., the energy associated with the (re)orientation of the magnetization relative to the lattice, as well as magnetostriction referring to the deformation of a magnetic material when subjected to an external magnetic field. In magnetostrictive thin films, the adaptation to new dimensions is inhibited by bonding to the substrate and the application of a magnetic field in the film plane does not result in an expansion or contraction along the field direction. Instead a magnetoelastic stress is created that leads to a change in magnetocrystalline anisotropy.

Here we study the magnetic as well as electronic properties of Ga$_x$Fe$_{1-x}$ films deposited on GaAs(001). In bulk form, Ga$_x$Fe$_{1-x}$ exhibits extraordinary magnetostrictive properties as function of composition implying a strong dependence of the underlying spin-orbit coupling on $x$. Ga$_x$Fe$_{1-x}$ thin films are therefore an ideal candidate to study the composition dependence of the magnetic anisotropy and to correlate this with its electronic structure.

Using soft x-ray absorption spectroscopy, we find that the magnetic as well as electronic properties of Ga$_x$Fe$_{1-x}$ films change significantly for Fe content near $x \approx 0.3$. The holes in the Fe majority-spin band disappear with increasing Ga content and the Fe majority-spin band is completely filled for $x \approx 0.3$. With further enhanced Ga content, the Fe moment as well as the x-ray magnetic linear dichroism (XMLD) angular dependence decrease strongly, which we attribute to the formation of D0$_3$ precipitates.

For this study, 20 nm single-crystalline Ga$_x$Fe$_{1-x}$ thin films, $0 \leq x \leq 0.6$, were deposited by molecular-beam epitaxy as described in Ref. 5. X-ray absorption (XA) spectra were measured at ALS beamline 4.0.2 at ambient temperature with either 90% circularly or 100% linearly polarized x rays in total electron yield mode by monitoring the sample drain current. External fields of 0.5 T were provided by an eight-pole electromagnet. X-ray magnetic circular dichroism (XMCD) was derived from XA spectra measured with the magnetic field collinearly aligned to the incident x-ray beam, which in turn impinged at an angle of 30$^\circ$ to the sample surface. XMLD spectra are obtained as the difference between XA spectra in external fields parallel and perpendicular to the x-ray polarization in normal-incidence geometry.

Figure 1(a) shows the average of XA spectra obtained from Ga$_x$Fe$_{1-x}$ films with $0 \leq x \leq 0.6$. In the concentration dependence of the Fe $L_3$, XMCD shown in Fig. 1(b), we can distinguish two regions exhibiting distinct magnetic charac-

![FIG. 1. (Color online) Fe $L_3$ XA and XMCD spectra for Ga$_x$Fe$_{1-x}$. (a) Average XA spectrum. Inset: change in photon energy of the Fe $L_3$ edge derived from the XA (solid symbols) and XMCD spectra (open symbols). (b) XMCD spectra for varying Ga concentrations, $0 \leq x \leq 0.6$. Lighter colored lines correspond to data obtained from samples with higher Ga concentration. Top inset: Fe XMCD spectra near 711 eV for $x \approx 0.28$. Bottom inset: concentration dependence of the Fe $L_3$ XMCD signal.](image-url)
teristics. For \( x \approx 0.28 \), the Fe \( L_3 \) XMCD signal—and with that to a good approximation the average Fe magnetic moment—is nearly constant and starts at 55% of the Fe \( L_3 \) XA intensity in agreement with previous results for Fe films.\(^8\) For \( x \gtrsim 0.3 \), the Fe \( L_3 \) XMCD decreases almost linearly with Fe concentration. It is generally accepted that hybridization of the Fe and Ga states reduces the Fe magnetic moment.\(^9\)\(^-\)\(^12\) Detailed calculations aimed at determining the concentration dependence of the Fe moment in different lattice structures and sites have been performed. However, there are significant variations in the results suggesting that an estimate of the average Fe moment as a function of Fe concentration using theory is not unambiguous at present.

The inset in Fig. 1(b) shows the spectral features at the high-photon-energy side of the Fe \( L_3 \) edge for \( x \approx 0.3 \). For the pure Fe film, a small positive XMCD feature is observed at \( \approx 711 \) eV, which can be ascribed to transitions from the \( 2p \) core level to the unoccupied majority-spin band.\(^13\)\(^-\)\(^14\) The main negative Fe XMCD \( L_3 \) feature is due to transitions to the unoccupied minority-spin band. The intensity of the XMCD feature at \( \approx 711 \) eV reduces with increasing Ga content, i.e., the majority-spin band is gradually filled. For \( x \approx 0.3 \), the Fe \( L_3 \) XMCD becomes fully negative indicating that the majority-spin band is completely filled. We also see a shift in both the XA and XMCD \( L_3 \) peak energy to lower photon energy [inset Fig. 1(a)] consistent with band filling argument.

The occupation of majority-spin states is closely correlated with the magnetostriction. Berger\(^15\) showed that the magnetoelastic energy—and hence the magnetostriction—is determined by the electronic states located near the Fermi level, \( E_F \), and that it is directly proportional to the spin-orbit splitting. Consequently, the magnetostriction changes sign, i.e., zero magnetostriction is observed if \( E_F \) is located between the spin-orbit split bands. Filling of the majority-spin band also affects the magneto crystalline anisotropy energy (MAE).\(^16\)\(^-\)\(^17\) Using second-order perturbation theory, i.e., considering the spin-orbit coupling to be small compared to energy differences caused by the crystalline electric field (CEF) and exchange interaction, one can distinguish two contributions to the MAE; those due to spin-conserved terms and those arising from spin-flip processes. An incompletely filled majority-spin band allows low-energy spin-flip excitations near \( E_F \). When the majority-spin band is full, however, such excitations require an energy in the order of the exchange interaction. Hence, unoccupied majority-spin states enable spin-flip excitations, thereby changing the MAE. While the spin-flip contributions are usually not the dominant contribution in the total MAE, they will lead to a change in MAE.\(^12\)\(^-\)\(^16\)\(^-\)\(^17\)

Previous studies on transition metal and lanthanide oxide thin films\(^18\)\(^-\)\(^21\) have shown that the angular dependence of the XMLD provides a sensitive measure of the CEF symmetry and strength as well as magnetic and electronic characteristics of the absorber. We determined the Fe \( L_{1,2} \) XMLD spectra, \( I_{\text{XMLD}}(\phi) \), for varying angle \( \phi \) between the linear x-ray polarization, \( \mathbf{E} \), and the [100] lattice direction of Ga\(_{0.2}\)Fe\(_{0.8}\). The XMLD spectrum is defined here as the difference between XA spectra with \( \mathbf{H} \) parallel and perpendicular to \( \mathbf{E} \), i.e., at angles \( \phi \) and \( \phi + 90^\circ \) to the [100] direction.

\[
I_{\text{XMLD}}(\phi) = I_{\text{XA}}(\mathbf{H}_\phi, \mathbf{E}_\phi) - I_{\text{XA}}(\mathbf{H}_{\phi+90^\circ}, \mathbf{E}_\phi).
\]

The averaged Fe XA signal is shown in Fig. 2(a) and the XMLD angular dependence for \( 0^\circ \leq \phi \leq 45^\circ \) in Fig. 2(b). A pronounced variation in the Fe XMCD signal with \( \phi \) is observed. In lattices with cubic symmetry, the angular-dependent XMLD can be very well described by a linear combination of two fundamental spectra accounting for the XA photon energy dependence in selected orientations.\(^19\)\(^-\)\(^21\) Their relative contributions are determined by the symmetry of the experimental geometry. In the (001) plane, the XMLD angular dependence is described by
where \( I_0 \) and \( I_{45} \) refer to the XMLD spectra obtained with the x-ray polarization \( E \) at 0° and 45° to the [100] direction, i.e., along the [100] and [110] directions, respectively. Equation (2) allows us to separate the angular-dependent XMLD into an angle-independent or isotropic contribution to the XMLD signal, \( \frac{1}{2}(I_0 + I_{45}) \), and an angular part or anisotropic contribution, \( \frac{1}{2}(I_0 - I_{45}) \), which gives the difference in the XMLD along two principal directions. Figure 2(c) shows the experimental results for \( \frac{1}{2}(I_0 + I_{45}) \) and \( \frac{1}{2}(I_0 - I_{45}) \). Applying Eq. (2) and using the results from Fig. 2(b), we normalized the XMLD signals to the square of the Fe \( L_3 \) XMCD signal that to good approximation is a measure for the Fe magnetic moment. To account for the observed change in Fe moment with Ga content [see Fig. 1(b)], we normalized the XMLD signals to the square of the Fe \( L_3 \) XMCD signal that to good approximation is a measure for the Fe magnetic moment. Small changes in the spectral shape are attributed to changes in the detailed electronic structure also seen in the Fe XMCD as discussed above. Using the same normalization for the angle-dependent signal \( \frac{1}{2}(I_0 - I_{45}) \) leads to a still strongly decreasing XMLD signal with decreasing Fe concentration. As shown in Fig. 3(c) normalizing \( \frac{1}{2}(I_0 - I_{45}) \) to the Fe concentration, i.e., scaling it by \((1 - x)^{-1}\) as well as the square of the average Fe moment, leads to a constant anisotropic XMLD signal for \( x \approx 0.28 \) as well as for \( x \approx 0.3 \). This indicates that additional factors beyond the concentration dependence of the Fe moment play a role in determining the XMLD angular dependence. One possible explanation is that for \( x \approx 0.3 \) a volume fraction of the Ga\(_{0.3}\)Fe\(_{1-x}\) film consists of a phase where the Fe XMLD signal does not exhibit an angular dependence, i.e., is highly isotropic. The volume fraction of Fe atoms in this phase increases with decreasing Fe content to account for the decrease in \( \frac{1}{2}(I_0 - I_{45}) \) with \( x \).

In case of a completely disordered or isotropic system, no angular dependence is expected, i.e., \( I_0 = I_{45} = 0 \). Our reflection high-energy electron-diffraction (RHEED) measurements during film deposition indicate that the structural quality of the Ga\(_{0.3}\)Fe\(_{1-x}\) does not depend significantly on the Fe concentration, i.e., structural disorder cannot account for the reduction in \( \frac{1}{2}(I_0 - I_{45}) \) with increasing Ga content.

In epitaxial EuO thin films, the Eu \( M_{5,4} \) XMLD shows only a very small angular dependence, which could be explained by the nearly perfect spherical shape of the half filled Eu 4f shell. For Ga\(_{0.3}\)Fe\(_{1-x}\), Lei et al. calculated the anisotropy of the Fe 3d orbitals for D0\(_3\), B2-like, and L1\(_2\) lattice structures. They showed that the D0\(_3\) structure has an almost isotropic 3d charge density and that there is hardly any difference between the two Fe sites in this lattice structure. By contrast, a pronounced anisotropy was found for the other two phases. This suggests that Fe in local D0\(_3\) environments could be responsible for the reduction in the Fe XMLD angular dependence with decreasing Fe concentration. Moreover, van’t Erve et al. concluded from RHEED measurements during deposition of Ga\(_{0.3}\)Fe\(_{1-x}\) films on AlGaAs/GaAs quantum-well structures that the D0\(_3\) phase is present for \( x \approx 0.25 \). These results lead us to conclude that with increasing Ga content the D0\(_3\) phase develops and the fraction of Fe in these D0\(_3\) precipitates increases with decreasing Fe concentration. This accounts for the change in the angular-dependent XMLD with Fe concentration \( 1-x \).

Having discussed concentration dependence of the mag-
nitude of the XMLD signal we now consider the variation in the Fe XMLD spectral shape with Ga concentration. The XMLD spectral shape is influenced by the CEF symmetry and strength as well as the magnetic and electronic characteristics of the absorber. In case of Ga$_{x}$Fe$_{1−x}$ alloys, the CEF created by the nearest neighbors of the Fe absorber is not expected to be very sensitive to change in the elemental species of between Fe and Ga for identical structural parameters. As a consequence the spectral shape of the Fe XMLD signal is expected to be the same for bcc Fe and a disordered Ga$_{x}$Fe$_{1−x}$ alloy with the same bcc structure, i.e., Ga$_{x}$Fe$_{1−x}$ in A2 phase. The experiments show that the spectral shape of $\frac{1}{2}(I_2−I_3)$ is the same for all Ga concentrations $x$, i.e., identical to pure bcc-Fe ($x=0$). Therefore, we attribute the $\frac{1}{2}(I_2−I_3)$ signal to Fe atoms in the A2 phase.

To determine the impact of the concentration dependence of the Fe electronic structure in Ga$_{x}$Fe$_{1−x}$ films on their magnetic anisotropy, we applied the XMLD sum rules to the experimental and theoretical studies are certainly warranted to explain the differences in structure, magnetic, and electronic between bulk and thin-film Ga$_{x}$Fe$_{1−x}$ samples.

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