SURFACE STABILITY OF EPITAXIAL LaNiO$_{3-\delta}$ THIN FILMS

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Thin LaNiO$_{3-\delta}$ films with pseudocubic (100) preferred orientation were prepared by reactive DC magnetron sputtering and in situ annealed in O$_2$ and vacuum. X-ray photoelectron spectroscopy (XPS) was used to determine the variation in composition of the films under high temperature annealing. The experimental O 1s and La 3d – Ni 2p$_{3/2}$ spectra of LaNiO$_{3-\delta}$ films was analysed in terms of O$^{2-}$, O$^{-}$/OH$^-$, and weakly adsorbed oxygen species. It was shown that the change in the type of conductivity from metallic to semiconducting one is accompanied by a marked increase in the intensity of the lateral (∼531 eV) peak of oxygen. The quantitative analyses of La 3d – Ni 2p$_{3/2}$ spectra show that the Ni/La concentration ratio significantly decreases after heating above the dehydration temperature. These variations in conductivity and surface composition were attributed to the loss of lattice oxygen with subsequent adsorption of O$^-$ and (OH)$^-$ anions and weakly adsorbed oxygen species from ambient air.

Keywords: rare earth alloys and compounds, oxide materials, XPS, surfaces and interfaces, LaNiO$_3$ thin film

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1. Introduction

In recent years, an increasing interest has been focused on metal oxides of oxygen octahedral structure due to their metal–insulator transition, high-temperature superconductivity, ferroelectricity, colossal magnetoresistance and on their heterostructures for future technological application potentials [1–4]. LaNiO$_3$ (LNO) is one of the few conductive oxides with a crystal structure suitable for integration in epitaxial heterostructures with perovskites because of its good transport properties, composition simplicity, and pseudocubic perovskite crystal structure with a lattice parameter $a = 3.84$ Å. In most studies, LNO films are considered as having good physical and chemical thermostabilities, and most metal oxide films, such as ferroelectric films, must be grown at a high temperature. Therefore, it is necessary to study the thermostability of these films.

It is known that considerable segregation of elements takes place in chemically synthesized LaNiO$_{3-\delta}$ samples, i.e. the surface concentration of various species differs from the volume one [5]. Another factor to be considered is the tendency of rare earth and nickel oxides to absorb water vapour and carbon dioxide from air, so that any ex situ exposure of these films to air will result in an uncontrolled reaction and stoichiometry [6].

Lanthanum hydroxide, in turn, tends to adsorb humidity at the surface and to react with carbon dioxide from the atmosphere in order to form lanthanum carbonate.

A study on the thermostability of LaNiO$_3$ films prepared by RF magnetron sputtering and in situ annealed at 265 °C were performed [7]. It was demonstrated that electric and optical properties of LNO films are unstable even at a relatively low temperature. But the annealing does not affect the lattice structure. The electronic structure of high-quality LNO thin films was investigated in situ using photoemission spectroscopy [8]. In that work it was shown that O 1s X-ray absorption spectrum is in good agreement with previous studies on polycrystalline LNO surfaces.

It was shown earlier that epitaxial LaNiO$_{3-\delta}$ films deposited onto (100)-plane oriented NdGdO$_3$ substrate demonstrated excellent in-plane orientation [9] and the surface La/Ni ratio was close to the stoichiometric one [10].
Angle dependent spectra, obtained with the Tunable High-Energy X-ray Photoelectron Spectrometer at the X-ray wiggler beam line BW2 of HASYLAB (Hamburg) reveal that significant variations of oxide versus hydroxide concentrations occur within the thin surface layer even after long-term (one year) exposure to the atmosphere. Estimated thickness of this hydroxide enriched layer is of about 2 nm [11].

In this paper, LNO films are deposited on NdGdO$_3$ substrates by a reactive DC magnetron sputtering system, and their composition and electrical conductivity dependence on high temperature annealing have been investigated.

2. Experimental details

Thin LaNiO$_{3-\delta}$ films were deposited onto monocrystalline (100)-plane oriented NdGaO$_3$ substrate by using a reactive DC magnetron sputtering technique. The ceramical LaNiO$_3$ target (25 mm in diameter and 2.5 mm thick) was prepared by pressing at 5·10$^8$ Pa and after sintering in air at 1000 °C for 10 h the La$_2$O$_3$ and NiO (99.99% purity from Aldrich-Chemie) powders in the stoichiometric ratio. The sputtering was performed in Ar and O$_2$ mixture (20:1) at pressure of about 15 Pa. To prevent the film bombardment by high energy ions during deposition, NdGaO$_3$ substrates were positioned in “off-axis” configuration at a distance of 15 mm from the symmetry axis of the discharge and 20 mm over the target plane. The substrate temperature was ∼750 °C. Under these conditions, the deposition rate was 25 nm/h, and the resultant thickness of LaNiO$_{3-\delta}$ film was about 0.1 μm. Films were annealed in situ in vacuum (10$^{-5}$ Pa) or in O$_2$ atmosphere (10$^5$ Pa).

X-ray photoelectron spectra were recorded using XSAM 800 (KRATOS Analytical, UK). The photoelectrons were excited using a non-monochromatized Al K$_\alpha$ (1486.6 eV) radiation source at 15 kV, 300 W. We were forced to use this relatively wide excitation to avoid interference from overlapping Ni(L$_3$M$_{2,3}$M$_{2,3}$) Auger features with XPS O 1s region using a Mg K$_\alpha$ source. The analyzer was used in a fixed retarding ratio mode with an energy resolution $\Delta E/E = 0.08\%$. The working pressure in the analysis chamber was maintained below 10$^{-7}$ Pa during the spectrum analysis.

Photoemission data were collected and processed by the KRATOS DS800 data system. After Al K$_\alpha$ satellites and background subtraction, the complex photoelectron spectra were decomposed into separate peaks by specifying the peak position – binding energy (BE), area, width, and Gaussian/Lorentzian ratio. Because any mathematical operation (such as data smoothing) on the raw data would distort the original physical picture, no such treatment was permitted in the quantitative data analysis. The accuracy of the measured lines BE and relative intensities were about 0.1 eV and 10% respectively. To control the sample contamination, the survey spectra of an air-exposed, fresh sample have been studied. Only lanthanum, nickel, oxygen, and carbon lines were identified within the probing depth of XPS. No more contamination related to the sample preparation was detected by XPS within the accuracy of our measurements. The adventitious carbon C 1s line was used for the fine correction of charging effects, supposing its BE is equal to 284.6 eV.

The influence of ion sputtering on the chemical composition of investigated compounds was treated by using Ar$^+$ ion bombardment at 2 keV and current density of 2 μA cm$^{-2}$ during 1 min.

The electrical conductivity measurements were performed by a standard DC four-probe method in the 78–300 K temperature range.

3. Results and discussion

According to the temperature dependence of resistance shown in Fig. 1, films may be divided into two groups with a metallic and semiconducting type of conductivity (see Table 1). Drastic changes in O 1s XPS spectra (Fig. 2) after high (above 700 °C) temperature annealing indicate strongly that the heat treatment induced changes in the films composition are responsible for the conversion of the conductivity.
Fig. 2. O 1s comparison spectra for the LNO films with metallic (as-grown and O750) and semiconducting (V700 and V750) type of conductivity. Annealing conditions are listed in Table 1.

Table 1. Annealing conditions and conductivity type of studied films.

<table>
<thead>
<tr>
<th>Film</th>
<th>Annealing condition</th>
<th>Conductivity type</th>
</tr>
</thead>
<tbody>
<tr>
<td>as-grown</td>
<td>—</td>
<td>metallic</td>
</tr>
<tr>
<td>O750</td>
<td>750  O₂</td>
<td>metallic</td>
</tr>
<tr>
<td>V450</td>
<td>450  vacuum</td>
<td>metallic</td>
</tr>
<tr>
<td>V700</td>
<td>700  vacuum</td>
<td>semiconducting</td>
</tr>
<tr>
<td>V750</td>
<td>750  vacuum</td>
<td>semiconducting</td>
</tr>
</tbody>
</table>

O 1s spectra are rather complex and may be roughly separated into three regions.

In all cases, an intense peak is observed at low energy around 529 eV. This peak seems to be very similar for all samples and may be attributed to O²⁻ anions of the crystalline network.

Beside this peak, a second component can be observed at higher energies. Its intensity is strongly affected by the heat treatment. For metallic films, this component only slightly varies in intensity and always appears as a shoulder around 531 eV and drastically increases for samples with dielectric conductivity. These lateral structures have been reported in many studies, but are not precisely attributed. The assumption of different valence states for the metal [12] or several final states [13] has been made, but in most cases this structure has been assigned to contamination surface species. The adsorption of (OH)⁻ or CO₂⁻ groups was suggested in comparison with the binding energy of the O 1s peak in hydroxide or carbonate compounds; other adsorbed species such as O₂²⁻ were also proposed.

Jimenez et al. [14] attribute the lateral peak to low coordinated oxygen ions at the surface of the samples. It was also suggested [15] that the oxygen partial integration might occur in the bulk and generate a high binding energy component, more or less intense, for the O 1s peak. In a mixed oxygen chemisorption-penetration situation, bond lengths and crystal potentials may vary significantly leading to a range of O 1s binding energies; this may also account for the large width and variable BE positions of the lateral structure. Moreover, (OH)⁻ in LNO may be in two forms: hydroxide and oxyhydroxides [16] with a close, but different O 1s core level BE in lateral peak energies range.

The high energy tail may include physisorbed (~534 eV), chemisorbed (~532 eV), structural H₂O and water in poor electrical contact with the films surface [17, 18]. In general, it may be concluded that numerous oxygen species with a BE distributed within a wide energy range may be involved in final O 1s spectra of LNO surface [19].

It is evident that these various oxygen contributions cannot be fitted in an unambiguous way even for XPS spectra obtained without considerable noise and with good resolution.

To obtain meaningful information about the surface composition from fitting results, one needs additional knowledge about the corresponding peaks nature and position. Such a priori information is highly desirable for the correct selection of an initial approximation in fitting experimental data, since the existence of local minima may lead to erroneous results.

Additional thermal treatments and ionic etching were realized in order to clarify the nature of species, contributing to O 1s spectra in the intermediate energy range. A highly resistive (R > 10 MΩ) film V750 with a semiconducting type of temperature behaviour was selected for this purpose. An in situ bombardment with Ar⁺ ions of 2 keV energy for 1 min was used for the surface cleaning.

After the deposition, the film was exposed to air for several days. Then it was placed in a vacuum chamber and XPS measurements were done for as-grown and Ar⁺ etched samples. Then the temperature was raised up to 550 °C and after a 30 hour exposure the XPS spectra were measured at this temperature and after slow (~100 °C/h) cooling down to room temperature. Finally, the film was exposed to air for 2 hours with a subsequent XPS analysis. The results are presented in Fig. 3. Note that spectra measured at 550 °C and after cooling were absolutely identical.

It is evident from the comparison spectra (Fig. 3, curves 1 and 2) that weakly absorbed species can be
effectively removed by low-intensity Ar$^{+}$ polishing. Further annealing leads to an additional decrease in the lateral peak (\(\sim 531 \text{ eV}\)) intensity and may be related to the film dehydration. It is known that thermal transformation from lanthanum hydroxide to oxide is a two-step mechanism \[16, 20\]. The intermediate product is lanthanum hydroxide oxide: \(\text{La(OH)}_3 \rightarrow \text{LaOOH} + \text{H}_2\text{O} \) (310–340$^\circ$C). Lanthanum hydroxide oxide decomposes in a second step in order to form the oxide according to \(2\text{LaOOH} \rightarrow \text{La}_2\text{O}_3 + \text{H}_2\text{O} \) (450–500$^\circ$C).

Thus, the single-peak difference spectra (Fig. 3, curve 5) may be attributed to hydroxide and/or oxyhydroxide at BE of about 531 eV and one would expect the full dehydration of LNO films. Then the shoulder (\(\sim 531 \text{ eV}\)) on the O 1s spectra may be attributed to tightly bonded oxygen species, probably surface and/or low coordinated oxygen. Unfortunately, the maximum available annealing temperature (550$^\circ$C) is not sufficient for the thermal decomposition of carbonate species: \(\text{La}_2\text{O}_2\text{CO}_3 \rightarrow \text{La}_2\text{O}_3 + \text{CO}_2 \) (600–800$^\circ$C) and possible surface contamination with these species cannot be excluded.

The XPS spectra of La 3d and Ni 2p$^{3/2}$ spectra region before and after film annealing at high temperature are presented in Fig. 4. Lanthanum oxide/hydroxide ratio was estimated from the relative areas of peaks corresponding to the oxide and hydroxide species, obtained from La 3d$^{3/2}$ spectra fitting. This ratio was 0.3 and 1.4 for as-grown and annealed sample, respectively. These results indicate that after annealing the lanthanum oxide/hydroxide ratio significantly increases.

The significant variations in the La 3d$^{3/2}$–Ni 2p$^{3/2}$ spectra region after the heating above dehydration temperature (Fig. 4) may be attributed to the partial disappearance of the nickel species as a result of the nickel hydroxide evaporation/decomposition at temperatures exceeding the melting point \(T_m = 230^\circ\text{C}\) \[21\]. The significant decrease of nickel/lanthanum concentration ratio from \(\text{Ni/La} = 1.02\) (close to stoichiometric one) for the as-grown sample to \(\text{Ni/La} = 0.34\) for the sample heated above the dehydration temperature evidently shows that the losses of nickel from the film surface take place under heat treatment.

Comparison of O 1s spectra measurements in situ after annealing in super-high vacuum and after exposure to ambient for a short (\(\sim 2\) hours) time demonstrate a rapid water absorption and LNO surface hydroxidation. The resulting difference spectrum (Fig. 3, curve 6) is rather broad and may be deconvoluted into two components centred at about \(\sim 531\) and \(\sim 533 \text{ eV}\) due to (OH)$^-$ and H$_2$O respectively. Finally, it may be concluded that even after surface cleaning and high temperature dehydration the intensity of the peak at \(\sim 531 \text{ eV}\) remains comparable with the O$^{2-}$–anion peak intensity. This lateral peak may be attributed to low coordinated oxygen \[13\] and/or to oxygen-deficient regions \[22\] in the LNO matrix. Unfortunately, the
spectral resolution of the data is insufficient to allow the observation of more than two spectral components even for dehydrated films measured with good statistics. Our attempt to fit the spectrum with an individual component expected for $\text{O}^-$ and (OH)$^-$ species demonstrates that these various contributions cannot be fitted in an unambiguous way; consequently, they are represented by one broad peak.

This is also true for the high energy tail composed of peaks corresponding to various types of absorbed water species \cite{17, 18}. That is the reason why we use the set of three peaks presenting the so-called O$^{2-}$, O$^-$ (including hydroxyl group (OH)$^-$), and H$_2$O species centred around 529, 531, and 533 eV respectively.

Several fitted spectra for dielectric LNO film before and after argon and temperature treatments (Fig. 5) demonstrate the effectiveness of such division for describing all spectra transformations due to film etching, annealing, and subsequent exposure to ambient atmosphere.

The same set of spectral components was used to fit a large number of spectra, recorded from different samples for further analysis of the effect of synthesis condition and subsequent annealing on the LNO films composition. For all treated samples, the energy separations (chemical shift) between the prime O$^{2-}$ and lateral peaks are in surprisingly good agreement (Fig. 6), and slight variations in the lateral peak BE may be attributed to the variation in the relative concentration of oxygen species contributing to peak broadening.

The effect of the thermal annealing on the relative concentration of various oxygen species concentration is shown in Fig. 7. It noteworthy that all films with dominating contribution of O$^{2-}$ species (the O$^{2-}$/O$^-$ concentration ratio $R_O > 1$) possess metallic type conductivity, while for films with $R_O < 1$ the temperature dependence of conductivity becomes semiconducting. Films remain metallic after high temperature ($\sim 750^\circ\text{C}$) annealing in oxygen atmosphere. This is also true for films annealed in vacuum at 450$^\circ\text{C}$. However, significant changes in the oxygen species ratio $R_O$ together with the accompanying conductivity conversion take place after high temperature annealing in vacuum. It is known that in LaNiO$_{3-\delta}$ transition
from metallic to insulating states takes place at $\delta = 0.25$ [23]. All this allows us to suppose that the loss of lattice $O^{2-}$ ions during the vacuum annealing is responsible for this transformation.

The sharp increase in the $O^{-}$ species concentration may be caused by the direct increase in the low coordinated oxygen content [14]. The increase in the number of oxygen vacancies, especially at the film surface, could be responsible for additional absorption of water, CO$_2$, and O$_2$ molecules with a subsequent formation of lower charge oxygen ions. All of them may participate in the increase of the lateral peak intensity.

However, spectral resolution of the system is too low for unambiguous interpretation of XPS spectra, and relative concentrations of various oxygen species cannot be determined with appropriate accuracy. Further investigations are necessary for a detailed and quantitative understanding of the nature of the metal–insulator transformation after high temperature annealing.

4. Summary and conclusions

In summary, we studied the changes in the chemical composition and electrical conductivity of LaNiO$_{3-\delta}$ films under high temperature annealing. The technique used in the study was mainly O 1s X-ray photoelectron spectroscopy. The experimental spectrum of LaNiO$_{3-\delta}$ films was analysed in terms of $O^{2-}$, $O^{-}/(OH)^{-}$, and weakly adsorbed oxygen species. As-grown films deposited onto (100)-plane oriented NdGdO$_3$ substrate demonstrated excellent in-plane orientation and metallic behaviour. Films remained metallic and no significant changes in their resistance and O 1s XPS spectra were observed after annealing in oxygen at $10^5$ Pa, 750°C. Conversion of the type of conductivity under high-temperature vacuum annealing may be attributed to the loss of lattice oxygen and reflects itself in the growth of the lateral peak at 531 eV in O 1s spectra. This increase in intensity may be attributed to the increasing number of oxygen vacancies with a subsequent increase in the concentration of low-coordinated oxygen species. These additional oxygen vacancies are likely to contribute greatly to rapid adsorption from ambient air and a significant increase in the surface contamination with $O^{-}$ and $(OH)^{-}$ and weakly adsorbed oxygen species. The quantitative analysis of the La 3d$_{5/2}$-Ni2p$_{3/2}$ spectra region before and after the heat treatment shows the significant decrease of Ni/La concentration ratio. The losses of nickel species may be attributed to the nickel hydroxide decomposition after heating above the dehydration temperature.

These changes in the composition and electric properties of LNO films must be taken into account when the films are applied in optical and electrical devices.

References


Santrauka

Plonieji daugiausia pseudokubine (100) kryptimi orientuoti \( \text{LaNiO}_3-\delta \) sluoksniai gaminti reaktyvinio nuolatinės srovės magnetroninio duškimo būdu ir \( \text{in situ} \) atkaitinti \( \text{O}_2 \) ir vakuume. Sluoksnų sudėties kitimas atkaitintu metu tirtas Rengeno fotoelektroninės spektroskopijos (RFS) metodu. Išsitikinti eksperimentiniuose \( \text{LaNiO}_{3-\delta} \) sluoksnų \( \text{O}^2- \), \( \text{O}^-/(\text{OH})^- \) ir silpnai surišto deguonies santykiais. Metalinio laidumo viršumą susidarydavo 

\[ \text{LaNiO}_{3-\delta} \] 


EPITAKSINIŲ \( \text{LaNiO}_{3-\delta} \) PLONŲ SKUOKSNIŲ PAVIRŠIAUS STABILUMAS

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