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Magnetic and spectroscopic characteristics of ZnMnO system

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Abstract: We report on the observation of room-temperature ferromagnetism in epitaxial (Zn,Mn)O films grown by a pulsed-laser deposition technique using high-density targets. The X-ray, microscopic, spectroscopic and magnetic properties of target material containing 6 at.% of Mn and films were compared. The target shows the presence of large clusters exhibiting paramagnetic behavior. However, ferromagnetic properties were observed in (Zn,Mn)O films grown at a substrate temperature of 500 °C and with an oxygen partial pressure of 1 mTorr. Although, crystalline quality of the film improves with increasing substrate temperature, the ferromagnetism becomes weaker.

The observation of ferromagnetic transition in (III, Mn)-V semiconductors [1], [2], [3], [4] and [5], such as GaMnAs, at higher temperatures (having Curie temperature >100 K) has attracted a lot of attention recently. The possibility of manipulating both charge and spin of electrons in semiconductors, its impact on information storage technology using ferromagnetic metals, and the recent demonstration of long spin-coherence time [6] have heightened research activities in dilute magnetic semiconductors

(DMS), including DMS oxides [7]. Following the theoretical prediction [8] that transition metals, especially Mn, doped with GaN and ZnO could show large ferromagnetic Curie temperature, numerous studies have been carried out on (Zn,Mn)O and (Ga,Mn)N systems. The recent discovery of ferromagnetism [4] and [9] in (Ga,Mn)N at temperatures much higher than the room temperature has initiated research interest that these materials can indeed have profound technological impact.

Recently, ferromagnetism in bulk, nanostructures and Mn ion-implanted ZnO films has been observed [10], [11], [12] and [13]. The reported ferromagnetic transition temperature varies from 50 to 300 K. On the other hand, ZnMnO films prepared by magnetron sputtering [14], pulsed laser deposition [15] and polycrystalline samples [16] did not show ferromagnetic behavior. Recently, ferromagnetism has been reported [17] in both ZnMnO bulk and thin film with ferromagnetic Curie temperature $T_c > 420$ K. This has aroused intense interest in this wide band gap semiconductor for possible spintronic applications. There are large controversies and difference in results in this system, which is attributed to the difference in preparation techniques. Apart from this, the metastable behavior of ferromagnetism, which not related to the carrier-mediated phenomenon [18], raised the questions on real origin of ferromagnetism in ZnMnO system. In this paper, we demonstrate the observation of roomtemperature ferromagnetism in epitaxial (Zn,Mn)O films grown by a pulsed-laser deposition technique using high-density targets. However, the target containing 6 at.% of Mn shows the presence of large clusters exhibiting paramagnetic behavior. The ferromagnetic properties were observed in (Zn,Mn)O films grown at an optimum substrate temperature of 500 °C and with an oxygen partial pressure of 1 mTorr. The ferromagnetism becomes weaker for films grown at higher substrate temperature despite higher crystalline quality of the film.

The $Zn_{1-r}Mn_rO$ (x = 0.01–0.07) precursors were prepared by mixing ZnO and MnO₂ (purity 99.999%) powders stoichiometrically and calcined at 400 °C for 8 h. The calcined powders were ground, palletized and isostatically pressed at 400 MPa in order to get a highly dense target. The target was sintered in air at 500 °C for 12-20 h. ZnMnO/Sapphire (0001) epitaxial films were grown by the pulsed-laser deposition (PLD) technique (KrF excimer, $\lambda = 248$ nm, laser repetition rate of 5 Hz) with a pulse energy density of 1-2 J/cm² and utilizing both target and substrate rotation facilities. High-density Zn_{0.94}Mn_{0.06}O target was used. The films were deposited with a substrate temperature $T_s = 500-600$ °C, keeping oxygen par-tial pressure $P_{O2} = 0.1-10$ mTorr. Clean single crystalline sapphire substrates were loaded to the chamber and heated just after the ultimate base pressure $<4 \times 10^{-8}$ Torr is reached. Phase identification was

performed by the X-ray powder diffraction (XRD) method on a Rigaku X-ray diffractometer (XRD) using Cu K α radiation. The Raman spectra were recorded using Infinity LabRam micro-Raman spectrometer built around an Olympus Microscope BH-2 and with He–Ne laser excitation (wavelength: 632.8 nm). The magnetization was measured using Quantum Design Superconducting Quantum Interference Device (SQUID-MPMS). The EPR experiments were done in an EMX Bruker spectrometer.

Fig. 1 shows the XRD patterns of ZnMnO bulk and thin film samples. The main XRD peaks for the bulk can be indexed to a hexagonal wurtzite ZnO structure. The XRD lines for ZnMnO bulk did not show any significant shifts compared to ZnO. Three prominent additional lines are seen for ZnMnO bulk.



Fig. 1. X-ray diffraction pattern of ZnMnO (a) bulk with Mn = 0 and 6 at.%, (b) film deposited at the substrate temperature of 500 and 600 °C. The inset in (b) shows the rocking curve for the film grown at a substrate temperature of 500 °C.

These peaks are arising from MnO₂ and can be indexed as shown in Fig. 1. The indexing corresponds to synthetic MnO₂ (pyrolusite), which is very stable with tetragonal rutile structure. It is also noted that similar XRD lines arise when the annealing temperature is increased for the lower concentration of Mn (not shown here). On the contrary, the XRD patterns of ZnMnO/Sapphire (0 0 0 1) films with a substrate temperature $T_s = 500$ and 600 °C reveal only one strong orientation $(0\ 0\ 2)$, illustrating the epitaxial nature of the film. The rocking curve for the epitaxial growth of the film is shown in the inset of Fig. 1. The full-width half-maxima (FWHM) calculated from X-ray (002) line broadening shows that FWHM is 0.4°. It is noted that the FWHM decreases from 0.4 to 0.19° with increasing T_s from 500 to 600 °C at an oxygen partial pressure (\vec{P}_{O2}) of 1 mTorr, illustrating the higher crystalline quality of the film grown at higher temperature. The XRD pattern also show that the $(0\ 0\ 2)$ peak of film for $T_s = 500$ °C shifts to lower angle compared to that of films grown at $T_s = 600$ °C. This clearly shows that Mn is incorporated into Zn lattice when films are synthesized at low T_s value. However, it is not clear yet that the broad peak observed at $T_s = 500$ °C as shown in the inset of Fig. 1 is the phenomenon of phase segregation due to ZnO (0 0 2) and MnO related phases. However, there is a possibility that the XRD peak broadening in films may be due the consequence of overlapping of two separate peaks arising from ZnO and MnZnO minority phase, whose presence is recently argued [19]. More studies are underway to explore this fact.

The scanning electron microscopic image (SEM) of the target surface is shown in Fig. 2(a). The SEM image shows that the larger clusters (bright spots) are embedded in the matrix, despite 12 h annealing at 500 °C. The atomic force microscopy (AFM) images of as-grown ZnMnO films are shown in Fig. 2(b) and (c). The remarkable improvement of the crystalline quality and surface morphology is very clear with the increasing T_s . The crystalline perfection of the surface is very consistent with the XRD results. The grain size increases from about 25–35 nm with increasing T_s from 500 to 600 °C, resulting in high-quality epitaxial film grown at higher T_s . The grains become very uniform and even coalesce in films grown at higher temperature.

In Fig. 3, we show the Raman spectra of ZnMnO for the target and films. ZnO belongs to the wurzite



Fig. 2. The (a) SEM image of ZnMnO bulk, and AFM images of as-grown ZnMnO films grown at the substrate temperature of (b) 500 $^{\circ}$ C and (c) 600 $^{\circ}$ C.

per primitive cell where all the atoms occupy $C_{3\nu}$ sites. Eight sets of zone center optical phonons [20], [21] and [22], $2A_1 + 2E_2 + 2E_1 + 2B_1$, are predicted according to the group theory analysis. The A_1 and



Fig. 3. Raman spectra of ZnMnO (a) bulk in the cluster region using 11 mW laser power, (b) in the normal region using 1.1 mW laser power, (c) film prepared at $T_s = 500$ °C, (d) film prepared at $T_s = 600$ °C, and (e) of sapphire substrate.

 E_1 modes are polar and split into transverse optical (TO) and longitudinal optical (LO) phonons, and E_{2} mode consists of two modes of low and high frequency phonons. On the other hand, the B_1 modes are not Raman active. To avoid sample photo transformation Raman spectra in the cluster region were recorded using low excitation power (1.1 mW). All normal region spectra in both bulk and films were recorded with 11 mW of power. We note that the beam spot size for the spectra at both normal and cluster regions was limited to only $<2 \mu m$. The normal and the cluster regions in bulk are clearly distinguishable by the microscope attached to Raman spectrometer. It is noted that giant clusters of $1 \, \mu m$ to several µm are seen in scanning electron microscopes. The cluster regions were very bright compared to the normal regions. In the normal region, the most intense peak at 437 cm⁻¹ corresponds to the vibrational mode of E_2^{high} , and it is a typical Raman peak of ZnO bulk. The additional low intensity peaks observed in Fig. 3 are assigned to their respective modes. The modes at 203, 333 and 664, and above 1000 cm⁻¹ are due to the multi-phonon

scattering process. The E_2 phonon mode centered around 437 cm⁻¹ is obviously a good choice in order to understand the stress-induced phenomena in wurzite ZnO. An increase in the E_2 phonon frequency is generally ascribed to compressive stress, where as a decrease is caused by the tensile stress. However, the E_2 phonon frequency observed at 437 cm⁻¹ did not show any significant change in the Raman shift. In addition, we did not observe any extra peaks related to Mn, suggesting that Mn is incorporated into Zn. The most remarkable feature in Raman spectra is the improvement in the crystalline quality with increasing T_s . This is clearly seen from the comparison of the spectrum (c) and (d), in which the $E_1(LO)$ mode in (d) becomes very broad.

In addition, the spectra in the cluster region of the target are significantly different compared to the spectra in the normal region. Surprisingly, the most intense E_2 phonon frequency at 437 cm⁻¹ as discussed earlier disappeared completely. Especially, in the higher frequency region, two remarkable changes happened: (1) two new peaks emerged at about 675 and 765 cm^{-1} , and (2) the multiphonon contribution observed in ZnO and for ZnMnO samples at normal spot completely disappeared. The Raman spectra of MnO₂ show a relatively sharp peak at ~765 cm⁻¹ (v_6 vibrational mode)¹⁸ indicative of a well-developed rutile-type MnO₂ structure with an interstitial space consisting of narrow 1D channels. It is generally characterized by the presence of one sharp peak at 665 cm⁻¹ along with four other bands at 319, 377, 486 and 750 cm^{-1} , those correspond to the skeletal vibrations. The bands at lower frequency region, such as 319, 377 and 486 cm⁻¹ in MnO₂ overlap with those of ZnO and could not be clearly resolved. The main contributions are attributed to the stretching mode of the Mn–O band in MnO₆ octahedra at higher frequency and to the deformation modes of the metal oxygen chain of Mn–O–Mn in the MnO₂ octahedra lattice at lower frequencies. However, the observation of giant clusters of MnO₂ in ZnO matrix is complex and there are possibilities that some of these bands are diluted or totally masked depending on the cluster size, configuration and distribution.

The temperature dependence of the EPR spectra is shown in Fig. 4 for the target. The nature of EPR spectra did not show any significant change of slope for T = 77, 240 and 300 K in the spectral characteristics. However, both slope and the nature



Fig. 4. Paramagnetic resonance spectra of 6 at.% Mn doping annealed at 500 °C for 12 h are shown for different temperatures.

of the EPR curves changes for T = 105 and 181 K. Surprisingly, the EPR spectra show a prominent paramagnetic behavior at 400 K. This is a contrasting behavior to the generally observed EPR signal for paramagnetic samples where temperature is varied. We would expect a diminished EPR signal at higher temperatures. Although the clusters are inhomogeneous and contribute to the paramagnetic signals observed in our experiments, the anomalous temperature dependence of EPR signal in ZnMnO could not be explained and needs further studies. However, EPR results clearly suggest that no ferromagnetism is present in ZnMnO bulk material. The EPR measurements were measured on the film synthesized from the same target. However, EPR did not show any significant signal, probably limited by the sensitivity of the instrument.

In Fig. 4(a), we present the magnetic field dependence of the magnetization of ZnMnO target and films at 300 K. The bulk target shows pronounced paramagnetic behavior. Fig. 4(b) shows the magnetic field dependence of magnetization curves of films at 300 K. The film grown at $T_s = 500$ °C exhibits a pronounced ferromagnetic behavior. The field at which the maximum in magnetization, H_m (low field to high field) is achieved decreases from 980 G at 5 K to 780 G at 300 K. The inset in Fig. 4(b) shows the temperature dependence of fieldcooled magnetization on the above film in an applied magnetic field of 100 G. The curve clearly illustrates that the Curie temperature is above 350 K. The rapid rise of magnetization below 100 K may be attributed due to the presence of small Mn clusters, which become dominating factor for enhanced paramagnetism. However, the room temperature ferromagnetic hysteresis shrinks with increasing T_s (for $T_s = 600$ °C), illustrating probably a dominant competition between ferromagnetic and anti-ferromagnetic states (Fig. 5).

In conclusion, we have demonstrated that the bulk material synthesized at 500 °C containing 6 at.% of



Fig. 5. The magnetization hysteresis loops of (a) ZnMnO bulk at 300 K, (b) films for two growth temperatures. The inset in (b) shows the temperature dependence of field-cooled magnetization of film grown at a substrate temperature of 500 $^{\circ}$ C.

Mn in ZnMnO displays paramagnetic behavior due to the presence of large Mn-related clusters. However, the epitaxial ZnMnO films grown by a pulsedlaser deposition technique using the same target shows pronounced ferromagnetism at room temperature when synthesized at a substrate temperature of 500 °C. The ferromagnetic properties become very week with increasing substrate temperature although the crystalline quality of the film improves. This contrasting behavior indicates that the Mn clustering, which is responsible for paramagnetic behavior, occurs at higher substrate temperature.

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