Growth and Magnetic Properties of Fe films on vicinal TO (001) substrates

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We report on investigations of the crystallographic structure and the magnetic anisotropies of epitaxial iron films deposited onto periodically stepped Ag(001) surfaces using low energy electron diffraction, x-ray diffraction, second harmonic generation (SHG), as well as the Brillouin light scattering (BLS) technique. The focus of the present study lies on the interrelation between the surface morphology of the buffer layers and the magnetic properties of the Fe films, epitaxially grown onto them. Especially the symmetry breaking at the atomic steps is found to create an uniaxial magnetic anisotropy measured by BLS and a very strong anisotropic signal in magnetic SHG.

The magnetic properties of ultrathin 3d transition metal films have been widely investigated in the last two decades. In recent years it was found that a reduction of the symmetry of these systems by depositing the films onto vicinal cut substrates leads to additional magnetic surface anisotropies [i,ii,iii,iv,v], their magnitude depending on the average step density [vi,vi]. These anisotropies originate from different contributions. Due to the broken symmetry at the step edges the magnetic environment changes. The reduced coordination number and the uniaxial strain induced by the lattice mismatch modify the band structure and thus the magnetic behavior. A great advantage of these step-induced anisotropies is that they can be artificially controlled by changing the vicinal miscut angle of the substrate, or, in other words, the average step density of the interface or surface [vi,vii]. On the other side these anisotropies can be used in future technical applications because of the artificially controllable character of the step density and so of the magnetic anisotropies.

Vicinal to (001) GaAs wafers with fcc Ag-buffers were used in this work as substrates for preparation of bcc Fe-films [viii]. The GaAs wafers of orientation (1 1 1) and (1 1 1), as well as (0 0 1)-oriented substrates for reference, were chemically cleaned and heated to 870 K for one hour in ultrahigh vacuum (UHV) to remove the oxides and adsorbates. The base pressure during preparation and deposition was typical lower than $5 \times 10^{-10}$ mbar. To achieve epitaxial growth of the 1500 Å thick fcc Ag-buffer layer the cleaned wafer was covered with a 10 Å thick Fe seed layer. After the deposition the Ag-buffer was annealed at 570 K for 1 hour to improve the surface morphology. bcc-Fe films of different thicknesses were deposited onto the Ag buffer with a deposition rate of 0.1 Å/s. The first 10 Å of Fe were grown at room temperature to avoid intermixing at the interface, followed by growth at T=370 K. To prevent the samples from oxidization and to insure symmetric interfaces both at the bottom and the top of the Fe-layer, the films were finally covered with 20 Å thick Ag film and 20 Å thick Cr cap layer.

The chemical cleanliness and the crystallographic structure of the films were controlled in-situ by Auger electron spectroscopy (AES) and low energy electron diffraction (LEED). The AES spectra indicate a typical amount of 5% or less of a monolayer Ag on top of the Fe-layer. After annealing the Ag-buffer the LEED images show a crossing of vicinal and lattice reflexes as a function of the electron energy (Fig. 1 a,b). This is a finger print of an ordered step structure on the surface of the Ag-buffer layer with an inclination angle relative to the atomic lattice. The periodicities of the steps [ix] with a terrace width of 11.5±2 Å and 18.0±2 Å determined from the LEED data deviate from periodicities resulting from the miscut angle measured by x-ray diffraction (XRD) (see next section) by less than 10%. LEED images taken from these Fe-films show that the energy depending spot splitting vanishes and the spots are elongated along the same direction (Fig. 1 c,d).

Fig. 1: LEED-images taken from the Ag-buffer (a, b) after annealing at 570 K showing an energy depending splitting of the spots (electron energy: a) 52.9 eV and b) 61.4 eV). The step density of the surface determined out of the images is in good agreement with the average step density determined from the XRD data. The images of the Fe-film (c, d) show elongated spots along the step direction. This points to a non-regular step distribution on the surface (electron energy: c) 96.1 eV and b) 122.0 eV). The (01)-spots of the surface unit cell are marked. The orientation of the sample (indicated by the coordinate system) is indicated.
This indicates a nonregular step distribution on the surface with an averaged terrace size of approximately 8 Å in the direction perpendicular to the steps.

The vicinal miscut of the substrate, the Ag-buffer layer and a 1000 Å Fe-film was verified by XRD. It was found, that the angles of crystal orientation with respect to the surface normal between the three materials deviate from each other. For the (1117)-oriented sample the vicinal angles of GaAs, Ag and Fe are 4.91°, 6.09° and 3.94° respectively. The angles for the (111)-oriented sample are 6.99°, 8.75° and 5.61°. As a matter of fact due to the different crystallographic structure of the substrate and the layers there is an out-of-plane lattice mismatch much larger than the in-plane lattice mismatch. The film-to-film variation of the vicinal angle can be understood by taking this into account [$x$] as follows. In Fig. 2 a sketch of a model is shown demonstrating the adoption of the out-of-plane lattice mismatch. For simplicity it is assumed that the adoption takes place just within one atomic layer (white layer in Fig. 2). The in-plane lattice mismatch is chosen to be zero. It is seen from Fig. 2 that the [001]-directions of the lower and the upper layer deviate from each other by an angle $\Delta \alpha$. The angle is given by geometric considerations: tan $\Delta \alpha = (a_1 - a_2)/t$, where $a_{1,2}$ are the out-of-plane atomic step heights (not to confuse with the lattice constants) and $t$ is the average terrace width. This additional inclination angle can be positive or negative depending on the sign of the out-of-plane lattice misfit. The linear step density $\rho$ depends on $\Delta \alpha$. The angle is given by geometric considerations: $\Delta \alpha = (a_1 - a_2)/t$, where $a_{1,2}$ are the out-of-plane atomic step heights (not to confuse with the lattice constants) and $t$ is the average terrace width. This additional inclination angle can be positive or negative depending on the sign of the out-of-plane lattice misfit. The linear step density $\rho$ is unchanged if this additional angle is small. This means that the tilt angle between the two materials generates only a small in-plane misfit. In the special case of GaAs-Ag and Ag-Fe interfaces the in-plane stress of the misfit parallel to the surface is even reduced. The stress can be reduced in an arbitrary number of layers in both materials depending on the elastic constants. In this way the total energy at the interface is minimized. In comparison with the experimental results we find that the theoretical value of the tilt angle is confirmed with an accuracy better than 5% for the Ag-Fe-interface. For the GaAs-Ag interface the measured value is only 55% of the theoretical value for both vicinal miscut angles, but the influence of the Fe-interlayer and the nonregular step distribution of the GaAs substrate on the additional miscut is unclear. Also the elastic constants of GaAs are much stiffer than those of the metal films, and the deformation takes place dominantly in the deposited layers. This strain is partly released by dislocation formation in the metal layers.

In order to determine the in-plane and the out-of-plane magnetic anisotropy contributions of twofold and fourfold symmetry, Brillouin light scattering (BLS) from spin waves in the Fe-films was performed. The anisotropy energy is given by:

$$E_{ui} = -K^{(2)}_2 \cos^2 \varphi + K^{(4)}_4 \sin^4 \varphi \cos^2 \varphi \sin^2 \varphi + K^{(2)}_y \sin^2 \varphi \cos^2 (\varphi - \varphi_0)$$

The in- and out-of-plane anisotropy constants $K^{(2)}$, $K^{(4)}$ with twofold symmetry and the in-plane anisotropy constant $K^{(4)}_y$ with fourfold symmetry were determined by fitting the measured spin wave frequencies as a function of the azimuthal angle $\varphi$ between the [100]-direction of the Fe-film and the applied external field $H$ with a least squares fit routine.

The details of the fit procedure and the underlying theory are described elsewhere [$x$]. Figure 3a shows the measured spin wave frequencies for a set of 20 Å thick Fe-layers as a function of $\varphi$. The maxima in the spin wave frequencies indicate magnetic easy directions. In addition to the fourfold anisotropy, characteristic for the sample without miscut, a...
The measurements were performed on the three different orientations of the layers \((1\ 1\ 11),\ (1\ 1\ 17)\) and, for reference, \((0\ 0\ 1)\) within a thickness range from 10 to 60 Å. While the out-of-plane anisotropy constant \(K^{(2)}_s\) and the in-plane anisotropy constant \(K^{(4)}_p\) are almost the same for all orientations, the twofold anisotropy constant \(K^{(2)}_p\) increases for the Fe-film thickness up to 30 Å with increasing film thickness (see Fig. 3b). The slope of the line is roughly proportional to the square of the vicinal angle. At about 40 Å the curves for both miscut angles show a characteristic maximum. This points to a relaxation behavior within the vicinal samples with increasing Fe-film thickness. This can be interpreted as a strain induced anisotropic effect. In combination with the additional miscut angle between the Ag and the Fe-layer we estimate that the out-of-plane lattice mismatch generates this anisotropy contribution, but with increasing thickness of the Fe-layer this stress is reduced or vanishes completely. This leads to a decreasing behavior of the twofold anisotropy constant. Generally the twofold anisotropy is of one order of magnitude larger than the values one would expect from the rotation of the magnetocrystalline tensor. This means that the anisotropy is likely to be a step induced anisotropy of the Fe-film. This result is confirmed by BLS measurements performed on 1000 Å thick Fe-layers displaying no twofold anisotropy contribution for all crystal orientations within the measurement accuracy.

Symmetry breaking at atomic steps on vicinal surfaces and interfaces make this system an ideal candidate for investigation with the magnetization induced second harmonic generation (MSHG) technique [xiii]. This technique is a nonlinear magneto-optical tool sensitive to both structural and magnetic properties of surfaces and interfaces. The strong surface/interface sensitivity is derived from the break in symmetry at boundaries between centrosymmetric media. The technique is non-invasive and therefore particularly suited for the study of buried interfaces. Measurements have been performed using a mode-locked Ti:sapphire laser operating at a repetition rate of 76 MHz with a pulse width of about 100 fs and a wavelength of 800 nm. The laser fluence was regulated such that the average laser power on the sample was 50 mW. The MSHG data were recorded in the transversal magnetic configuration at an angle of light incidence of 45°. After filtering the outgoing specular second-harmonic light, in order to suppress the fundamental frequency, the light was detected with a cooled photomultiplier as a function of the azimuthal rotation of the sample. Figure 4 demonstrates typical MSHG patterns from the 30 Å Fe-film for PP-polarization combinations. Shown is the magnetic contrast \(\rho\), defined by the difference of the MSHG-signal for two opposite field directions divided by the sum of the two signals. These rotational anisotropy patterns shows a very distinct asymmetry in the lobe structure and a 1-fold symmetry due to the vicinal nature of the Fe/Ag interface.

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