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Bi-substitution-induced magnetic moment distribution in spinel $Bi_x Co_{2-x} MnO_4$ multiferroic

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Abstract

We report the near-edge x-ray absorption spectroscopy (NEXAFS) at the Co/Mn $L_{3,2}$ edge and oxygen K edge of the well-characterized Bi-substituted Co_2MnO_4 multiferroic samples. The evolution of peak features in NEXAFS spectra of the Co/Mn $L_{3,2}$ edge and O K edge show the Bi-induced redistribution of magnetic cations (Co/Mn). The variation in valence states of Co and Mn in all the substituted compositions is consistent with the observed ferrimagnetic behaviour of the samples. Magnetization data show the decrease in molecular field complementing the ferrimagnetism. The role of Bi in the enhancement of magnetic interactions as well as the appearance of ferroelectricity in $Bi_xCo_{2-x}MnO_4$ ($0 \le x \le 0.3$) is discussed.

(Some figures in this article are in colour only in the electronic version)

1. Introduction

Multiferroics (MF) are the materials that enfold physics for various future technological applications and have received worldwide attention from the scientific community working in the field of materials sciences [1, 2]. Coexisting magnetism and ferroelectricity along with coupling between them ensures the possibility of technological uses for these materials but it is hard to achieve the above order parameters in a single phase due to their contrasting origins. Various alternative mechanisms have been proposed as a solution to this problem, including an approach to achieving the ferromagnetism with non-metallicity or ferroelectricity in centrosymmetric systems [3, 4]. These materials are timehonoured systems owing to their extraordinary magnetic and electric properties, which appeared as the source of the most fascinating tunable magnetoelectric behaviour, by changing the chemical composition and cation redistribution [4-11].

The crystal symmetry in spinel oxides offers ample opportunities for testing/designing new functional properties with numerous permutations and combinations possible at A and B sites. The charge distribution of the normal spinel is represented by $[A^{2+}]_{8a}[B_2^{3+}]_{16d}[O_4^{2-}]_{32e}$, in which Wyckoff positions 8a denote the tetrahedral sites and 16d the octahedral sites surrounded by O^{2-} ions at 32e sites. Empty interstitial space is comprised of 16 octahedral sites (16c) and 56 tetrahedral sites (8b and 48f). In the Co-based spinel oxides the substituted trivalent metal ion occupies the octahedral sites, while cobalt ions are distributed over both octahedral (Co³⁺ ions in low spin $t_{2g}^6 e_g^0$ state) and tetrahedral sites (magnetic Co²⁺ ions in high spin $e_g^4 t_{2g}^3$ state) [12]. Recently we have reported the magnetoelectric multiferroic properties of Bi-substituted Co₂MnO₄, which crystallizes into

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a normal spinel with O_h^7 - $Fd\bar{3}m$ space group [13, 14]. In $Bi_x Co_{2-x} MnO_4$, Bi^{3+} with $6s^2$ lone pair electrons at the octahedral sites instead of magnetic ions, distorts the oxygen octahedron and consequently introduces ferroelectricity and influences the ferrimagnetic ordering of the parent system. Mixed valence states of this spinel oxide plays a key role in determining the electrical and magnetic characteristics and requires further investigations to understand the intriguing nature of spin state distribution and hybridization states. The deeper understanding of the electronic structure of these spinel oxides would be desirable to design new MF materials and understand their complex magnetic ordering, which has a profound effect on ferroelectric behaviour. It necessitates the use of powerful microscopic experimental tools like near-edge x-ray absorption fine structure (NEXAFS) and extended xray absorption fine structure (EXAFS). Owing to its simplicity and universality, the NEXAFS technique is most widely used in determining the valence states of atoms and local site symmetries in solids. Here, we report the electronic structure of multiferroic-magnetoelectric $Bi_x Co_{2-x} MnO_4$ (0 $\leq x \leq$ 0.3) samples using x-ray absorption spectroscopy (XAS). In this study, the NEXAFS spectra at the O K, Mn L_{3,2} and Co L_{3,2} edges of Bi_xCo_{2-x}MnO₄ ($0 \le x \le 0.3$) along with the reference compounds Bi2O3, CoO, LiCoO2, MnO and MnO2 are presented. Magnetization results are correlated to the cation distribution.

2. Experimental details

Polycrystalline single-phase $Bi_x Co_{2-x} MnO_4$ (0.0 $\leq x \leq 0.3$) samples were synthesized by the standard solid state reaction technique. The details of materials synthesis together with structural, magnetic, ferroelectric and coupling studies are presented elsewhere [13, 14]. The NEXAFS measurements of $Bi_x Co_{2-x} MnO_4$ (0.0 $\leq x \leq$ 0.3) along with Co₃O₄, CoO, MnO, MnO₂, and Bi₂O₃ at O K, Co L_{3,2} and Mn L_{3,2} edges were performed at the soft x-ray beamline 7B1 XAS KIST of the Pohang Light Source and some parts of the measurements were repeated at the ESRF's ID08 beamline, which uses an APPLE II type undulator giving $\sim 100\%$ linear/circular polarization. All the samples were scraped with diamond foil prior to the measurements in order to remove the surface contaminants. The spectra were collected simultaneously in both total electron yield (TEY) and total fluorescence yield (TFY) modes at room temperature, ensuring both surface and bulk sensitivities. The spectra were normalized to incident photon flux and the base pressure of the experimental chamber was better than 5×10^{-10} Torr. The Raman spectra were recorded at room temperature in backscattering configuration using a HR800 Jobin-Yvon spectrometer having a resolution of 1 cm⁻¹. An He–Ne laser (488 nm) was used as an excitation source at 9 mW power. Magnetization measurements were performed using the vibrating sample magnetometer (VSM) option of the Quantum Design (PPMS) set-up.

3. Results and discussions

Spinel oxides exhibit a variety of exciting physical properties due to the active role of the orbital degrees of freedom within

its complicated structure, which can be typically observed in the lattice and electronic response. As a matter of fact, such properties appear to have their origin in the unique electronic structure derived from the hybridized transition metal ions' (TMI) 3d and O 2p orbitals in the structural and chemical environment of a spinel oxide [15]. It is reported that the substitution of Bi3+ (non-magnetic) ion with 6s2 lone pair introduces the non-centrosymmetric charge ordering, leading to ferroelectricity in Co₂MnO₄ simultaneously increasing the net magnetic moment [13, 14]. This can be attributed to the hybridization effects of Mn/Co 3d-O 2p orbitals with Bi 6sp orbitals. Possibly due to the larger ionic radii of Bi³⁺, which may cause the redistribution of cations among the occupied 8a (tetrahedral- T_D) and 16d (octahedral- O_H) sites, and the participation of unoccupied 16 octahedral sites (16c) and 56 tetrahedral sites (8b and 48f) in a distorted environment due to Bi-induced chemical pressure.

NEXAFS spectra were assessed at the Mn/Co L3,2 and O K edges for $Bi_x Co_{2-x} MnO_4$ (x = 0.0, 0.1, 0.2 and 0.3) samples. The Mn/Co L_{3.2} edges directly proves the unoccupied Mn/Co 3d states via $2p \rightarrow 3d$ transitions, whereas the O K edges establish the unoccupied O 2p states via O 1s \rightarrow 2p dipolar transitions [16], together providing the information on the Mn/Co 3d occupancy and the effect of Bi³⁺ ion substitution on the hybridization between the O 2p and Mn/Co 3d orbitals. It is well known that the peak positions and the spectral shapes of the Co/Mn L_{3,2} edge NEXAFS spectrum depend on the local electronic structure of Co/Mn ions, and are sensitive to the crystal field symmetry due to the 2p-3d transitions. So the L edge spectrum provides information on the valence state of the Co/Mn ions [17, 18]. Figure 1 shows the normalized Co L_{3,2} edge NEXAFS spectra of $Bi_x Co_{2-x} MnO_4$ (0.0 $\leq x \leq$ 0.3) samples. The spin-orbit interaction of the Co 2p core states splits the spectrum into two broad multiplets, namely the L_3 (2p_{3/2}) and L_2 (2p_{1/2}) ~15 eV apart. Each of these two regions further splits into t_{2g} and e_g orbital features because of the crystal field effect of neighbouring ions. The intensity of these peaks is the direct measure of total unoccupied Co 3d states. In Co₂MnO₄, the Co ions distributed at A and B sites (in the AB₂O₄ structure) are in T_D (Co²⁺) and O_H (Co³⁺) symmetries. Co^{3+} at the B (O_H) site has the higher crystal field splitting energy, so all the valence electrons are in a lower orbital t_{2g} ($t_{2g}^6 e_g^0$) only. In tetrahedral symmetry, the situation is reversed; the e_g subbands lie at lower energy than that of t_{2g} . The associated lower crystal field splitting energy results in the high spin state of Co^{2+} ($e_g^4 t_{2g}^3$). The L_3 region contains two spectral features, A_1 and B_1 , at 775.8 eV and 777.2 eV, which correspond to Co^{2+} and Co^{3+} valence state, respectively. To determine the valence states of Co ions in $Bi_x Co_{2-x} MnO_4$, Co $L_{3,2}$ edges have been compared with those of CoO and LiCoO₂ spectra. For the starting composition Co_2MnO_4 (x = 0.0), the intensity ratio B_1/A_1 is almost equal to one, indicating that distribution of Co^{2+} ions at T_D and Co^{3+} ions at O_H , respectively, are equal. The B_1/A_1 ratio is found to increase with increase in Bi substitution, clearly showing that Co²⁺ content is reduced. This reveals that substitution of Bi at O_H sites enforces the relocation of Mn ions between $O_{\rm H}$ and $T_{\rm D}$ sites.



Figure 1. Normalized Co $L_{3,2}$ spectra of $Bi_x Co_{2-x} MnO_4$ (x = 0.0, 0.1, 0.2 and 0.3) along with the spectra of CoO and LiCoO₂.

Figure 2 shows the Mn $L_{3,2}$ spectra of $Bi_x Co_{2-x} MnO_4$ along with the reference compounds MnO and MnO₂. Similar to Co L_{3,2} edge spectra (figure 1) due to the spin-orbit interaction, Mn 2p core states split the spectrum into two broad multiplets, namely the L₃ $(2p_{3/2})$ and L₂ $(2p_{1/2}) \sim 11 \text{ eV}$ apart. Each of these two regions further splits into t_{2g} and e_g orbital features because of the crystal field effect among the neighbouring ions. The intensity of these peaks measures the total unoccupied Mn 3d states. These spectra show a valencespecific multiplet structure with a chemical shift due to changes in oxidation state. The inflection points of the Mn L3 edge for manganese oxides (Mn²⁺O and Mn⁴⁺O₂ shown in the plot and $Mn_2^{3+}O_3$ from [19–21]) are subsequently at higher energies, corresponding to the increasing valence states of Mn. The L₃ region in both MnO₂ and MnO contains two spectral features, which are assigned to Mn t_{2g} and Mn e_g subbands. Mn ions in MnO have a 2+ valence state with orbitals each filled by majority spin electrons, so that the feature at lower energy can be attributed to t_{2g} and the higher energy feature is attributed to eg subbands. The larger intensity of the lower energy peak is due to the fact that there are three t_{2g} orbitals and only two eg orbitals and t2g have more unoccupied states. In MnO2, Mn ions have a 4+ oxidation state with the eg subband empty. So the unoccupied part of the eg subband will contribute to the higher energy peak of the Mn L_{3,2}-edge spectrum, and is more intense. Similarly, in the case of Mn_2O_3 the L_3 edge is also dominated by e_g subbands only [19–21]. The L₃ region in Mn $L_{3,2}$ spectra of substituted $Bi_x Co_{2-x} MnO_4$ contains three



Figure 2. Normalized Mn $L_{3,2}$ spectra of $Bi_x Co_{2-x} MnO_4$ (x = 0.0, 0.1, 0.2 and 0.3) along with the spectra of MnO and MnO₂. Inset: zoomed-normalized Mn L_3 -edge region.

spectral features, A₂ (639 eV), B₂ (640.5 eV) and C₂ (642 eV), which can be directly compared with the three valence states (2+, 3+ and 4+) of Mn ions (see figure 2). Spectra of the parent composition (x = 0) match well with those to the spectra of Mn³⁺ [19–21]. However, the increase in the intensity of peak features A₂ and C₂ with Bi substitution at the O_H site reveal that more of the Mn³⁺ ions are converted into Mn²⁺/Mn⁴⁺. The zoomed-normalized Mn L₃-edge region is shown as an inset in figure 2, where increased Mn²⁺ and Mn⁴⁺ at the expense of Mn³⁺ is clearly displayed. Mn²⁺ occupies the T_D sites, complementing our observation of the reduced Co²⁺ in Co L_{3,2} spectra, whereas Mn⁴⁺ ions are in O_H sites only, as the signature of Mn⁴⁺ is similar to MnO₂. The increase in Mn²⁺ and Mn⁴⁺ at the expense of Mn³⁺ is clearly displayed in the enhancement of net magnetic moment.

Figure 3 shows the measured O K-edge NEXAFS spectra of the Bi_xCo_{2-x}MnO₄, along with the spectra of reference compounds CoO, LiCoO₂, Bi₂O₃, MnO and MnO₂. O Kedge NEXAFS spectra, which represents the orbital nature of the spectral features of the O 2p unoccupied states in the conduction bands, can be efficiently used to explore all kinds of possible hybridizations with different cations (Co, Mn and Bi). Based on the existing literature and band structure calculations [22–25], four spectral features marked by A₃-D₃ in Bi_xCo_{2-x}MnO₄ (x = 0.1) are identified. Except for the spectral feature B₃, all other features are present in the parent composition (x = 0.0). This is due to the reason



Figure 3. Normalized O K-edge spectra of $\text{Bi}_x \text{Co}_{2-x} \text{MnO}_4$ (x = 0.0, 0.1, 0.2 and 0.3) along with that of reference compounds CoO, LiCoO₂, MnO, MnO₂ and Bi₂O₃.

that B₃ originates from Bi 6s/O 2p hybridizations. The O Kedge spectra of the pure (x = 0.0) sample indicates that low energy features resemble Co_3O_4 [26] and the spectral feature B_3 at ~532 eV observed for doped compositions follows the behaviour of a highly polarizable 6s² lone pair of electrons of the Bi^{3+} ion. The orientation of the $Bi 6s^2$ lone pair towards a surrounding O^{2-} ion can produce local distortion and hybridization between the 6s² Bi orbital and O 2p orbital, resulting in ferroelectric properties in doped compounds. As a matter of fact, increase of Bi content at the O_H site induces more Mn²⁺/Mn⁴⁺ crystal symmetries, which can be clearly observed in the O K-edge spectra of an optimally doped (x =(0.3) sample, where A_3 is more prominent. On the whole, the spectral feature of peaks A_3 , B_3 , C_3 and D_3 clearly represents the presence of Bi^{3+} , $Mn^{2+/3+/4+}$ and $Co^{2+/3+}$ ions in Bisubstituted samples. The above observation is in conformity to the $L_{3,2}$ spectra of Co/Mn.

The element-specific observation discussed above complements our structural measurement XRD [13] and Raman scattering. The parent compound, Co_3O_4 , which crystallizes in the normal spinel structure with O_h^7 spectroscopic symmetry, was shown to possess five Raman-active modes, as $A_{1g} + E_g +$ $3F_{2g}$, with wavenumbers 194.4 cm⁻¹ (F_{2g}^1), 482.4 cm⁻¹ (E_g), 521.6 cm⁻¹ (F_{2g}^2), 618.4 cm⁻¹ (F_{2g}^3) and 691 cm⁻¹ (A_{1g}) [27]. The Raman mode at ~691 cm⁻¹ (A_{1g}) is attributed to the characteristics of O_H sites (CoO₆) in the O_h^7 spectroscopic symmetry and the mode at ~195 cm⁻¹ (F_{2g}^1) is attributed to the T_D



Figure 4. Raman scattering spectra collected at room temperature for $Bi_x Co_{2-x} MnO_4$ (x = 0.0, 0.1, 0.2 and 0.3).

sites (CoO₄). The broadening and shifting of Raman modes on Mn substitution into the Co_3O_4 lattice is already reported [28]. Raman spectral studies of the $Bi_x Co_{2-x} MnO_4$ (x = 0.0, 0.1, 0.2 and 0.3) samples show a shift in peak positions towards lower wavenumbers (A_{1g} shifted from 664 to 650 cm⁻¹ and F_{2g}^1 shifted to 188–183 cm⁻¹ for x = 0.0-0.3) due to increased unit cell parameters with Bi substitution (figure 4). The greater shift in the A1g mode indicates that the substituted Bi and most of the Mn cations are occupying the octahedral sites (Co/Bi/MnO₆). The shift in A_{1g} further indicates the domination of the octahedral environment, as Bi³⁺ ions with greater ionic radius occupy the octahedral sites. At the same time, the shift in the $F_{2\alpha}^1$ mode is not that more prominent, since the transition metal cations occupying the tetrahedral sites are of comparable radii. In general, the Raman spectra with a flat background are indicative of only a phonon structure present in the system, but the increased background at higher wavenumbers has been observed in large numbers of reports on superconductors [29] due to increased electronic scattering. $Bi_x Co_{2-x} MnO_4$ samples exhibited a considerable increase in the background at higher wavenumbers, which indicates that the multiferroic property of this material, which is strongly dependent on electron-phonon coupling, is evident at lower temperatures as well.

Regarding the magnetic behaviour of spinel oxides, there are mainly three types of magnetic interactions possible between the ions at A site and B sites through the intermediate oxygen ions (O^{2-}) via intrasite A–O–A (J_{AA}) and B–O–B (J_{BB}), and intersite A–O–B (J_{AB}) superexchange interactions. In Co₂MnO₄ these interaction (J_{AA} and J_{BB}) energies are negative, favouring AFM. Simultaneously a positive, J_{AB} , superexchange interaction results in ferrimagnetism. In addition, with Bi substitution, a weak, J_{AA} , superexchange interaction between A sites (Co²⁺–Co²⁺) mediated through O^{2–} and Co³⁺ that maintains the AFM ordering is affected due to the redistribution of Co/Mn ions. The ferrimagnetic behaviour appears to evolve due to the Bi-substitution-



Figure 5. $1/\chi_M - T$ plots Bi_xCo_{2-x}MnO₄ (x = 0.0 and 0.3). Inset: M-H loops Bi_xCo_{2-x}MnO₄ (x = 0.0 and 0.3) measured at 150 K.

induced competition among J_{AB} (Co²⁺–O–Mn⁴⁺, Co²⁺– O–Mn³⁺, Mn²⁺–O–Mn³⁺, Mn²⁺–O–Mn⁴⁺), J_{AA} (Co²⁺– O–Co²⁺, Mn²⁺–O–Mn²⁺, Co²⁺–O–Mn²⁺), J_{BB} (Mn³⁺–O– Mn³⁺, Mn⁴⁺–O–Mn⁴⁺ and Mn³⁺–O–Mn⁴⁺) superexchange interactions and consequent magnetic frustration [14, 30]. The improvement in the ferrimagnetic property was confirmed through the rise in the ratio T_c/θ_{CW} [31], where θ_{CW} is the Curie-Weiss constant (which changes from 0.364 to 0.502 for x = 0.0-0.3 as observed from $1/\chi_M$ versus T plots, figure 5), which represents an increase in the frustration among the AFM ordering in the system, as Bi content increases. In other words, this can be interpreted by directly looking at the fall in negative Curie–Weiss constant, θ_{CW} (-494 to -370, for x = 0.0-0.3; obtained as the intersecting point of extrapolation of the high temperature region of $1/\chi_M$ versus T plots), which represent the fall in negative molecular field and consequent rise in AFM frustration, finally leading to improved ferrimagnetic behaviour. The increase in net magnetic moment because of the evolution of Mn²⁺ and Mn^{4+} (at the expense of Mn^{3+}) due to the Bi substitution is evident in M-H plots (inset of figure 5). However, it has to be mentioned that the well-defined ferrimagnetic behaviour exhibited by these compositions cannot be explained completely by the AFM frustration discussed so far. The canting of the spins by the Bi-induced structural distortion also contributes to ferrimagnetism. Therefore the source of increasing ferrimagnetic ordering with the Bi substitution in $Bi_x Co_{2-x} MnO_4$ (0.0 $\leq x \leq 0.3$) originates from the complex magnetic ordering attained and consequent enhancement in magnetic interactions. The NEXAFS study substantiates that the complex magnetic ordering is induced in the Bi-substituted spinel Co₂MnO₄.

4. Conclusions

In conclusion, we have investigated the electronic structure of $\text{Bi}_x \text{Co}_{2-x} \text{MnO}_4$ (0.0 $\leq x \leq$ 0.3) using NEXAFS spectra at

the O K edge and $L_{3,2}$ edge of Co/Mn. It has been shown that an increase in net magnetic moment and more ordered ferrimagnetism is due to Bi-induced cationic redistribution. More of the Mn³⁺ ions are converted into Mn²⁺ and Mn⁴⁺ and redistributed to T_D and O_H sites. Raman spectra of the samples confirm the greater occupation of substituted cations in the octahedral sites. A lowered negative molecular field is depicted by the decrease in Curie–Weiss constant, which indicates the increased AFM frustration with Bi substitution, supporting ferrimagnetic behaviour. O K edge spectra confirm the highly polarizable behaviour of the Bi 6s² lone pair of electrons. Finally, Bi substitution introduces the radical change in cationic distribution and canting of the spins, favouring the ferrimagnetism along with the ferroelectricity.

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