Off-axis sputter deposition of ZnO films on c-sapphire substrates by utilizing nitrogen-mediated crystallization method

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Abstract. High-quality epitaxial ZnO films on c-plane sapphire substrates have been obtained by utilizing an off-axis sputtering configuration together with buffer layers prepared via nitrogen-mediated crystallization (NMC). The role of NMC buffer layers is to provide high density of nucleation site and, thus, to reduce the strain energy caused by the large lattice mismatch (18%) between ZnO and sapphire. The NMC buffer layers allow two-dimensional growth of subsequently grown ZnO films, being particularly enhanced by employing an off-axis sputtering configuration in which the substrate is positioned out of the high-energy particles, such as negative oxygen ions originating from the targets. As a result, ZnO films with smooth surfaces (root-mean-square roughness: 0.76 nm) and a high electron mobility of 88 cm²/V·s are fabricated. Photoluminescence spectra of the ZnO films show strong near-band-edge emission, and the intensity of the orange-red defect emission significantly decreases with increasing horizontal distance between the target and the substrate. From these results, we conclude that off-axis sputtering together with NMC buffer layers is a promising method for obtaining high-quality epitaxial ZnO films. © The Authors. Published by SPIE under a Creative Commons Attribution 3.0 Unported License. Distribution or reproduction of this work in whole or in part requires full attribution of the original publication, including its DOI. [DOI: 10.1117/1.OE.53.8.087109]

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1 Introduction

ZnO is a remarkable multifunctional material with a distinctive property set and a huge range of existing and emerging applications, such as varistors, transparent conducting electrodes, thin-film transistor, and gas sensors.1–10 ZnO has recently been recognized as a candidate for high-performance ultraviolet light-emitting diodes (LED) and laser diodes (LD) that take advantage of its high exciton binding energy of 60 meV,11–13 which is much larger than that of the commercial LED material GaN (25 meV). Furthermore, due to the material abundance and low toxicity of ZnO, the replacement of GaN will bring a significant reduction in the cost as well as the reduction of damage to human health and the environment. Since such devices require single crystals with low defect density, ZnO-based LEDs have been fabricated on lattice-matched but expensive substrates, such as bulk ZnO and ScAlMgO4 substrates.11,12 Furthermore, these ZnO films have been fabricated by a molecular beam epitaxy method that needs an ultrahigh vacuum of 10⁻⁷ Pa. Even though the material cost of ZnO itself is much lower than that of GaN, it is essential to fabricate single crystalline ZnO films on cost-effective substrates using a mass-productive method for the replacement of GaN. C-plane sapphire has a great potential as an epitaxial substrate for ZnO because of its low cost and availability in large-area wafers. Due to a large lattice mismatch of 18%, however, ZnO films prepared on sapphire substrates have large crystal mosaics, high residual carrier concentrations, and low mobility, all of which make optoelectronic applications challenging.14

One of the most promising means to improve the crystal quality of ZnO films is to prepare buffer layers prior to the crystal growth. Various types of ZnO buffer layers have been reported. Khranovsky et al. fabricated highly oriented ZnO (002) films on c-sapphire substrates by incorporating low-temperature (LT) ZnO buffer layers prepared by metal-organic chemical vapor deposition.15 In that work, x-ray diffraction (XRD) 2θ–ω patterns showed only the (002) diffraction peak, whereas ZnO films without the buffer layers showed the (100) and (101) diffraction peaks aside from the (002) peak. Similarly, Nakamura et al. fabricated ZnO films by pulsed-laser deposition on LT buffer layers on c-plane substrates.16 With buffer layers deposited at 500°C, the full-width half-maximum (FWHM) of the (002) rocking curve decreased to 0.09 deg from 0.2 to 0.3 deg, indicating that well-aligned ZnO films were fabricated. These results certainly demonstrate that buffer layers improve the crystal quality of ZnO films. Single crystalline ZnO films with desired properties for homojunction LED/LD have, however, not been obtained yet. The idea of LT buffer layers was first reported by Amano et al., having been practically used for fabrication of GaN-based LED on sapphire substrates, where the lattice mismatch between GaN and sapphire is 16%.17 The role of the LT buffer layers is to provide high density of the nucleation site and smooth surface and, thus, to reduce the interfacial free energy between GaN and the substrates that come from the large lattice mismatch. The nucleation control of the LT buffer layers is carried out by depositing amorphous films at a temperature sufficiently lower than the crystal growth temperature and annealing them. Such buffer
layers enhance the lateral growth of subsequently grown GaN; as a result, single crystalline GaN films are fabricated. In the case of ZnO, however, LT buffer layers have limited effects on the improvement of the crystal quality because the amorphous phase of ZnO is difficult to obtain even at room temperature because of the low crystallization temperature. This is apparent when a sputtering method is employed for film deposition, where the high energy of impingement of the sputtered species as well as negative oxygen ions and recoil argons exist.

We have recently reported a new type of buffer layer fabricated via nitrogen-mediated crystallization (NMC), which enables us to make high-quality ZnO films. During epitaxial growth of ZnO films on unbuffered c-plane sapphire substrates, highly strained two-dimensional (2-D) layers are initially formed due to the large lattice mismatch between ZnO and sapphire. On the 2-D layers, subsequent growth of three-dimensional (3-D) columnar grains with poor alignment occurs so that the strain energy stored in the film is released (Stranski-Krastanov mode). In contrast, our NMC buffer layers possess a high surface concentration of discrete ZnO islands on the sapphire substrates even at the very early stage of crystal growth because nitrogen atoms introduced into the deposition atmosphere disturb the growth of ZnO islands. Since such buffer layers have good in-plane alignment, ZnO crystals that originate from the grains of the buffer layers are expected to be grown laterally and to be coalesced if secondary nucleation is suppressed.

In this study, we demonstrate the epitaxial growth of ZnO films on NMC-ZnO buffered c-plane sapphire substrates, where radio frequency (RF) magnetron sputtering, which has great advantages in terms of mass production, is employed. In magnetron sputtering, there are many high-energy particles, such as negative oxygen ions and recoil Ar, which cause defect formation in ZnO films. Since both particles are accelerated in the cathode sheath, the flux of the energetic particles can be reduced by taking the off-axis configuration where the substrate is positioned out of the high-energy particles originating from the targets. Here, aiming at realizing high-quality ZnO films with a low defect density, the effects of off-axis configuration together with NMC buffer layers are investigated through the analysis of the properties of ZnO films fabricated at various substrate/target configurations.

2 Experimental

2.1 Fabrication of NMC Buffer Layers

NMC-ZnO buffer layers were deposited on 10 × 10 mm² c-plane sapphire substrates by RF magnetron sputtering. The schematic view of the configuration of the magnetron sputtering system is shown in Fig. 1. For preparation of NMC buffer layers, the substrate holder was rotated at 10 rpm, the rotation axis of which was set at x = 45 mm, in order to fabricate the films with uniform distribution of the film properties. The target-substrate vertical distance was 74.5 mm. The supplied RF power was 100 W and the deposition temperature was 700°C. Ar – N₂ mixed gas with a N₂/(Ar + N₂) flow rate ratio of 0.08 was used and the total gas pressure was 0.30 Pa. The thickness of the buffer layers was 10 nm, confirmed by x-ray reflectometry (D8 Discover, Bruker AXS K.K., Kanagawa, Japan).

2.2 Fabrication of ZnO Films on NMC Buffer Layers

ZnO films were deposited on NMC-ZnO buffer layers by RF magnetron sputtering in the geometry as shown in Fig. 1. Here, the substrate holder was not rotated in order to study the effects of target/substrate configuration. The vertical distance from the target center to the substrate was 74.5 mm and the horizontal distance from the center of the target to the substrate was 0 to 60.6 mm. The substrate temperature was kept at 700°C and no postannealing of ZnO was performed. ZnO ceramic targets (2 in. in diameter) were used and the supplied RF power was 60 W. Ar – O₂ mixed gas with an O₂/(Ar + O₂) flow ratio of 0.10 was used, and the total deposition pressure was 0.70 Pa. It has been reported that high-energy particles impinging the growth surfaces, such as negative oxygen ions and recoil Ar, originate from the erosion race track, the state of which plays an important role in crystal growth, especially at such a low gas pressure. In this study, the erosion depth was in the range between 1 and 2 mm. From scanning electron microscopy observation, it was confirmed that the film thickness at the center of each sample was 1 μm, where the thickness gradient across one sample was < ± 6%. For comparison, ZnO films deposited directly on sapphire substrates (without buffer layers) were fabricated under the same deposition conditions. The crystallinity of all the films was evaluated by XRD using a four-circle texture diffractometer and a Cu Kα source (λ = 0.154 nm). The surface morphology was investigated by tapping mode atomic force microscopy (AFM). Optical properties of the thin films were characterized by photoluminescence (PL) spectra excited by 325-nm He-Cd laser at room temperature. Electrical properties were evaluated by Hall-effect measurement by using the four-point van der Pauw configuration at room temperature.

![Fig. 1 Schematic view of the configuration of the magnetron sputtering system.](https://www.spiedigitallibrary.org/journals/Optical-Engineering/v53i8/087109-2/image1.png)
3 Results and Discussion

3.1 Morphology and Electrical Properties of ZnO Films on NMC Buffer Layers

Figure 2(a) shows AFM images of ZnO films deposited on NMC buffer layers, where \( x \) denotes the sample position shown in Fig. 1. For comparison, AFM images of ZnO films deposited directly on the sapphire substrates are shown in Fig. 2(b). For ZnO films on NMC buffer layers, as \( x \) increases, an apparent change in the growth mode from 3-D island growth to 2-D lateral growth of trapezoid islands is observed, whereas there is no significant change in the growth mode for ZnO films without buffer layers. Similarly, the surface roughness, which is shown in Fig. 3, varies with the sample position for ZnO films with buffer layers, while ZnO films without buffer layers possess a large surface roughness of 20 to 30 nm, independent of \( x \) (except for \( x = 90 \) mm). For \( x \) > 55 mm, coalesced trapezoid islands are observed for ZnO films with buffer layers, and the root-mean-square (RMS) roughness reaches the minimum value of 0.76 nm at \( x = 65 \) mm, which is significantly smaller than the RMS roughness of 11 nm for the films deposited at \( x = 30 \) mm. The change in the growth mode results in a change in the electronic properties. Figure 4 shows electron Hall mobility and carrier density at room temperature as a function of sample position \( x \). The electron mobility of ZnO films with buffer layers monotonically increases with increasing \( x \) and reaches 88 cm²/V·s at \( x = 90 \) mm, whereas the mobility is 66 cm²/V·s for the films deposited at \( x = 30 \) mm. The residual carrier density of ZnO films with buffer layers decreases with increasing \( x \), indicating the decrease in the crystal defects at a large \( x \). This improvement in the surface morphology as well as in the electrical properties is attributed to both the NMC buffer layers and off-axis sputtering geometry. The buffer layers enhance the lateral growth of ZnO islands because of the low interface energy between ZnO and the NMC buffer layers. Besides, off-axis sputtering (\( x \) > 55 mm) suppresses secondary nucleation on the growth surfaces because of the reduced high-energy flux of negative oxygen ions and/or recoil Ar (Ref. 31) as well as the enhancement in the surface migration of sputtered particles.32,33 In our experimental conditions, such high-energy particles originating from the target erosion are accelerated throughout the cathode sheath of \( \sim 100 \) V and impinge the growth surface with little or no loss of energy because of a low gas pressure of 0.7 Pa.30 The eroded target with a depth of a few millimeters in our experiments provides the flux of the high-energy particles with a somewhat broad angular distribution; the particles bombard the growth surface facing the whole area of the target (\( x = 0 \) to 45 mm).30,34 This bombardment by high-energy particles can cause secondary nucleation,31 crystal-defect formation, and surface roughening, which result in 3-D island crystal growth with large RMS roughness observed at \( x < 50 \) mm even when the buffer layers are utilized. On the other hand, at \( x > 55 \) mm, ZnO films on NMC buffer layers grow in 2-D mode, which is attributed to the enhancement in the surface mobility of sputtered particles along with the reduced flux of high-energy particles. In an off-axis configuration, it has been reported that the surface mobility of sputtered particles is higher than that in on-axis configuration due to increase in the momentum component parallel to the growth surfaces.32,33 The higher mobility of sputtered particles as well as the reduced flux of high-energy particles are considered to lead to the lateral growth of ZnO crystals and the smooth surfaces observed at \( x = 50 \) to 65 mm. When \( x \) exceeds 65 mm, however, the RMS roughness increases with increasing \( x \) (Fig. 3). Since not only the flux of high-energy particles but also the kinetic energy of the sputtered species is reduced at a large \( x \), surface migration of Zn and O atoms, which enhances lateral growth and coalescence of the islands, is suppressed, resulting in films with a higher density of large-size pit defects as well as rougher surfaces.35,36

3.2 Photoluminescence of ZnO Films on NMC Buffer Layers

Because the sample position \( x \) was found to have a significant influence on the crystal growth mode of ZnO, especially

![AFM images of ZnO films](https://www.spiedigitallibrary.org/journals/Optical-Engineering/2014/087109-3/fig2a.png)

**Fig. 2** Atomic force microscopy (AFM) images of ZnO films fabricated at various sample positions \( x \).
(a) ZnO films deposited on nitrogen-mediated crystallization (NMC) buffer layers fabricated by NMC method. (b) ZnO films deposited directly on c-plane sapphire substrate.
when using NMC buffer layers, we next studied the PL spectra from ZnO films, which provides an insight into the nature of the crystal defects. Figure 5 shows the room-temperature PL spectra of ZnO films deposited on NMC buffer layers at various sample positions $x$. All spectra have a narrow UV peak at 380 nm. This near-band-edge emission is attributed to the recombination of free excitons and donor-bond excitons. As $x$ increases, the intensity of near-band-edge emission increases, indicating the decrease of nonradiative recombination centers that originate from crystal defects, such as dislocations, zinc vacancy related complex, and surface/interface states. Recently, Matsumoto et al. reported that nonradiative recombination centers are also attributed to crystal grain boundaries, the number of which increases with increasing grain density. In our ZnO films, the grain boundaries are clearly seen when the films were fabricated at $x < 55$ mm; by contrast, there is no grain boundary observed at a larger $x$ (Fig. 2), suggesting that the decrease in the grain boundary may cause an increase in the near-band-edge emission intensity. On the other hand, the intensity of the broad orange emissions centered at 650 and 750 nm, which are mainly related to deep-level emissions through oxygen interstitial, decreases at a large $x$. This is because the flux of energetic negative oxygen ions, which cause the formation of oxygen interstitial in ZnO, decreases with increasing $x$.

From these results, we conclude that by utilizing off-axis sputtering, and thereby suppressing the flux of energetic particles, such as negative oxygen, the density of lattice defects, including interstitial atoms and grain boundaries, can be reduced.

### 3.3 XRD Results

Although a variation of sample position $x$ brings a significant change in the crystal growth mode as well as the surface roughness of ZnO, we clarified that fluctuation of the crystal orientation in both in-plane and out-of-plane directions is independent of sample position $x$ when NMC buffer layers are utilized. Figure 6(a) shows FWHM of the (0002) symmetric rocking curves for ZnO films with and without NMC buffer layers as a function of the sample position $x$. Here, all ZnO films are epitaxially grown on c-sapphire substrates, where the epitaxial relationship between ZnO and sapphire is $\frac{1}{2}[0001]_{\text{ZnO}} // \frac{1}{2}[0001]_{\text{sapphire}}$ and $\frac{1}{2}[10-10]_{\text{ZnO}} // [11-20]_{\text{sapphire}}$. ZnO films with high out-of-plane alignment are obtained by using NMC buffer layers, independent of the sputtering configuration. FWHM of the films deposited on the buffer layers ranges from 0.089 to 0.11 deg, which is much smaller than the value of 0.18 to 0.43 deg for the films deposited without buffer layers. Furthermore, as can be seen in
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Naho Itagaki is an associate professor at Kyushu University, Japan. She received her BS, MS, and PhD degrees in plasma physics from Kyushu University in 1998, 2000, and 2003, respectively. She is the author of more than 60 journal papers and has more than 40 patents. Her current research interests include oxide and oxynitride semiconductors for advanced optoelectronic devices.

Biographies for the other authors are not available.