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Permalink
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Journal
The Journal of chemical physics, 152(3)

ISSN
0021-9606

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Publication Date
2020

DOI
10.1063/1.5134653

Peer reviewed
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Cite as: J. Chem. Phys. 152, 034704 (2020); doi: 10.1063/1.5134653
Submitted: 31 October 2019 • Accepted: 23 December 2019 • Published Online: 16 January 2020

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Note: This article is part of the JCP Special Topic on Oxide Chemistry and Catalysis.
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ABSTRACT
The correlation between the structural phase transition (SPT) and oxygen vacancy in SrRuO$_3$ (SRO) thin films was investigated by in situ X-ray diffraction (XRD) and ambient pressure X-ray photoelectron spectroscopy (AP-XPS). In situ XRD shows that the SPT occurs from a monoclinic SRO phase to a tetragonal SRO phase near $\sim$200 °C, regardless of the pressure environment. On the other hand, significant core level shifts in both the Ru and Sr photoemission spectra are found under ultrahigh vacuum, but not under the oxygen pressure environment. The directions and behavior of the core level shift of Ru and Sr are attributed to the formation of oxygen vacancy across the SPT temperature, indicating the close relationship between oxygen vacancy and SPT in SRO thin films.

INTRODUCTION
Perovskite materials with the ABO$_3$ structure have been extensively investigated for future applications in memory devices, energy storage devices, and electrodes for water-splitting due to their rich electronic/magnetic properties, i.e., ferroelectricity, ferromagnetism, superconductivity, etc.$^{1-12}$ In the selection of ideal perovskite materials for such applications, it is necessary to identify key physical parameters that determine the characteristics of perovskites. Among the key parameters, structural properties are known to have very close relation with diverse electronic/magnetic properties of perovskites.$^3$ In general, the structural factor in perovskites
is divided into 3 categories, (i) Jahn-Teller distortion (LaMnO$_3$), (ii) cation displacement (BaTiO$_3$, PbTiO$_3$), and (iii) octahedral tilting (SrRuO$_3$ and La$_{1-x}$Sr$_x$MnO$_3$). Especially, the octahedral tilting, the most common structural factor among perovskites, influences strongly the physical properties such as ferroelectricity, magnetism, and charge ordering. In perovskites with octahedral tilting, the B cation occupies the center of the BO$_6$ octahedron, and the distorted 6 surrounding oxygen atoms maintain the periodicity and corner connectivity relative to one or more symmetric axes.

SrRuO$_3$ (SRO) is one of the representative perovskite materials, which shows the octahedral tilting. Due to the chemical/thermal stability as well as good conductivity, SRO has been a popular choice for electrode materials of functional oxide films. In addition, the exotic transport properties of SRO have attracted much interest in basic research, e.g., superconductivity, ferromagnetism, and topological hall effect. Previously, it is reported that bulk SRO shows a structural phase transition (SPT) from the orthorhombic to tetragonal phase at 547°C, and from the tetragonal to cubic phase at 677°C. Interestingly, when prepared as a thin film (~less than 40 nm thick), SRO makes the SPT from the monoclinic to tetragonal phase at 200°C, which is significantly lower than the bulk SRO. Although a substrate-induced strain is pointed out as the cause of the low temperature SPT of the SRO thin film, the exact mechanism of the SPT has not been answered. Many fabrication factors are known to have effects on the onset of SPT of SRO thin films, which include film thickness, oxygen partial pressure during film growth, substrate strain, and temperature. Among these, oxygen partial pressure during film growth is closely related to the oxygen vacancy, i.e., lesser oxygen gas pressure during sample fabrication increases the oxygen vacancy in the film. Since oxygen vacancy can be considered as a defect state, the conductivity, optical transparency, and other important physical properties of films are closely related to the amount of oxygen vacancy. Previously, it is reported that the existence of oxygen vacancy in the SRO thin film varies the orbital hybridization between Ru 4d and O 2p, affecting the electronic structure and transport properties of thin films. It is also found that the formation of oxygen vacancy at the Sr-O plane is more energetically favorable than the Ru-O plane. Apparently, the concentration of oxygen vacancy in oxide materials has become a critical adjusting parameter for engineering the desired properties.

To illustrate the SPT of the SRO thin films, two different structures of SRO thin films are shown in Fig. 1, a tetragonal SRO (T-SRO), and monoclinic SRO (M-SRO). As the oxygen vacancies are generated at the Sr-O planes in T-SRO, the SRO thin film exhibits a tetragonal structure owing to the increase in Ru-Ru repulsion that suppresses the octahedral tilting, as shown in Fig. 1(a). On the other hand, in the case of M-SRO, the concentration of the oxygen vacancy is low and octahedral tilting prevails with the tilting angle of ∼0.45° due to a substrate induced strain, as shown in Fig. 1(b).

To understand the role of oxygen vacancy, substantial efforts have been devoted with diverse analytic techniques, e.g., positron annihilation lifetime spectroscopy and scanning transmission electron microscopy. Unfortunately, due to their poor sensitivity of probing techniques and highly diluted contents, the direct observation of oxygen vacancy still remains as a great challenge and its role in transport properties of thin films is not well understood. In this report, we investigated the correlation between the SPT of the SRO thin film and oxygen vacancy with in situ X-ray diffraction (XRD) and ambient pressure X-ray photoelectron spectroscopy (AP-XPS). In situ XRD shows that the SPT occurs from M-SRO to T-SRO near ~200°C, regardless of the pressure environment. When XPS is applied to monitor the surface chemical properties of the SRO film during the SPT temperature, Ru core level spectra shift toward the lower binding energy direction, whereas the Sr core level spectra move to the opposite direction under ultra-high vacuum (UHV) conditions. The directions of the core level shift of Ru and Sr can be interpreted with the formation of oxygen vacancy during the SPT of the SRO thin film. The features of valence band spectra of SRO thin films across the SPT temperature also reveal the evidence of oxygen vacancy formation. Then, when the oxygen vacancy is filled, the observed peak changes of XPS disappear. Combined analysis of in situ XRD and XPS results suggests that oxygen atoms migrated to the surface to form the surface oxide across the SPT temperature, revealing the close correlation between oxygen vacancy and SPT of SRO thin films.

**EXPERIMENT**

SRO thin films were epitaxially grown on TiO$_2$ terminated SrTiO$_3$ (STO) (001) substrates by pulsed laser deposition (PLD) with a KrF excimer laser (λ = 248 nm) pulses. The temperature of the STO substrate during the deposition was fixed at 960°C with an oxygen partial pressure of 20 mTorr and 100 mTorr for T-SRO and M-SRO thin films, respectively. Details on sample fabrication and characterization can be found in elsewhere.
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 measurements were carried out at beamline 5D at Pohang Light Source in Korea. The X-ray energy was tuned to 10 keV by using a double bounce Si (111) monochromator. The thickness of T-SRO and M-SRO films estimated by the oscillation fringes period in the XRD pattern was approximately 40 nm. AP-XPS measurements were carried out at SOLEIL synchrotron (TEMPO beamline) and PLS-II (8A2 beamline). In the AP-XPS systems (PHOIBOS NAP 150, SPECS) consisting of the differential pumping scheme and advanced electrostatic lens design, it was possible to measure photoemission spectra up to the gas pressure of 25 Torr. Repeated cycles of oxygen annealing were carried out to achieve a clean surface of the sample was superb.

RESULTS AND DISCUSSION

Prior to XPS measurements, the crystal structure of the SRO thin film was confirmed by XRD. In Fig. 2(a), displaying the (002) specular reflection of T-SRO and M-SRO, the momentum transfer in the film normal z direction, q_z values of the T-SRO (red) peak are smaller than that of M-SRO (black) in reciprocal space, indicating that the lattice parameter of T-SRO along the a-axis is larger than that of M-SRO. The lattice parameters of T-SRO and M-SRO along the surface normal direction obtained from the (002) Bragg reflections are 3.969 Å and 3.952 Å, respectively, as shown in Fig. 2(b). The lattice parameters of T-SRO, M-SRO, and STO along the a-axis are all 3.905 Å due to compressive strain of the STO substrate exerted on the epitaxially stacked SRO thin film. The oscillatory patterns in Fig. 2(a) are thickness fringes originated from the interference between X-rays reflected from the substrate and thin film. The sharp fringes patterns indicate that the crystalline quality of the thin film sample was superb.

First, the XRD measurements of the off-specular (103), (013), (103), and (013) Bragg reflections were carried out at elevated temperatures under UHV condition, as shown in Fig. 2(c). The sample temperature is indicated just below the spectra, e.g., room temperature RT (I), 150 °C (II), and 220 °C (III), for T-SRO and M-SRO. In Fig. 2(c), at RT and 150 °C, the q_z values of the M-SRO, (013) and (013) peaks are different from the (103) and (103) peaks, whereas T-SRO shows that q_z values of all [103] family peaks are the same. In the off-specular geometry, the identical q_z values of all [103] family peaks of the T-SRO film indicated the absence of octahedral tilting. On the other hand, in the case of M-SRO, the values of q_z of (103) and (103) reflections are 4.756 Å⁻¹ and 4.780 Å⁻¹, respectively. Previously, in M-SRO, having different q_z values in the off-specular geometry has been interpreted as the presence of octahedral tilting. On the other hand, at 220 °C (III) (yellow shaded), all [103] family peaks of M-SRO have the same q_z values, suggesting that octahedral tilting disappear. In short, the M-SRO undergoes SPT from the monoclinic to tetragonal phase near ~200 °C, whereas T-SRO shows no change under UHV. We note that the shift toward the lower q_z observed in all samples as temperature increases was due to the thermal expansion.

As mentioned previously, the role of oxygen vacancy becomes important as the phase transition takes place in the M-SRO thin film. To observe any surface modification due to the changes of oxygen vacancy at the onset of the phase transition, XPS measurements were carried out under the identical pressure and temperature conditions as used in Fig. 2. It is necessary to note that XRD provides bulk sensitive information, whereas XPS delivers surface sensitive information. Nevertheless, if there is any change of oxygen vacancy states at the bulk and interface of the SRO film, there is a possibility that the oxygen atoms out of the vacancies can migrate to the surface, resulting in the changes on surface chemical states.

Figure 3(a) shows the XPS spectra of Ru 3p, Sr 3d, and O 1s of T-SRO and M-SRO films obtained under UHV conditions from RT to 220 °C. The measurements were performed in the order of RT (I), 150 °C (II), 220 °C (III), and RT (IV). As shown in Fig. 3(a), no

FIG. 2. (a) X-ray diffraction (XRD) measurement of T-SRO (red) and M-SRO (black) along the surface normal direction. (b) Schematic view of T-SRO (left) and M-SRO (right). The lattice parameters of T-SRO and M-SRO along the c-axis are 3.969 Å and 3.952 Å, respectively, and the lattice parameter of T-SRO and M-SRO along the a-axis is equal to 3.905 Å due to epitaxial strain by the STO substrate. (c) XRD measurements of T-SRO (top) and M-SRO (bottom) along the off-specular direction. Each black, red, blue, and green lines indicate the XRD results along the off-specular (103), (013), (103), and (013) directions, respectively. There are no changes found in T-SRO, but M-SRO shows that SPT occurs at 220 °C.
changes are found in the Ru 3p spectra in both T-SRO and M-SRO during I → II step. On the other hand, as the temperature increases above the SPT temperature, ∼220 °C, Ru 3p spectra of both films show the core level shift toward the lower binding energy direction. The binding energy of Ru 3p is shifted from 464.0 eV to 462.6 eV in T-SRO, and from 464.1 eV to 463.6 eV in M-SRO, respectively. The core level shift of Ru 3p can be interpreted as following. In SRO, Ru atoms share the electron with oxygen atoms in the SRO lattice. As mentioned previously, during the SPT, the presence of oxygen vacancy at the Sr-O plane is energetically more stable than the Ru-O plane. Additionally, Ru atoms share less electrons with oxygen atoms along the z direction as oxygen vacancy is being generated. Also, the lattice along the z direction is being stretched during SPT, which changes the Coulomb interaction with neighboring oxygen atoms, can induce core level shift of Ru to lower binding energy direction. As the sign of oxygen vacancy starts to show near SPT temperature, it can be assumed that the origin of oxygen vacancies formed on surface is due to the strain release between STO and SRO across the SPT temperature.

In the case of Sr 3d spectra in Fig. 3(a), T-SRO and M-SRO show slightly different response to temperature than Ru 3p spectra. In the T-SRO film, the Sr 3d spectra show little change during I → II step. Then, as temperature reaches 220 °C, the peak at higher binding energy side, ∼133.5 eV, shows a significant increase. On the other hand, the M-SRO film shows similar, but smaller changes, as the T-SRO film is at temperature of 150 °C (I → II) and remains unchanged at 220 °C. The Sr 3d film returns to the original lineshape at RT, showing the reversibility during SPT. On the other hand, the line-shape after annealing is only slightly different from the lineshape at RT in the T-SRO film since only the M-SRO film shows SPT during the annealing process. According to the previous reports, the binding energies of Sr 3d5/2 in the SRO lattice and SrO1+x oxide are ∼132.0 eV and ∼133.5 eV, respectively. That is, in both films, the spectral weight of the SrO1+x oxide is increasing as the temperature increases. This observation can be possibly interpreted with the migration of oxygen atoms to the surfaces. As oxygen vacancy starts to form near the SPT temperature, the oxygen inside the film starts to migrate and moves to the Sr-terminated surface, which is shown as the increased oxidation state of the Sr 3d spectra in Fig. 3(a). In other words, the spectral change of Sr 3d shows that the metastable Sr oxide layer is being formed on the surface through oxygen migration across the SPT temperature. The core level shift of the Sr 3d spectra toward the lower binding energy direction is negligible compared to Ru 3p spectra even if the oxygen vacancy is generated in the Sr-O plane. As the distance between Sr atoms and oxygen vacancy is much larger than that of Ru-O while the symmetric array of the Sr atoms is unchanged, the Coulomb potential that Sr experiences from the oxygen vacancy is much less than Ru. Additionally, the electronegativity of Sr is much lower than Ru. The comparison of Sr 3d spectra between RT and T = 220 °C is shown in Fig. S1 of the supplementary material, which clearly shows the presence of SrO1+x oxide. It is important to note that the enhanced spectral contribution of oxides at T = 220 °C reduced to original when the temperature reduces to RT, i.e., a clear indication that oxygen comes from the inside of the films.

Since the number of oxygen vacancy is higher in the T-SRO film, the presence of migrating oxygen, i.e., surface SrO1+x oxide, is more significant than M-SRO. In the case of the M-SRO film, the sign of oxygen vacancy exists from the increase in SrO1+x oxide, yet the amount of surface oxide is not higher than T-SRO, resulting in the smaller increase in SrO1+x oxide. As stated previously, the structure of the SRO thin film becomes tetragonal (monoclinic) below (above) 60 mTorr of oxygen partial pressure. The T-SRO used in this experiment was fabricated at 20 mTorr of oxygen partial pressure, whereas M-SRO was fabricated at 100 mTorr of oxygen partial pressure. Thus, the oxygen vacancy concentration of T-SRO is much larger than the M-SRO, explaining the result of Sr 3d spectra. In addition, as the oxygen in T-SRO can move easily due to larger oxygen vacancies, the peak shift of T-SRO becomes large when the same thermal energy is exerted on both T-SRO and M-SRO. The variation of surface SrO1+x oxide and oxygen vacancy of Sr 3d spectra can be also found in O 1s spectra in Fig. 3(a). The amount of
The evidence of oxygen vacancy can further be found in the valence band spectra. In Fig. 3(b), the valence band spectra at each temperature and the comparison of the valence band spectra between RT and 220 °C are shown for both T-SRO and M-SRO films. In the case of the T-SRO film, the difference of the valence band spectra between RT and 220 °C shows the sign of enhancement near the Fermi level and ∼2.5 eV (blue line), which matches well with the oxygen vacancy features of density of states estimated from first-principle calculation in the previous report.19 However, the difference of valence band spectra of the M-SRO film does not show much resemblance to that of T-SRO, which is possibly due to the lower concentration of the oxygen vacancy. It is consistent with the results of XPS core level spectra. The Ru 3d and Sr 3p spectra also exhibit an almost identical trend, as shown in Fig. S2. When the temperature is reduced back to room temperature, all the positions of XPS in M-SRO are back to the initial state, showing the reversibility of SPT. In the case of T-SRO, the lineshape of spectra is almost recovered during the annealing process with only slight difference. It is to be noted that this measurement was carried out under UHV, and no extra oxygen was being added to the film. This explains that the generated oxygen vacancy returns to the equilibrium position, and the structure is regained.

To check out the role of oxygen vacancy, the external source of oxygen was introduced by utilizing AP-XPS. Figures 4(a) and 4(b) show the Ru 3p, Sr 3d, O 1s, and valence band spectra under O2 pressure of 100 mTorr at elevated temperatures. As shown in Fig. 4, the measurements of AP-XPS were conducted in the order of RT (V), 150 °C (VI), 220 °C (VII), and RT (VIII). The XPS spectra in Figs. 4(a) and 4(b) showed no visible changes during the SPT temperature. There is a small change in the Sr 3d in T-SRO film, but the amount of change is much smaller compared to the UHV condition. As the oxygen vacancy created during the SPT temperature has been filled up with the external oxygen source, the formation of surface SrO1+x oxides is reduced significantly. Both T-SRO and M-SRO spectra VIII are almost identical to the spectra V, indicating annealing at 220 °C does not create additional surface oxide. On the other hand, the surface adsorbed oxygen was detected, which was different from that of Fig. 3 (UHV condition). In Fig. 4(a), the positions of the higher binding shoulder of O 1s and Ru 3p spectra at 220 °C are different from that of Fig. 3(a). In the case of O 1s, the shoulder position, i.e., the surface oxide peak, is located at higher
binding energy side, ~ 0.7 eV, than Fig. 3, which is the indication of a different origin of oxygen. The binding energy of the surface oxide peak under UHV is 531.7 eV, whereas 531.0 eV in the oxygen pressure environment, as shown in Fig. S3. The origin of the O 1s shoulder peak in Fig. 3(a) is due to the internal oxygen migration, whereas that in Fig. 4(a) originates from external gaseous oxygen. It is to note that the peak difference of M-SRO between UHV and oxygen pressure condition is not clearly visible due to the relatively lower concentration of oxygen vacancy than T-SRO.

To confirm if our observation in Figs. 2 and 3 is coming from the surface defect or the degradation of the sample, the SPT was checked again under oxygen condition. As shown in Fig. 5, we monitored the off-specular (103), (013), (103), and (013) reflections of T-SRO and M-SRO films under 100 mTorr of oxygen pressure at elevated temperature. From the bottom, the measurements were performed in the order of RT (V), 150 °C (VI), and 220 °C (VII). As previously observed under the UHV conditions, M-SRO shows SPT at 220 °C, but T-SRO shows no change. That is, the SPT of the SRO thin film takes place regardless of the external gaseous pressure. The result of Fig. 5 confirms that the validity of samples and the observation of XPS measurements is mostly from the surface effect, indicating that the surface structure during SPT could be different from that of bulk properties.

CONCLUSION

In this paper, with in situ XRD and AP-XPS, we investigated the presence of oxygen vacancy of T-SRO and M-SRO films across the SPT temperature. Across the SPT temperature, surface chemical shifts were observed in both T-SRO and M-SRO films, and the analysis of AP-XPS spectra suggested the sign of surface migration of lattice oxygen. In addition, the emerging valence band features during the SPT temperature showed good agreement with previously reported theoretical electronic structure calculations. Our finding suggests that there exists a close relation between oxygen vacancy and SPT. Our observation also suggests that bulk SPT of the SRO thin film can be possibly observed with XPS in an indirect manner. As a further study, the in situ surface XRD can be helpful for identifying surface structural changes during SPT.

SUPPLEMENTARY MATERIAL

See the supplementary material for comparison and analysis of core level photoemission spectra displaying surface oxide formation across structural phase transition on SRO thin films.

ACKNOWLEDGMENTS

This work was supported by the National Research Foundation of Korea (Grant Nos. NRF-2015R1A5A1009962, NRF-2017K1A3A7A90916316, NRF-2019R1A2C2080852, NRF-2017K1A3A7A90916395, and NRF-2019K1A3A1A21030984) and the GIST Research Institute Grant funded by the Gwangju Institute of Science and Technology (GIST) 2019. The Advanced Light Source is supported by the Office of Basic Energy Sciences of the U.S. DOE under Contract No. DE-AC02-05CH11231.

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