Article

Analysis of Metal-Insulator Crossover in Strained SrRuO$_3$ Thin Films by X-ray Photoelectron Spectroscopy

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Abstract: The electronic properties of strontium ruthenate SrRuO$_3$ perovskite oxide thin films are modified by epitaxial strain, as determined by growing on different substrates by pulsed laser deposition. Temperature dependence of the transport properties indicates that tensile strain deformation of the SrRuO$_3$ unit cell reduces the metallicity of the material as well as its metal-insulator-transition (MIT) temperature. On the contrary, the shrinkage of the Ru–O–Ru buckling angle due to compressive strain is counterweighted by the increased overlap of the conduction Ru-4d orbitals with the O-2p ones due to the smaller interatomic distances resulting into an increased MIT temperature, i.e., a more conducting material. In particular, in the more metallic samples, the core level X-ray photoemission spectroscopy lineshapes show the occurrence of an extra-peak at the lower binding energies of the main Ru-3d peak that is attributed to screening, as observed in volume sensitive photoemission of the unstrained material.

Keywords: metal-insulator-transition; perovskite oxides; laser deposition; stress-strain relations; angular resolved photoemission spectroscopy; X-ray photoemission spectroscopy

1. Introduction

Electron hybridization in solids in competition with Coulomb interactions plays a fundamental role in the quantum properties of these transition metal oxide materials [1–7]. In the context of spintronics, magnetic and electronic reconstructions at the interface have been often reported, with their origin lying in the delicate interplay between the charge, spin and orbital degrees of freedom. Looking at the strength of electron hybridization and localization, near a surface or interface the reduced translational symmetry modifies the electronic properties with important consequences on the magnetic order parameter, the transition temperature and the metallic vs. insulating behaviour, thus hindering the achievement of the desired performance of interface-based devices.

Strontium ruthenate SrRuO$_3$ (SRO) is an itinerant ferromagnet and a prototypical spintronics system [8,9]. Because of the structural and chemical similarities of all oxide perovskites, the growth of very high-quality epitaxial SRO-based heterostructures is indeed possible, thus allowing to explore new perspectives in the field of electronic, magnetic and optical devices [10–12]. Yet, interest in SRO goes well beyond its use as a functional layer embedded in oxide heterostructures. As a matter of fact, several works have provided evidence of a strong correlation between transport, magnetic and structural properties of SRO making it a very interesting model material for the study of itinerant magnetism in oxide systems [8,13–18]. However, as in many other oxide materials, surface-
in general interfacial-) and defect states play critical roles in mediating ferromagnetism, due to the modified chemistry of the first top—or interfacial—layers.

Here we report on the influence of the epitaxial strain on the electronic properties of SRO thin films at the surface. Epitaxial thin films were grown on different substrates by pulsed laser deposition (PLD) technique, characterized by crystallography and their properties were probed by electron spectroscopy. As expected, the metal-insulator-temperature $T_{MI}$ turns out to be modulated by the in-plane biaxial strain (it increases in compressively strained films, while it decreases in tensile strained ones). Core level soft X-ray photoemission spectroscopy with tuneable synchrotron radiation reveals the occurrence of an extra peak at Ru-3d, which is the signature of bulk-like hybridization and metallicity, only in the most metallic SRO films. In particular, only compressively strained SRO thin films display bulk-like electronic properties also in proximity of the surface (i.e., within 1 nm) as opposite to SRO films under tensile strain. This evidence that the surface/interface electronic charge distribution can be effectively controlled via atomic-precision growth techniques, is of importance for possible applications in the field of integrated spintronics.

2. Materials and Methods

PLD growth of epitaxial SRO thin films was performed using a KrF excimer pulsed laser source, with a typical energy density per laser shot of about 3 J/cm$^2$ and a laser repetition rate of 3 Hz ablating from a SRO stoichiometric target (purity at 99.99%). The films growth was performed in an ultra-pure oxygen background atmosphere (purity at 99.9999%). All of the investigated samples were grown at 0.1 mbar with a substrate temperature at 500 °C. After the film growth, the samples were cooled down to room temperature in about 15 min in oxygen at the same deposition pressure. Structural characterization was carried out using a four-circle Panalytical X’pert diffractometer with a Cu K$_\alpha$ radiation source. Surface morphology of the SRO samples was investigated by a Supra 40 field-emission gun (FEG) scanning electron microscope (SEM) equipped with a Gemini column and an In-lens detector yielding increased signal to noise ratio. EDS experiments were carried out by Oxford LN$_2$-free X-Act Silicon Drift Detector and chemical composition were calculated by Aztec software. Soft X-ray radiation spectroscopy (XPS and ARPES) and LEED experiments were carried out at the APE-IOM beamlines at Elettra. The measurements were recorded on the samples that were transferred in situ directly after growth under UHV conditions (base pressure < 2 × 10$^{-10}$ mbar) to the APE spectrometer end stations. XPS spectra were taken with a photon energy of 900 eV with an Omicron E125 hemispherical analyzer. The ARPES data were acquired with linearly polarized synchrotron radiation and a Scienta DA-30 hemispherical analyser.

3. Structural Properties

SRO possesses an orthorhombic crystal structure with lattice constants $a = 0.556$ nm, $b = 0.553$ nm and $c = 0.784$ nm. Yet, it shows a perovskite-like sub-unit cell with lattice constants $a = b = c = 0.393$ nm for bulk material [19,20]. For the sake of simplicity in this paper, the Miller indices of SRO will be referred to the perovskite cell. In order to induce different in-plane biaxial strain, SRO thin films were grown on (110) NdGaO$_3$ (NGO), (001) SrTiO$_3$ (STO) and (110) GdScO$_3$ (GSO) single crystal substrates. Considering the SRO bulk lattice parameter of 0.393 nm, SRO films are expected to grow strained under compressive (i.e., $-1.8\%$ and $-0.6\%$ on NGO and STO, respectively) or tensile (i.e., $+0.9\%$ on GSO) conditions. Deposition of SRO thin films was optimized at 0.1 mbar of ultra-pure oxygen pressure without any post-annealing process of the samples. Since the high-pressure background condition during the growth (i.e., 0.1 mbar) could possibly affect the stoichiometric ratio of the heavy-ion elements [21–23], energy dispersive spectroscopy (EDS) analysis was performed. Because of the overlap between the characteristic peaks of SRO film with some elements present in the substrates, a reliable stoichiometric analysis was obtained by performing the EDS analysis on SRO films grown on NGO substrates (Figure 1).
EDS experiments confirm that SRO films grown at 0.1 mbar, with a target-to-substrate of 5 cm and by using high energy laser pulses of 300 mJ show the optimal heavy-ion stoichiometric ratio (namely, Sr:Ru = 1:1—with an experimental error of about 5%). X-ray diffraction (XRD) characterization was routinely performed for all of the samples. The θ–2θ scans in symmetrical Bragg-Brentano configuration for optimized SRO films only contain (00l) peaks, indicating the preferential c-axis orientation of the films (in left a) of Figure 2 data refers to a SRO sample grown on NGO substrate). As expected, the out-of-plane lattice parameters evolve as a consequence of the substrate-induced strain mechanism (data are reported in panel b) of Figure 2).

An in-plane compressive (tensile) strain induces the elongation (contraction) of the out-of-plane lattice parameters. In particular, the elastic response of the lattice parameters as a function of a biaxial strain can be obtained by the formula

$$\frac{\Delta c}{c} = \frac{2\nu}{1-\nu} \cdot \frac{\Delta a}{a}$$  \hspace{1cm} (1)

where \(\nu\) is the Poisson ratio which can range between 0 and 0.5 (typical values found in oxides is 0.33). All of the measured structural deformations are compatible with elastic strain mechanism only (Figure 2b). The film thickness and surface roughness were probed by low-angle X-ray reflectivity (XRR). Representative XRR measurement of a SRO sample is shown in the Figure 3a.
Simulations of the low-angle XRR data were performed by means of the IMD 4 package of XOP 2.3 software [24,25]. The fitted curves (red curves included in Figure 3a) nicely match the expected value of 15 nm. XRR oscillations are recorded up to 2θ values of 5°, while, above this angle, the oscillations fall below the experimental sensitivity of our X-ray diffractometer [26,27]. Surface long range order was probed by in situ low-energy electron diffraction (LEED). Figure 3b reports the LEED pattern of a SRO film grown on STO substrate, measured by electrons with 80 eV kinetic energy, which shows sharp diffraction spots arranged in an in-plane squared coordination of the pseudocubic SRO structure.

4. Transport Properties

Electrical transport measurements were carried out by standard four-probe dc technique, with a bias pulsed and reversed current. Here we report the resistivity versus temperature behaviour of SRO films grown on NGO (−1.8% compressive strained), STO (−0.6% compressive strained) and GSO (+0.9% tensile strained) substrates (Figure 4a).

Among these, SRO film deposited on GSO substrates shows the highest resistivity values and a very small residual resistivity ratio (RRR) value (slightly larger than 1), indicating a higher degree of structural disorder in the film [28–30]. The negative slope of the resistivity curve in the low temperature range (T < 50 K) is consistent with the presence of impurities that act as scattering centres determining the insulating character of SRO film on GSO at low temperatures [31]. Conversely, SRO film grown on NGO substrate is the most metallic one, across the whole temperature range, with room temperature resistivity of about 600 µΩ·cm, which is very close to those reported for SRO thin films [32] while
thicker films are characterized by lower (i.e., factor of 2) values [13,14,33]. Moreover, the RRR of SRO film grown on NGO is a few percent smaller than the STO sample, which can indicate that the STO substrate (i.e., low strain imposed) might favorize better structural properties (i.e., less defects). The paramagnetic to ferromagnetic transition in SRO is associated to a change in the resistivity curve, namely from a linear-$T$ to a parabolic-$T^2$ behaviour at high and low temperatures, respectively [8,34]. In this respect, the analysis of the derivative $\Delta \rho / \Delta T$ provides the evaluation of the $T_{MI}$ temperature by the onset of its increase at low temperatures (see arrows in Figure 4b). The $T_{MI}$ temperature is severely reduced in the case of the film grown on GSO (91 K) film with respect to the SRO samples grown on STO (i.e., 106 K) and NGO (i.e., 127 K).

5. ARPES Experiments

Photoemission investigation was performed with polarized synchrotron radiation on SRO films transferred in situ (base pressure during the transfer never exceeds $2 \times 10^{-10}$ mbar) directly after the growth, to the two end-stations of the APE beamline [35]. Such a strategy allowed us to perform spectroscopic experiments on pristine uncontaminated samples [36–38]. In particular, the most metallic SRO film (i.e., the one grown on NGO substrate) was investigated by means of ARPES. Measurements were performed at room temperature. Figure 5 shows the ARPES energy dispersion map obtained with linearly polarized light of $h\nu = 70$ eV. Two arcs dispersing near ±0.5 confirm.

![Figure 5](image-url)

**Figure 5.** ARPES energy dispersion map of SrRuO$_3$ measured with 70 eV photon energy at room temperature. (a) Valence band dispersion and angle-integrated DOS showing spectral intensity at the Fermi level (white profile superimposed to ARPES spectrum). (b) Zoom to the area close to the Fermi level indicating the states responsible for the metallic nature of the film.

The integrated density of state (DOS) is superimposed to ARPES map of Figure 5 (white solid line). In particular, the spectral weight close to the Fermi level is ascribed to the Ru-4d $t_{2g}$ orbital, while the valence band (VB) is mainly associated to O-2p orbitals [39]. The overall DOS shape is consistent with those reported in the literature, showing the VB bandwidth spans between 8 and 1.8 eV binding energy [39,40]. Some discrepancies in the ARPES data are ascribable to the different experimental conditions. In particular, a tiny electron pocket centred at $k_x = 0$ is missing in our spectra, while it is reported in refs [40,41] but not in ref [42]. However, at difference to our case, the data in other reports were mainly acquired using unpolarized He–I radiation (21.2 eV) at $T < 15$ K [40–42]. The observed discrepancies are therefore consistent either with a different Brillouin zone sampling due to the excitation energy either to matrix element effects induced by light polarization or experimental geometry. The overall good consistency between our data and those reported in literature [40–43] is indicative of optimal stoichiometry bulk-like electronic properties of SRO samples grown on NGO. In particular, due to the high surface sensitivity intrinsic to ARPES technique, such properties are maintained up to the very surface of the films (i.e., 3–4 Å).
6. Core Level Photoemission Spectroscopy Experiments

Soft X-ray photoemission spectroscopy (XPS) spectra were measured at room temperature with linearly polarized synchrotron radiation on SRO samples transferred in situ directly after growth. The near surface properties of the SRO thin films were quantitatively explored by measuring the relative intensities of core-level photoemission spectra as well as the valence density of states (Figure 6) [44–48].

The XPS survey scan displays the main peaks of both Sr and Ru with no trace of material segregation. Moreover, the XPS spectra of the valence band show differences in the density of states in the Fermi level region that reflect the transport results discussed above. The connection between strain and metallic character can be also seen from the changes at the Fermi level: the density of states becomes larger for the compressively strained samples (i.e., NGO and STO) compared with the tensile strained one (GSO) thus confirming that SRO grown with the most compressive strain possesses has a clear metallic character up to the surface. XPS of the Ru-3d, Sr-3p and O1s peaks were investigated in the most metallic sample. Even though Sr-3p1/2 peak partially overlap Ru-3d3/2, a doublet structure can be clearly seen at Ru-3d. These peaks were attributed to well-screened (lower binding energy peak) and poorly screened (unscreened—higher binding energy peak) components arising in the photoemission process [44,49–51]. Consistently, O-1s peak also shows a double-peaks structure, with the satellite peak attributed to an electron screening mechanism [52]. In order to correlate the presence of the Ru-3d extra-peak with the metallicity of the system, lineshapes of Ru-3d and Sr-3p peaks were analyzed in all SRO samples (Figure 7).

![Figure 6.](image-url)

Figure 6. (a) XPS survey scan of a SRO film grown on NGO substrate (excitation photon energy was \( h\nu = 850 \text{ eV} \)); (b) high-resolution XPS scans of O-1s (b-1) and Ru-3d/Sr-3p (b-2) core levels, along with the valence band (b-3); In particular, valence band scans are reported for all of investigated SRO samples (i.e., namely grown on NGO (blue), STO (red) and GSO (green), respectively; in the inset, a zooming out of the Fermi’s edge).
Figure 7. High-resolution XPS scans of Ru-3d/Sr-3p peaks for SRO films grown on NGO (a), STO (b) and GSO (c); photon energy was $h\nu = 850$ eV. Results of fitting procedures are also reported (area of well-screened peak illuminated by red filling color).

The best fit was obtained allowing for two energy shifted components for Ruthenium, consistently with the reference papers [44,45,51]. The intensity ratios between the two components of each spin-orbit doublet are 4/3 for Ru-3d and 1/2 for Sr-3p. By comparing the Ru-3d spectra with those reported for nearly perfectly stoichiometric SRO [44–46]. Moreover, in tensile-strained SRO sample with low metallicity (i.e., SRO on GSO), the well-screened peak is suppressed while it is well-defined in compressive-strained highly metallic ones (i.e., SRO grown on NGO and STO).

7. Conclusions

The ensemble of photoelectron spectroscopy results shows how both the buckling of the bond angle and the changes in the bond length must both be considered to explain the electronic transport properties. The width of the conduction band strongly depends on the superimposition of the oxygen $O-2p_\sigma$ and $O-2p_\pi$ orbitals and Ru-4d orbitals. On one hand, tensile strain—occurring in SRO samples grown on GSO—tends to make the structure near-cubic (i.e., the Ru-O-Ru buckling angle decreases) thus better aligning Ru-4d and O-2p orbitals. On the other hand, the increased inter-atomic distance makes such a superimposition smaller thus resulting in a less conductive sample, as shown from the increase in resistivity. Differently, compressive strain—occurring in SRO samples grown on NGO and STO—increases the conductivity albeit it shrinks the Ru-O-Ru angle, meaning that the overlap of the conduction orbitals is increased by the smaller interatomic distance in spite of their axes becoming more misaligned. The very presence of the Ru-3d well-screened peak in the soft X-ray photoemission spectra demonstrates that compressively strained SRO thin films display bulk-like electronic properties also in proximity of the surface.

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