Epitaxial and layer–by–layer growth of EuO thin films on yttria–stabilized cubic zirconia (001) using MBE distillation

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We have succeeded in growing epitaxial and highly stoichiometric films of EuO on yttria–stabilized cubic zirconia (YSZ) (001). The use of the Eu–distillation process during the molecular beam epitaxy assisted growth enables the consistent achievement of stoichiometry. We have also succeeded in growing the films in a layer–by–layer fashion by fine tuning the Eu vs. oxygen deposition rates. The initial stages of growth involve the limited supply of oxygen from the YSZ substrate, but the EuO stoichiometry can still be well maintained. The films grown were sufficiently smooth so that the capping with a thin layer of aluminum was leak tight and enabled ex situ experiments free from trivalent Eu species. The findings were used to obtain recipes for better epitaxial growth of EuO on MgO (001).

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I. INTRODUCTION

Stoichiometric EuO is a ferromagnetic semiconductor with a Curie temperature ($T_C$) of 69 K and a band gap of about 1.2 eV at room temperature. Upon electron doping, the material shows a wealth of spectacular phenomena, including a metal–to–insulator transition and colossal magnetoresistance, where the change in resistivity can exceed 8–10 orders of magnitude. The Curie temperature can also be doubled by electron doping and even almost tripled by pressure. In the ferromagnetic state the conduction band shows a splitting of about 0.6 eV between the spin–up and spin–down states leading to an almost 100% spin polarization of the charge carriers in electron doped EuO. These properties make EuO a very attractive candidate for fundamental research in the field of spintronics.

EuO in thin film form has been studied already in the late 1960s and early 1970s. Use was made of coevaporation of Eu and Eu$_2$O$_3$ under technical vacuum conditions, i.e., pressures in the range of $10^{-6}$–$10^{-5}$ mbar. The preparation of thin films was considered as a convenient alternative synthesis route for EuO, alternative to that of the bulk synthesis which required very high temperatures with a delicate phase diagram. The use of the thin–film preparation route specifically facilitated doping dependence studies using rare–earth and transition–metal impurities.

After a pause of two decades, a strong renewed interest in EuO thin films emerged in recent years. This time it is the thin–film community which is making the attempt to utilize the extraordinary properties of EuO for device applications. Part of the motivation also originates from the fact that tremendous progress has been made in preparation technologies, e.g., molecular beam epitaxy (MBE) under ultra–high–vacuum conditions, and that new analysis methods have become available, e.g., synchrotron–based spectroscopies. All in all, these new efforts have culminated in a high point such that EuO thin films can be grown epitaxially on Si, demonstrating its potential for spintronics applications.

It is remarkable in the recent EuO research that control of the stoichiometry is nevertheless still a serious issue. Many studies reported that Eu$^{3+}$ ions were present in their films and/or that the magnetic moment per f.u. was not close to the expected $7\mu_B$ for a 4$f^7$ system. It is not clear in what precision the relative supply rates of oxygen and europium were controlled in these works. We will show below that this control need not be precise as long as one is in the so–called Eu–distillation condition during growth.

Also layer–by–layer growth has – to our knowledge – never been mentioned, although epitaxy has been often reported. We therefore set out to do a renewed growth study. We have chosen for yttria–stabilized cubic zirconia (YSZ) as substrate, the lattice constant of YSZ is 5.142 Å, practically identical to the 5.144 Å value for EuO at room temperature, and epitaxy has been reported already. Yet, it was also claimed that control stoichiometry is extremely difficult related to the fact that YSZ acts as a source of oxygen during the MBE growth process. We will show below that we do have achieved full control of the growth process, i.e., with stoichiometry control and layer–by–layer epitaxial growth, and that we can use these results as a firm basis for further studies including the doping dependence and the specific use of well–defined interfaces.
The EuO films were grown in an ultra–high–vacuum MBE facility with a base pressure of 2×10⁻¹⁰ mbar, maintained by a cryopump. High purity Eu metal from AMES Laboratory was evaporated from an EPI effusion cell with a BN crucible at temperatures between 460°C and 525°C. Proper degassing of the Eu material (mostly hydrogen gas) ensured that during Eu evaporation the pressure was kept below 3×10⁻⁹ mbar. The Eu deposition rate (4–8 Å/min) was calibrated using a quartz–crystal monitor which was moved to the sample growth position prior and after each growth. Molecular oxygen was supplied through a leak valve, and its pressure (4–16×10⁻⁸ mbar) was monitored using an ion–gauge and a mass–spectrometer. The growth was terminated by closing first the oxygen leak valve and then the Eu shutter after 30 s.

As substrates we used epipolished single crystals of YSZ from SurfaceNet GmbH and cleaved single crystals of MgO from TBL–Kelpin. The surface normal of the substrates are all the (001). The lattice constant of YSZ is 5.142 Å, very close to the 5.144 Å value for EuO at room temperature. The lattice constant of MgO is 4.21 Å. Prior to growth the substrates were annealed in situ at T = 600°C in an oxygen atmosphere of 5×10⁻⁷ mbar for at least 120 min in the case of YSZ, and in 1×10⁻⁷ mbar for at least 60 min for MgO, in order to obtain clean and well–ordered substrate surfaces. The substrates were kept at T = 400°C during growth.

The MBE facility is supplied with the EK–35–R reflection high–energy electron diffraction (RHEED) system from STAIB Instruments for in situ and online monitoring of the growth. The MBE facility is attached to an ultra–high–vacuum μ–metal photoemission chamber equipped with a Scienta SES–100 electron energy analyzer and a Vacuum Generators twin crystal monochromatized Al–Kα (hν = 1486.6 eV) source for in situ x–ray photoelectron spectroscopic (XPS) analysis. The overall energy resolution was set to 0.35 eV and the Fermi level $E_F$ was calibrated using a polycrystalline Au reference. The μ–metal chamber is also equipped with a Vacuum Generators Scientific T191 rear–view low–energy electron diffraction (LEED) system for further in situ structural characterization. The base pressure of the μ–metal chamber is 1×10⁻¹⁰ mbar, and all characterizations herein were carried out at room temperature.

The MBE facility is also connected to a separate ultra–high–vacuum chamber for the evaporation of aluminum as protective capping layer of the air–sensitive EuO films. This allows the ex situ characterizations using x–ray reflectivity (XRR), superconducting quantum interference device (SQUID), and x–ray absorption spectroscopy (XAS). The thickness of the aluminum capping is about 20–40 Å. The XRR measurements were carried out using a Siemens D5000 diffractometer. The magnetic properties of the films were determined using a Quantum Design MPMS–XL7 SQUID magnetometer. The XAS measurements were performed at the Dragon beamline of the National Synchrotron Radiation Research Center (NSRRC) in Taiwan. The spectra were recorded using the total–electron–yield method and the photon–energy resolution at the Eu $M_{4,5}$ edges (hν ≈ 1100–1180 eV) was set at ≈ 0.6 eV.

II. EXPERIMENT

The EuO films were grown in an ultra–high–vacuum MBE facility with a base pressure of 2×10⁻¹⁰ mbar, maintained by a cryopump. High purity Eu metal from AMES Laboratory was evaporated from an EPI effusion cell with a BN crucible at temperatures between 460°C and 525°C. Proper degassing of the Eu material (mostly hydrogen gas) ensured that during Eu evaporation the pressure was kept below 3×10⁻⁹ mbar. The Eu deposition rate (4–8 Å/min) was calibrated using a quartz–crystal monitor which was moved to the sample growth position prior and after each growth. Molecular oxygen was supplied through a leak valve, and its pressure (4–16×10⁻⁸ mbar) was monitored using an ion–gauge and a mass–spectrometer. The growth was terminated by closing first the oxygen leak valve and then the Eu shutter after 30 s.

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III. INITIAL STAGES OF GROWTH ON YSZ

Figure 1(a) shows the RHEED photograph of the clean and annealed YSZ (001) before growth, and Figs. 1(b)–1(c) show the photographs after 10 min of EuO growth. The Eu flux rates were 8.1–8.2 Å/min for (b)–(d) and 4.2–4.3 Å/min for (e)–(f). The oxygen pressure in the chamber was 4×10⁻⁸ mbar for (b), and 2×10⁻⁸ mbar for (c) and (e). No oxygen was supplied into the chamber for (d) and (f). The RHEED electron energy was 20 keV with the beam incident along the [100] direction.

![RHEED photographs](image)

FIG. 1: (Color online) RHEED photographs of (a) clean and annealed YSZ (001), [(b)–(d)] EuO films on YSZ (001) after 10 min of growth at 400°C with 8.1–8.2 Å/min Eu flux rates, and [(e)–(f)] with 4.2–4.3 Å/min Eu flux rates. The oxygen pressure in the chamber was 4×10⁻⁸ mbar for (b), and 2×10⁻⁸ mbar for (c) and (e). No oxygen was supplied into the chamber for (d) and (f). The RHEED electron energy was 20 keV with the beam incident along the [100] direction.
indicated for a two-dimensional (2D) layer–by–layer or Frank–van der Merwe growth mode. It is surprising that there are only five to six oscillations for all deposition conditions as indicated in Fig. 2 and that these oscillations even exist in the absence of oxygen in the MBE chamber. It is important to note that the oscillation period does not depend on the oxygen pressure $P_{\text{O}_2}$, thus also in the case of no oxygen in the chamber. This indicates that the oxygen needed for the formation of EuO must also come from the YSZ substrate. The $T = 400^\circ\text{C}$ substrate temperature apparently provides sufficient mobility for the oxygen ions to migrate to form at least five or six EuO layers. The oscillation period, which represents a formation of a new atomic single layer, is determined only by the Eu flux rate $\Phi_{\text{Eu}}$; reducing it by a factor of 2, from 8.1–8.2 to 4.2–4.3 Å/min, doubles the period, from 25 to 50 s.

LEED photographs for all these films displayed a good single crystallinity. Figure 3 depicts examples for the case of no oxygen in the MBE chamber during growth. Also here we can observe a perfect (001) surface of the EuO rock–salt structure, consistent with the RHEED results. The LEED photographs were taken at electron beam energies of 213–215 eV since lower energies did not provide stable patterns due to charging.

To investigate the implications of observing only five to six oscillations, we also carried out photoemission experiments on those films. Figure 4, left panel (a), shows the Zr 3$d$ and Eu 4$d$ core level XPS spectra which were collected at normal emission. It can be clearly seen that the Zr signal is reduced when comparing the clean YSZ (bottom curve – black) with the EuO–covered YSZ (top three curves – blue, green, and red). The EuO films here were grown without oxygen in the MBE chamber. Different Eu flux rates and total time of growth are indicated in the figure. It is remarkable that the EuO–covered YSZ spectra have very similar Zr signals, and also equal Eu intensity, despite the fact that the total amount of Eu exposure is twice as large in the two middle curves (green and red) than in the top one (blue). This indicates that in the absence of oxygen in the MBE chamber, the growth of EuO is limited to five to six monolayers only and that the rest of the deposited Eu metal is re–evaporated back into the vacuum (the substrate temperature is 400°C). In other words, the sticking coefficient for Eu after the completion of five to six monolayers is reduced to zero, suggesting that oxygen transport through EuO is much more difficult than in YSZ.

It is also important to investigate the chemical state of the Eu. Figure 4, right panel (b), depicts the O 2$p$ and Eu 4$f$ valence band spectra together with the Zr 4$p$, O 2$s$, and Eu 5$p$ core levels. The Eu 4$f$ lineshape in all the films is very characteristic for a Eu$^{2+}$ system. The multiplet structure typical for Eu$^{3+}$ is not visible. One can also observe that the O 2$p$ spectrum at 6–10 eV binding energy for YSZ is converted into the O 2$p$ valence band at 4–7 eV typical for EuO. All these demonstrate that only EuO has been formed, free from Eu$_2$O$_3$ or Eu$_3$O$_4$ contaminants. This also means that YSZ can only oxidize Eu into the 2+ state, and definitely not into the 3+.

We have also carried out take–off angle–dependent XPS experiments on the films. Figure 5 shows the Zr 3$d$ and Eu 4$d$ core level XPS spectra of one of the EuO films of Fig. 4 (a) collected at grazing emission, i.e., $\Theta = 70^\circ$ with respect to the surface normal, and at normal emission. One can clearly see that the Eu signal is not significantly reduced but the Zr signal has almost disappeared in the grazing–emission geometry. Since grazing emission means more surface sensitivity, this result confirms not only that the EuO film is on top of the YSZ substrate with negligible intermixing of the cations but also that the film is closed and flat.

Figure 6 depicts the Zr 3$d$ and Eu 4$d$ core level spectra
clearly observe that the Zr 3d signal is getting smaller when the oxygen pressure is increased, indicating that the thickness of the EuO film becomes larger. The line-shapes of the Eu 4d and Eu 4f levels are those of divalent Eu for pressures up to $12\times10^{-8}$ mbar. For a pressure of $12\times10^{-8}$ mbar or higher, however, the Eu 4d and Eu 4f spectral shapes start to change and show characteristics which indicate the presence of trivalent Eu. Apparently, for a Eu flux rate of 8.2 Å/min and substrate temperature of 400°C, $10–12\times10^{-8}$ mbar is the critical oxygen pressure below which EuO films can be made on YSZ (001) free from any Eu$_2$O$_3$ or Eu$_3$O$_4$ type of impurity phases.

We have also measured the photoemission spectra of these EuO films under grazing take-off angle conditions. Again, the Eu 4d core level and Eu 4f valence band spectra are all 2+ as long as the oxygen pressures in the MBE chamber are below the critical 10–12$\times10^{-8}$ mbar value. One example is shown in Fig. 7, where an oxygen pressure of $4\times10^{-8}$ mbar was used. This figure demonstrates that also the surface region of the films is free from Eu$_3^+$ species. It is interesting to compare the grazing with the normal emission spectra and also with the spectra displayed in Fig. 5. One can clearly see in Fig. 7 that the Zr 3d signal is very much suppressed in grazing emission, even more suppressed than in the grazing emission spectrum of Fig. 5. This shows that the EuO films grown with the supply of oxygen in the MBE chamber are thicker, a not so surprising and yet very consistent observation since without oxygen we have found that the EuO film growth is limited to five to six monolayers.

The following picture can now be drawn about the initial stages of growth of EuO film on YSZ (001). In case there is no oxygen in the MBE chamber, YSZ is supplying all the oxygen required to form EuO. The film is per-
grown at 400°C and the surface normal. The film was perfectly free from Eu\(^{3+}\) species. In case oxygen is present in the MBE chamber, YSZ is supplying only the amount of oxygen that is needed to complete the formation of EuO. This also explains why the growth rate of the first five to six monolayers is determined only by the Eu flux rate and is totally independent of the supply of oxygen pressure in the MBE chamber, see Fig. 2. It is important that the pressure is kept below the critical value of 8.2 Å/min Eu flux rate for 10 min.

FIG. 6: (Color online) (a) Zr 3d – Eu 4d core level XPS spectra and (b) Zr 4p – O 2s - Eu 5p core level and O 2p – Eu 4f valence band XPS spectra of EuO films on YSZ (001), grown at 400°C with a 8.2 Å/min Eu flux rate for 10 min. The spectra were collected at normal emission. From top to bottom: EuO films grown under oxygen pressures of 16, 12, 10, 8, and 4 × 10\(^{-8}\) mbar, and clean YSZ substrate.

FIG. 7: (Color online) Take-off angle dependence of the Zr 3d – Eu 4d core level XPS spectra of a EuO film on YSZ (001). Top: grazing emission, i.e., Θ = 70° with respect to the surface normal. Bottom: normal emission. The film was grown at 400°C under a 4 × 10\(^{-8}\) mbar oxygen pressure and a 8.2 Å/min Eu flux rate for 10 min.

10–12×10\(^{-8}\) mbar as we will discuss in more detail in Sec. IV.

Based on the comprehensive set of RHEED, LEED, and XPS data, including the RHEED intensity oscillations, we have now demonstrated that EuO thin films can be grown epitaxially in a layer–by–layer fashion with good control of its chemical state. The supply of oxygen from the YSZ does not do any harm, and in fact, it can be utilized as a verifying method to calibrate the Eu flux rate accurately, e.g., the 8.1–8.2 Å/min from the quartz crystal monitor corresponds to the growth of one monolayer EuO per 25 s.

IV. SUSTAINED GROWTH OF EuO ON YSZ

Having shown that the initial stages of growth of EuO on YSZ (001) can be made quite perfect, we now investigate whether thicker EuO films can be prepared while keeping the epitaxy and, especially, the layer–by–layer growth mode. We therefore have grown films for longer deposition times, e.g., between 100 and 200 min, using a series of finely intervalled pressures for the oxygen, e.g., 4, 8, 10, 12, and 16 ×10\(^{-8}\) mbar. The Eu flux rates were kept at 8.0–8.3 Å/min. The resulting thickness of the films varies between roughly 300 and 800 Å as will be discussed later. The RHEED and LEED results are plotted in Fig. 8. One can clearly see that excellent epitaxial growth has been achieved for (a) 4, (b) 8, and (c) 10 ×10\(^{-8}\) mbar oxygen pressures. For 12 ×10\(^{-8}\) mbar (d) or higher pressures, however, the appearance of additional spots in the RHEED indicates that the surface structure starts to change, and the absence of a pattern in
The LEED electron energies were 369, 368, 370, and 266 eV, respectively. The LEED photographs were taken at electron beam energies of 368–370 eV since lower energies did not provide stable patterns due to charging. The LEED even suggests appreciable surface roughness. The LEED photographs were taken at electron beam energies of 368–370 eV since lower energies did not provide stable patterns due to charging.

As discussed in Sec. III, the initial five to six oscillations of the specular reflected RHEED beam intensity are a unique feature for the initial stages of EuO growth on the YSZ (001). These initial oscillations do always occur, i.e., independent of the oxygen pressure in the MBE chamber, unless the pressure exceeds a critical value above which trivalent Eu species are formed. Remarkable is that no more oscillations can be observed beyond these five to six when growing thicker films. This is the case for a wide range of oxygen pressures. There is one exception: for a pressure of $10 \times 10^{-8}$ mbar, we were able to see further RHEED intensity oscillations. Figure 9 shows that the initial five to six oscillations are then followed by at least 50 more oscillations. It demonstrates that a layer–by–layer growth mode for EuO is possible during the sustained growth. Interestingly, the oscillation period during the sustained growth is similar and yet a little bit larger than during the initial stages of growth: 30 s against 25 s. Apparently, the oxygen pressure must be close to and yet a little less than the critical value in order to maintain the layer–by–layer growth mode, while the Eu flux determines the growth rate in the initial stages, i.e., $8.0–8.2 \, \text{Å/min}$ Eu flux corresponding to 25 s per EuO layer, it is the limited oxygen supply from the MBE environment which dictates the speed during the sustained growth, i.e., to 30 s per EuO layer.

To elucidate further the growth process, we have measured the thickness of the films using ex situ XRR measurements. Since EuO deteriorates rapidly under ambient conditions, the films need to be capped. To this end, an aluminum layer with a thickness of 20–40 Å has been evaporated on top of the EuO. This thickness turns out to be sufficient for the aluminum to be a good protective overlayer as will be discussed later. Figure 10 exhibits the XRR profile of the EuO film which was grown at an oxygen pressure of $4 \times 10^{-8}$ mbar for 200 min. The corresponding RHEED and LEED patterns of the EuO film are displayed in Fig. 8(a). From the period of interference fringes, we deduce that the thickness of the EuO film is about 350 Å.

Thickenss of the other films are also determined from their XRR profiles. The results are displayed in Fig. 11, where we plot the thickness against the product of oxygen pressure and total deposition time. For oxygen pressure
up to $12 \times 10^{-8}$ mbar we can observe a clear and direct linear relationship between them, strongly suggesting that the thickness is determined by the amount of oxygen incorporated. In other words, the growth is limited by the amount of oxygen made available. This in turn means that the Eu flux rate is higher than necessary and that the excess Eu must be re-evaporated into the vacuum. Figure 11 essentially confirms the distillation process needed to maintain good control of the stoichiometry as reported in our earlier studies by Steeneken et al.\cite{82} and Tjeng et al.\cite{83}.

We now investigate to what extent the growth conditions affect the magnetic properties of the EuO films using a SQUID magnetometer. The results are shown in Fig. 12. The films grown with 4, 8, 8.5, and $10 \times 10^{-8}$ mbar oxygen pressures all have a Curie temperature of 69 K with a magnetic moment of 7 $\mu_B$/f.u., as expected for a $4f^7$ system. The inset shows the field dependence of the magnetization at 5 K for the film grown with a $8 \times 10^{-8}$ mbar oxygen pressure. Here one can observe a hysteresis behavior with a saturation magnetization of $7\mu_B$. These results are in agreement with the RHEED and LEED results as displayed in Figs. 8(a)–8(c), in the sense that the proper ferromagnetic properties are always maintained as long as good epitaxial growth is also achieved. On the other hand, films grown with 12 and $16 \times 10^{-8}$ mbar have completely lost their ferromagnetic properties. It is remarkable that exceeding the $10 \times 10^{-8}$ mbar value just a little bit causes such a dramatic change. This very abrupt change is also consistent with the RHEED and LEED results as displayed in Fig. 8 (d). Considerable film roughness starts to develop for oxygen pressures higher than $10 \times 10^{-8}$ mbar. We would like to infer that having only a $T_C$ of about 69 K is not a sufficient characteristic to conclude that the film is homogeneous and stoichiometric. One also needs to establish that the film has a full saturation magnetization of $7\mu_B$. The measure-
oretical spectra for Eu$^{2+}$ (top) and Eu$^{3+}$ (bottom). It is clear that the Eu spectra are very similar to the theoretical spectrum for Eu$^{2+}$, meaning that the EuO films with 4, 8, and 10 $\times$ 10$^{-8}$ mbar oxygen pressures are completely free from Eu$^{3+}$ species. This in turn implies that an aluminum overlayer as thin as 20–40 Å works well to protect the EuO films against air, contrary to the claims made elsewhere that one needs very thick capping layers. We attribute this to the fact that the epitaxial growth of EuO on YSZ (001) yields such a smooth film so that a very thin aluminum film is sufficient to make a closed capping overlayer. The smoothness of the films as well as the complete absence of Eu$^{3+}$ impurities forms a good starting point for the fabrication of well-defined interfaces with other metals or oxide materials, thereby opening up new opportunities to study or even generate new phenomena related to interface physics.

We conclude that the critical oxygen pressure is around 10–12 $\times$ 10$^{-8}$ mbar for a 8.0–8.3 Å/min Eu flux rate. Only below this pressure one has the distillation process taking place so that good epitaxial growth can be achieved with the proper stoichiometry and ferromagnetic properties. Apparently, layer-by-layer growth can be obtained only if one is close to, but not exceeding, the critical pressure.

V. EPITAXIAL GROWTH OF EuO ON MgO

Having understood the growth process of EuO on YSZ (001), and having found the recipe to obtain excellent epitaxial growth of EuO on YSZ (001), we now turn our attention to the preparation of EuO on MgO (001). MgO as a substrate for EuO is interesting since several studies have reported good epitaxial growth despite the very large lattice mismatch of about 20%. The RHEED and LEED photographs of EuO films grown on MgO using different oxygen pressures are displayed in Fig. 14. EuO films with oxygen pressure below 10 $\times$ 10$^{-8}$ mbar all show excellent epitaxial growth. The relationship between the distances of the MgO and EuO streaks is consistent with the ratio of their lattice constants of 1.22. On the other hand, a higher oxygen pressure creates additional spots in the RHEED image and even causes the LEED pattern to disappear. Similar to the EuO on YSZ case, this indicates that 10–12 $\times$ 10$^{-8}$ is the critical value for the oxygen pressure above which one no longer gets crystalline and stoichiometric EuO.

Our attempts to observe RHEED intensity oscillations of EuO films grown on MgO (001) were yet unsuccessful. The RHEED streaks and specular spot suddenly disappeared right after the deposition of EuO has been initiated. In approximately 30 s, new streak lines appear, whose spacing conforms to the EuO lattice parameter. However, the specular spot was never recovered. A more detailed growth study is required to determine the growth process of EuO on MgO (001).

To investigate the quality of these EuO films in terms of their magnetic properties, we have performed SQUID measurement for films grown with oxygen pressures of 8, 10, and 12 $\times$ 10$^{-8}$ mbar. The results are shown in Fig. 15. Similar to the EuO on YSZ case, the films that were grown below 10 $\times$ 10$^{-8}$ mbar have a Curie temperature of 69 K with a magnetic moment close to $7\mu_B$/f.u. Conversely, the film grown under a higher oxygen pressure has completely lost its ferromagnetic character. The small peak at roughly 5 K indicates the typical antiferromagnetic ordering temperature of Eu$_2$O$_4$, meaning that the dramatic loss of the ferromagnetism for the growth using slightly above the critical oxygen pressure is due to the presence of Eu$^{2+}$ species.

FIG. 14: (Color online) Top panels: RHEED photographs of EuO films on MgO (001) grown at 400°C with 8.1–8.2 Å/min Eu flux rates under oxygen pressures of – from left to right – (a) 4, (b) 8, (c) 10, (d) 12, and (e) 16 $\times$ 10$^{-8}$ mbar. The deposition times were all 100 min. The RHEED electron energy was 20 keV with the beam incident along the [100] direction. Bottom panels: corresponding LEED photographs. The LEED electron energy was set at 360 eV.

FIG. 15: (Color online) Temperature dependence of the magnetization of epitaxial EuO films on MgO (001) grown at 400°C with a 8.1 Å/min Eu flux rate under various oxygen pressures as indicated. The small magnetization contribution from the substrate has been subtracted. The applied magnetic field was 1000 G.
VI. CONCLUSION

We have successfully grown epitaxial and highly stoichiometric EuO films on YSZ (001). The initial stages of growth involve the limited supply of oxygen from the YSZ substrate, but the EuO stoichiometry can still be well maintained. We have also observed RHEED intensity oscillations during the sustained stages of growth, which demonstrate that the layer–by–layer growth mode can be achieved for EuO films on YSZ (001). The EuO films were sufficiently smooth so that capping with a thin layer of aluminum enabled ex situ experiments free from trivalent Eu species. The excellent epitaxial growth of EuO films is always accompanied by equally excellent ferromagnetic properties: a saturation magnetization of 7\(\mu_B\) and a \(T_C\) of 69 K. We have also confirmed that the use of the Eu-distillation process during the MBE-assisted growth enables the consistent achievement of stoichiometry. The layer–by–layer growth, the smoothness of the film, and the excellent stoichiometry provide an excellent basis for the fabrication of well-defined interfaces for further studies.

VII. ACKNOWLEDGMENTS

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39 R. W. Ulbricht, A. Schmehl, T. Heeg, J. Schubert, and D. G. Schlom, Appl. Phys. Lett. 93, 102105 (2008).
40 R. P. Panguluri, T. S. Santos, E. Negusse, J. Dvorak, Y. Idzerda, J. S. Moodera, and B. Nadgorny, Phys. Rev. B 78, 125307 (2008).
41 N. J. C. Ingle and I. S. Elfimov, Phys. Rev. B 77, 121202(R) (2008).
42 R. P. Ingel and D. Lewis III, J. Am. Ceram. Soc. 69, 325 (1986).
43 M. Yashima, S. Sasaki, M. Kakhana, Y. Yamaguchi, H. Arashi, and M. Yoshimura, Acta Cryst. B 50, 663 (1994).
44 V. E. Henrich and P. A. Cox, The Surface Science of Metal Oxides (Cambridge University Press, 1994). Table 2.2.
45 N. Q. Minh, J. Am. Ceram. Soc. 76, 563 (1993).
46 B. T. Thole, G. van der Laan, J. C. Fuggle, G. A. Sawatzky, R. C. Karnatak, and J.-M. Esteva, Phys. Rev. B 32, 5107 (1985).
47 J. B. Goedkoop, B. T. Thole, G. van der Laan, G. A. Sawatzky, F. M. F. de Groot, and J. C. Fuggle, Phys. Rev. B 37, 2086 (1988).