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Magneto-optical properties of MnBiAl thin films

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Mn-Bi-Al thin films were produced by sequential evaporation of the constituents, followed by an anneal at 300 °C. The temperature and composition dependencies of the Kerr rotation angle, absolute reflectivity, and magnetic anisotropy were measured. The results show that, up to 30 at. % Al concentration, the thin films retain the pure MnBi hexagonal structure. Further, for suitable Al content, the films have the same large Kerr rotation as MnBi. Pure MnBi films exhibit perpendicular anisotropy at room temperature and in-plane anisotropy for temperatures below 142 K. In contrast, the Al-doped films prepared here have perpendicular anisotropy down to at least 85 K. The increased coercivities of the Al-doped films are attributed to the occupation of grain-boundary and interstitial sites of the NiAs-type hexagonal structure by the Al atoms.

I. INTRODUCTION

High-density magneto-optical storage technology requires the recording medium have a large polar Kerr rotation. Currently, the most promising magneto-optical materials, rare-earth–transition-metal amorphous thin films and Pt/Co multilayer films, only have a polar Kerr rotation of 0.1°–0.4°.1,2 Although there are methods to enhance the Kerr rotation, such as dielectric antireflection coatings3 and optimization of the optical properties of the substrate,4 it is still of great interest to look for a material that possesses a large intrinsic Kerr rotation and good related physical properties.

So far, the materials with the largest Kerr rotations are certain U-based compounds,5 PtMnSb,6 and MnBi.7 The Curie temperatures of the U compounds are mostly well below room temperature, while PtMnSb has very weak or no perpendicular anisotropy, so that these materials are not suitable for practical application. On the other hand, MnBi has a large Kerr rotation and very strong perpendicular anisotropy.7 Unfortunately, it is polycrystalline and has a first-order phase transition near its Curie temperature. In order to overcome these disadvantages, considerable work has already been done on MnBi films modified by doping with, for example, Ti,8 Cu,9 and Dy.10 In some of these experiments, doping improved the thermal stability and some reduced the Curie temperature, but at the same time it reduced the Faraday or Kerr rotation as well.

Recently, Wang studied the effects of (Si, Al) doping on the MnBi thin films.11 He found that (Al, Si) doping increased the Kerr rotation, improved the thermal stability, and reduced the polycrystalline grain size, all important considerations for magneto-optical recording. In this paper, we will discuss the effect of Al on the magneto-optical properties and perpendicular anisotropy of MnBiAl thin films.

II. EXPERIMENT

Thin films of MnBi0.8Al0.2 (0 < x < 1.0) were made by sequential evaporation of Al, Bi, and Mn from a tungsten boat onto a room-temperature glass substrate. The base pressure of the vacuum chamber was 2 × 10⁻⁷ Torr, and was 3 × 10⁻⁶ Torr during evaporation. The film thickness was typically 800 Å. After evaporation, the thin films were covered with a SiO antireflection coating of thickness 1500 Å (suitable for 632.8 nm). All of the samples were subsequently annealed in vacuum at about 300 °C for at least 4 h. X-ray-diffraction measurements confirmed that all the samples with x < 1.0 retained the NiAs-type hexagonal structure of pure MnBi, with the c axis being highly oriented along the film normal direction. The lattice parameter is the same as that of MnBi within our measurement error for all the samples. As x was increased above 1.0, the hexagonal structure gradually disappeared and the films became amorphous. The Kerr rotation was measured using apparatus described elsewhere.12 The temperature dependencies of the properties were measured using a dewar that can be used over the temperature range 80 to 600 K. The maximum magnetic field in this measurement was 8 kOe.

III. RESULTS AND DISCUSSION

Figure 1 shows the Al concentration dependence of the Kerr rotation at room temperature. The Kerr rotation first decreases with increasing Al concentration, then increases for further Al doping. At about x = 0.9 the Kerr rotation reaches 3.15°, the same value as that of MnBi.8 The thickness of the SiO overcoat is the same for all the samples, so this phenomenon is not due to the coating. Figures 2 and 3 show the saturation Kerr rotation and absolute reflectivity wavelength dependencies of MnBi0.8, MnBi0.8Al0.2, and MnBi0.8Al0.8, respectively. The reflectivity oscillates with wavelength because of interference in the SiO overcoating. Besides the oscillation, there is little structure between 300 and 800 nm. The Kerr rotation has a broad peak around 640 nm for all the samples, due mostly to the enhancement caused by the antireflection coating. The wavelength dependencies of the Kerr rotation of MnBi0.8 and MnBiAl have similar shapes, suggesting that the Al doping does not
gives rise to any additional transitions in the visible and near-infrared regions. Because of the overcoating, it is difficult to determine the intrinsic Kerr rotation for the MnBiAl films, but the intrinsic saturation Kerr rotation of the MnBi at room temperature is about 0.7°, so it is reasonable to estimate that the intrinsic Kerr rotation for MnBi0.8Al0.2 is of similar magnitude. There is no Kerr rotation enhancement due to the Al doping while such enhancement was found in the (Al, Si) doping of MnBi thin films. The difference may be due to the Si content in the earlier films, since in our MnBiAl films the overcoating is SiO2, which likely bonds tightly enough to avoid diffusing into the thin films. From the band structure of MnBi calculated by Coehoorn and de Groot,13 we can see that a little shift of the Fermi level due to Al doping in the MnBiAl system will not affect the interband transitions very much. In other words, we do not expect a large change of the Kerr rotation through Al doping, at least in a rigid-band model. Although there is no Kerr rotation enhancement, the Kerr rotation of the MnBiAl system still is larger than that of the rare-earth–transition-metal amorphous thin films.

Another purpose of doping MnBi films with Al is to reduce the polycrystalline grain size and to study how the modified microstructure affects the perpendicular anisotropy and the coercivity. Figure 4 shows Kerr hysteresis loops at 85 K for MnBi0.8 and MnBi0.8Al0.2. From the figure we can see that at that temperature, MnBi0.8 no longer has perpendicular anisotropy, in agreement with the results of Chen and Gondo.7 However MnBi0.8Al0.2 maintains its perpendicular anisotropy down to at least 85 K, as is evidenced by the square hysteresis loop. Figure 5 shows the temperature dependence of the saturation and remanent Kerr rotations for MnBi0.8, MnBi0.8Al0.2, and MnBi0.8Al0.8. The saturation Kerr rotation of MnBi0.8 increases with decreasing temperature a little faster than those of MnBi0.8Al0.2 and MnBi0.8Al0.8, but the behaviors are qualitatively the same. We do not find such a dramatic increase (to 4.5° at 80 K) of the saturation Kerr rotation when the temperature lowered as was reported in Ref. 7. However our results are in agreement with the results of the Faraday rotation measurements of Ref. 14. The increase in Kerr rotation as the temperature decreases can be attributed mostly to the increase of the saturation magnetization. The remanent Kerr rotation temperature dependence has very different features for MnBi and MnBiAl films. For the MnBiAl films, from 600 to 80 K, the hysteresis loops are always perfectly square, meaning there is no deviation of the magnetic moment from the c axis. Whereas at T = 200 K, the remanent Kerr rotation of
MnBi gradually changes from along the c axis to the basal plane. This result also was found in other experiments. Therefore at low temperature, the pure MnBi films have no perpendicular anisotropy.

Figure 6 shows the temperature dependence of the coercivity for MnBi, MnBiAl, and MnBiSb. The coercivity increases dramatically with an increase in the Al concentration. The temperature dependence of the coercivity is similar for all the samples. As the temperature is increased from 85 K, the coercivity increases with increasing temperature, presumably because of the increase in the magnetocrystalline perpendicular anisotropy. The coercivity reaches a maximum at about 450 K and then decreases at higher temperature as the Curie temperature is approached.

MnBi, MnBiAl, and the related system MnSb all have the NiAs-type hexagonal structure. MnBi and MnSb flip their magnetic moment from along the c axis into the basal plane at 142 and 510 K, respectively. The perpendicular anisotropy in these systems presumably results from magnetocrystalline anisotropy, so that the anisotropy is closely associated with the crystal field. When the temperature decreases, the lattice will contract, which should cause a change of the local crystal field and therefore the anisotropy. When MnBi is doped with Al, some Al atoms occupy interstitial sites of the hexagonal structure, and these atoms may block the contraction of the lattice. This may be the reason why in MnBiAl thin films there is no magnetic moment reorientation down to 80 K. To test these speculations, more experimental studies are needed.

In summary, the temperature dependence of the magneto-optical and perpendicular anisotropy properties of the MnBiAl system have been investigated. Suitable Al doping does not affect the intrinsic Kerr rotation, but it changes the microstructure and anisotropy of the MnBi film. Between 80 and 600 K, thin films of MnBiAl are thermally stable and have a large Kerr rotation. Thus MnBiAl may be a good candidate for magneto-optical recording applications.

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