Role of swift heavy ion irradiation on the structural and magnetic properties of Ti0.95Co0.05O2−*δ* epitaxial thin films

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**Abstract**. Defects such as oxygen vacancies are found to play a vital role in determining the physical and magnetic properties of Ti1−*x*Co*x*O2−*δ*. These defects can be created during growth or may be induced by ion irradiation or implantation. In this work, the structural and magnetic properties of epitaxial Ti0.95Co0.05O2−*δ*thin films deposited on LaAlO3 single crystal substrates under dense electronic excitation are discussed. Films were deposited by pulsed laser deposition (PLD) technique. X-ray diffraction (XRD) studies revealed that reflections, beside those corresponding to the planes (004) and (008) of the anatase phase of TiO2, are suppressed, indicating epitaxial growth of the films along the *c*-axis. In order to investigate the important role of defects on the physical properties of the thin films, these were irradiated with 100 MeV Ag7+ ions with fluences 1×1011, 1×1012 and 1×1013 ions.cm-2. XRD of the irradiated films indicate successive amorphization of the films with increasing ion dose. The magnetic measurements indicate a significant enhancement in the magnetization of the film irradiated with a fluence of 1×1013 ions.cm-2. This unexpected increase in magnetization is explained on the basis of the bound magnetic polaron (BMP) model. The findings suggest the pivotal role of ion irradiation on tailoring the structural as well as the magnetic properties.

1. Introduction

Semiconductor spintronics unites both logic-processing functionalities (electronic charge) of a semiconductor and information-storage functionalities (electronic spin) of a ferromagnetic material. The main obstacle to realize such spintronics devices are the low Curie temperature (*T*c) and controversial origin of magnetism in these systems [1].

In oxide based systems, room temperature ferromagnetism (RTFM) was first discovered by Matsumato *et al.* [2] in Co doped TiO2 thin films making it a potential diluted magnetic semiconductor (DMS) material. Several models or mechanisms have been used to explain the origin of ferromagnetism in these systems, including the bound magnetic polaron (BMP) model, Ruderman-Kittel-Kasuya-Yosida (RKKY) exchange interaction, Stoner type model and charge transfer mechanisms [3, 4]. All these models or mechanisms could not explain the origin of magnetism satisfactorily. Although the exact mechanism of the magnetic correlation is still unclear, the possible coupling between charge carriers created by defects like oxygen vacancies and magnetic moment is unanimously accepted [5]. However, clustering of Co in the TiO2 matrix resulting in ferromagnetic order is not accepted [6]. Ferromagnetism in Co doped TiO2 polycrystalline thin films deposited by pulsed laser deposition (PLD) technique refuting the presence of Co clusters in the films has also been reported [7]. Films deposited under low oxygen partial pressure or annealing of the films under reduced atmosphere are found to demonstrate ferromagnetic behaviour at room temperature. Defects like oxygen vacancies also greatly influence the ferromagnetic properties of undoped TiO2 thin films [8]. The challenge is therefore to create defects in the material systematically.

Ion irradiation used for defect engineering has been employed as a potential tool in material science [9]. Irradiation may dissolve the magnetic clusters in the host lattice and create disorder like amorphization in the system depending on the energy of the incident ion beam [9]. It can also create change in the crystal structure, enhance phase formation and defect annihilation with suitable selection of projectile ion, energy and fluence [10]. The material modiﬁcation can be complex depending upon the magnitude of electronic energy loss (*S*e) as compared with a material-dependent threshold value, *S*eth, beyond which swift heavy ions (SHI) can create amorphized latent tracks or can induce crystalline-to-crystalline phase transition [11, 12]. Recently, the role of SHI on structural and magnetic properties of Co doped TiO2 polycrystalline thin films deposited on Si substrates have been reported [13].

To investigate the effect of ion irradiation on epitaxial thin films of Co doped TiO2, this study utilized *c*-axis oriented LaAlO3 (LAO) as the substrate, because of the small lattice mismatch it has with TiO2 [14]. Secondly, owing to the important role of defects in tuning the magnetic property, the thin films were irradiated with 100 MeV Ag7+ ions with fluences 1×1011, 1×1012 and 1×1013 ions.cm-2. The present work focuses on the modification in structural and magnetic properties of Ti0.95Co0.05O2−*δ*thin films with the use of swift heavy ion irradiation.

2. Experimental

Co doped TiO2 powdered samples were used to prepare the target for the pulsed laser deposition (PLD) by pressing the samples together in pellets and sintering these at 900 oC for 24 hours. The target was ablated using a *KrF* excimer laser (Lambda Physik COMPex 201 Model, Germany) at constant laser energy of 240 mJ and 10 Hz repetition rate. Commercial LaAlO3 (LAO) single crystal substrates were used for film deposition. The substrate temperature was maintained at 700 oC during film deposition. Films were fabricated at 10 mTorr oxygen partial pressure. The target was rotated continuously during laser ablation. The deposited films were irradiated with 100 MeV Ag7+ ions at room temperature using a 15 UD tandem pelletron accelerator at IUAC, New Delhi, India with fluences 1×1011, 1×1012, and 1×1013 ions.cm-2. The magnetic measurements of all the samples were investigated using a superconducting quantum interference device (SQUID) vibrating sample magnetometer (VSM) from Quantum Design in a magnetic field up to 1 T and temperature ranging from 5 to 300 K. Structural characterization was performed using a D8 Advance x-ray diffractometer from Bruker with LynxEye 1D-PSD detector in *θ*-2*θ* geometry. Film thickness and elemental compositions were measured with Rutherford’s backscattering spectrometry using stream of *α* particles (He2+). The measured thickness of the pristine film was calculated to be ~ 100 nm.

3. Results and discussion

The XRD pattern of the as deposited thin film indicates the anatase phase of TiO2, as seen in Fig. 1. All the reflections except (004) and (008) are suppressed due to the epitaxial growth of the film along *c*-axis. After 100 MeV Ag7+ ion irradiation with fluences 1×1011, 1×1012 and 1×1013 ions.cm-2, successive amorphization is clearly seen from the XRD profile shown in Fig. 1.

The interaction of ion beams with material modifies its physical properties. Ion beam irradiation is found to induce defects, anneal pre-existing defects, induce structural phase transformation and also influence the crystallinity of the target material [10]. SHI passes through the surface with velocity comparable to an electron’s Bohr velocity and loses its energy while traversing through matter. The

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| **Figure 1.** XRD pattern of pristine and irradiated Ti0.95Co0.05O2−*δ* thin films deposited at 10 mTorr oxygen partial pressure on LaAlO3 substrate. The fluence used is indicated in each panel. ‘S’ denotes the reflections related to LaAlO3 substrate. |

total energy loss can be expressed as the contribution of two independent energy loss processes: (i) electronic energy loss, (dE/dx)e ~ *Se*, attributed to electronic excitation and ionization due to the inelastic collision with electrons dominating at high energy regime and (ii) nuclear energy loss, (dE/dx)n ~*Sn*, due to the elastic collisions with the atoms of the solid with the nuclei of the projectile ion. The former process is dominant for ions in the mega electron volt (MeV) energy range and the latter is valid for ions in the kilo electron volt (keV) energy range. In the present study, the energy of the projectile ions is selected such that it can pass through the film thickness and get buried inside the substrate. The 100 MeV Ag7+ ions deposit ~ 18.16 keV.nm-1 in inelastic collisions (electronic energy loss *Se*) with electrons, ~ 90.86 eV/nm in elastic collisions (nuclear energy loss *Sn*) with nuclei as estimated using SRIM (The Stopping and Range of Ions in Matter) simulations code [15]. The projected range of the ions is approximately 9650 Å which is greater than the film thickness i.e. ~ 1000 Å. Since the threshold for track formation in TiO2 film is approximately 6.2 keV.nm-1, it is expected that the changes produced in the films are dominantly due to the electronic energy loss [16]. From the XRD it is clear that the film gets completely amorphized at a fluence of 1×1013 ions.cm-2, as shown in Fig. 1.

In order to investigate the magnetic properties of the pristine and irradiated films, magnetization measurements as a function of applied magnetic field under zero filed cooling (ZFC) and field cooling (FC) conditions, with a probing field of 50 Oe and temperature ranging from 5 to 300 K, were performed. These results are shown in Fig. 2. As Co is a magnetic metal and can contribute to the magnetization of the film in the form of Co clusters instead of substituting the Ti site, it is necessary to determine the Co concentration in TiO2. This was performed by using Rutherford’s backscattering

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| **Figure 2.** Magnetization as a function of temperature of pristine and irradiated Ti0.95Co0.05O2−*δ* thin films deposited at 10 mTorr oxygen partial pressure on LaAlO3 substrate. |

spectrometry (RBS). RBS measurements indicated that the actual Co concentration is 5 atomic %. In order to form Co clusters in TiO2 matrix, a Co concentration of greater than 7 atomic % is needed [2]. In the measured films the Co concentration is less than the critical concentration and thus the contribution to magnetization because of Co cluster is ruled out. From Fig. 2 it is clear that the magnetization relative to ZFC and FC measurements are irreversible, as the two measurements do not coincide. From Fig. 2 it is also evident that the *T*c of the films is above 300 K. Fig. 3 illustrates the magnetization as a function of applied magnetic field at 300 K and 5 K for the pristine as well as irradiated thin films. From the obtained magnetization values the diamagnetic contribution from the LAO substrate was carefully deducted. The magnetization increases with increasing magnetic field and almost saturates. While decreasing the field, the magnetization did not retrace the same path and show a prominent hysteresis for the pristine as well as irradiated thin films. The saturation magnetization (*M*s) increases for the fluence 1×1011 ions.cm-2. However, it decreases with fluence 1×1012 ions.cm-2. Surprisingly, the *Ms* increased abruptly for the film irradiated with 1×1013 ions.cm-2 that shows complete amorphization at this fluence. A similar trend is obtained in *M-H* behaviour measured at 5 K (see Fig. 3 (b). Thakur *et al.* [17] reported ferromagnetism in undoped TiO2 when irradiated with 200 MeV Ag ions with a phase change from anatase to brookite. Sanyal *et al.* [18] reported ferromagnetism with low energy Ar irradiated TiO2 thin films. In case of Co doped TiO2 polycrystalline thin films, the saturation magnetization *M*s decreases with increasing ion fluence [13]. The decrease in *M*s was explained as a consequence of formation of magnetically disordered region around the ion track that does not contribute to the ferromagnetic order [13]. However, in the present

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| **Figure 3.** *M*-*H* plots for the pristine and irradiated Ti0.95Co0.05O2−*δ* thin films deposited at 10 mTorr oxygen partial pressure on LaAlO3 substrate and measured at temperatures: (a) 300 K and (b) 5 K. The insets show the zoomed view of the hysteresis loops in low field region. |

case the *M*s is increased with ion irradiation which is quite different from the response of the polycrystalline films towards swift heavy ions.

There are several mechanisms proposed for the observed magnetism in Co doped TiO2 system: (i) Bound magnetic polaron percolation (BMP) model, (ii) indirect RKKY exchange interaction, and (iii) impurity band exchange model. Among these models or mechanisms BMP and RKKY models are quite convincing. Bound magnetic polarons are formed due to Coulomb and magnetic exchange interactions surrounding a defect like oxygen vacancy [4, 19]. In the BMP model carriers are localized due to electrostatic interaction with some defect [19]. The magnetic polarons are well deﬁned, non-overlapping, isolated entities, only at low carrier densities and suﬃciently large temperatures. The size of the polarons increase as the temperature decreases, eventually overlapping with neighbouring BMPs. This overlap causes the alignment of their spins, therefore forming ferromagnetic (FM) clusters. The FM transition takes place when an ‘inﬁnite cluster’ (of the size of the system) is formed, i.e. when the percolation of BMP occurs. The model is valid in the low carrier density regime and when the magnetic impurities are larger than the density of carriers [19]. In case of RKKY model, the itinerant electron plays a vital role. The itinerant electron interacts with the localized moments and tends to order them. For RKKY model to be applicable, the system should have sufficient itinerant electrons. With ion irradiation it has been found that the conductivity of TiO2 films decreases and films become insulating [20]. Thus, the mechanisms leading to ferromagnetism in these films at lower and higher fluence are definitely different. The competing effect between these two mechanisms leading to ferromagnetic ordering results in the anomaly seen in the magnetization for the intermediate fluence as observed from the *M*-*H* measurements at 5 and 300 K. Thus, bound magnetic polaron (BMP) mechanism is mostly dominant for the amorphous film irradiated with fluence 1×1013 ions.cm-2 demonstrating maximum magnetization.

4. Conclusions

Ti0.95Co0.05O2−*δ* thin films were grown on single crystal LaAlO3 substrates. The anatase phase of the film was retained with ion irradiation. However, the crystallinty degraded with ion irradiation and the films became completely amorphous at fluence 1×1013 ions.cm-2. ZFC-FC magnetization revealed prominent bifurcation below 300 K, indicating that the *T*c of the film is well above room temperature. Magnetization as a function of applied magnetic field measured at 300 K and 5 K confirms the ferromagnetic behaviour of the films. The *M*s initially increased for the film irradiated with fluence 1×1011 ions.cm-2 then decreased for the fluence 1×1012 ions.cm-2 and finally the magnetization increases dramatically for the amorphous film irradiated with fluence 1×1013 ions.cm-2. This unusual increase in the magnetization can be explained on the basis of BMP model.

Acknowledgements

PM gratefully acknowledge Dr. R. J. Choudhary, UGC-DAE Consortium for Scientific Research, Indore, India for the measurement facilities. AREP and CJS acknowledge the financial support of UJ and the NRF (grant number 93551).

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