Optical and structural properties of ZnO films deposited on GaAs by pulsed laser deposition

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ZnO films were synthesized on GaAs substrates at different growth conditions by pulse laser deposition. High-purity (99.999\%) oxygen was used as the ambient gas. The pressure of the ambient oxygen gas for ZnO film growth was varied from 20 to 50 mTorr, and the growth temperature from 300 to 450 °C. ZnO films showed very strong bound exciton peaks located between 3.37 and 3.35 eV. The full width at half maximum of the bound exciton peak is less than 5 meV. These results indicate ZnO films on GaAs substrates can be used for optical devices such as light-emitting diodes. The other significant properties of textured ZnO films on GaAs substrates are described. © 2000 American Institute of Physics. [S0021-8979(00)2013-2]

I. INTRODUCTION

ZnO films have been studied extensively for use in piezoelectric and waveguide devices. Recently, ZnO films have become attractive candidates as substrates for GaN films because both materials possess the same crystal structure (wurtzite) with small lattice mismatch. GaN is known as a good material for fabrication of optical devices such as blue light emitting diodes (LEDs) and blue laser diodes (LDs). Nakamura has used GaN in an effort to increase the average lifetime of blue LDs toward a goal of 10000 h at room temperature.\textsuperscript{1}

This research was motivated by the possibility that ZnO could be a good candidate material for optical devices such as blue LEDs. This supposition is based on the fact that the crystal structure of ZnO is the same as that of GaN, and that the optical properties are similar to each other.\textsuperscript{2} ZnO has other physical features that make it attractive. For example, as a wide band-gap semiconductor, ZnO is a relatively hard material. The strength of the Zn–to–O bond is larger than that of the Ga–to–N bond. Their difference in bonding strength can lead to significantly different results for p-type doping. ZnO has a melting temperature of about 2000 °C. Thus, ZnO is sufficiently stable that it can withstand the high temperature annealing and treatment processes associated with doping and forming ohmic contacts. The hardness of ZnO, its resistance to mechanical stress, and its high melting temperature are features that could conceivably expand the lifetime of devices such as blue LEDs and blue LDs fabricated with this material, and for some applications it is possible that ZnO could be superior to GaN.

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II. FILM GROWTH

ZnO films were synthesized on (001)-GaAs substrates by PLD using an ArF excimer laser. Experiments were performed in a high-vacuum chamber maintained at $8 \times 10^{-7}$ Torr by a turbopump. The focused laser radiation illuminated a rotating ZnO target at an incident angle of 40°. The target was 99.999\% pure polycrystalline ZnO. The target-to-substrate distance was 7 cm.
ZnO films were in the range 0.7–1.1 μm. The results can be understood by considering the energy density of the focused laser beam was roughly 80 mJ, the repetition rate was 20 Hz and the power per pulse was 80 mJ. While maintaining laser illumination of the target. The laser was immediately after the target had been preablated. The growth chamber was filled with argon gas at the time laser illumination was begun to create a (Zn,O) plasma. The use of an ambient gas of sufficient pressure helps avoid particulate deposition on the film. Keeping a designated ambient gas pressure, argon gas was gradually replaced with oxygen gas while maintaining laser illumination of the target. The laser repetition rate was 20 Hz and the power per pulse was 80 mJ. The energy density of the focused laser beam was roughly estimated to be 2 J/cm² at the target. The thicknesses of the ZnO films were in the range 0.7–1.1 μm.

III. MEASUREMENTS AND RESULTS

ZnO films were measured by XRD to study crystal structure. The x-ray θ–2θ rocking curve for a ZnO film on GaAs is shown in Fig. 1. The strong intensity of the (0002)-ZnO peak indicates a self-texture preference in the c-axis direction, even on a GaAs substrate. X-ray in-plane measurements show that ZnO films deposited by PLD have hexagonal structures. Figure 2 shows the variation of the full width at half maximum (FWHM) of the x-ray θ-rocking curve for the (0002)-ZnO peak with oxygen gas pressure and the substrate temperature. The results can be understood by considering the (Zn,O) plume size, the reactive oxygen density, the density of particulates generated at the target that reach the substrate, the surface mobility, and As diffusion into the newly formed ZnO layers. Oxygen ambient molecules can be transformed into reactive elements through inelastic collisions within the initial (Zn,O) plasma that is created from the ZnO target by the focused laser beam. ZnO films, regardless of the growth technique used, possess some oxygen vacancies. Thus, it is very important for ZnO film growth that the amount of available reactive oxygen created from the ambient oxygen gas be sufficient to help reduce the number of oxygen vacancies. It is also known that the number of particulates arriving at the substrate is decreased as the pressure of an ambient gas increases. Deposited particulates in a ZnO film degrade film quality. For these reasons, generally speaking, the crystal quality is improved as the ambient gas pressure increases.

At sufficiently high ambient gas pressures, however, the (Zn,O) plasma will not reach the substrate and the ZnO film will not be well crystallized. The plasma density that might be expected from a Beer’s law behavior is given by
\[ I = I_0 \exp(-P\sigma x), \]
where \( I_0 \), \( P \), \( \sigma \), and \( x \) are the initial plasma density at the target, the oxygen ambient gas pressure, the total cross section for plasma-to-oxygen collisions, and the distance from the target, respectively. The density of available source elements for ZnO film growth might be exponentially decreased as the ambient gas pressure increases, and the relative density ratio for the Zn and O elements may not be good for high-quality film growth at a high ambient pressure such as 50 mTorr. The plume size \( R \) is related to the ambient gas pressure and to the laser-pulse energy \( E \). It can be roughly calculated by
\[ R \approx (E/P)^{1/3}, \]
where \( \gamma \) is the ratio of specific heats of the elements in the plume. If the substrate is located beyond the plume size \( R \), the adhesion coefficient for source elements that arrive at the substrate is drastically decreased. Therefore, the crystal quality of ZnO becomes worse as the ambient gas pressure increases beyond an optimal pressure. The ambient oxygen gas pressure should be neither too low nor too high. An acceptable value is 35 mTorr for ZnO film growth on GaAs.

Arsenic atoms from the GaAs substrate can evaporate

FIG. 1. The x-ray θ–2θ rocking curve for a ZnO film on GaAs. A (0002)-ZnO peak indicates the c-axis oriented growth direction.

FIG. 2. (a) and (b) show the effects of growth temperature and oxygen gas pressure on the crystalline quality of ZnO film grown by PLD. The FWHM values of the x-ray θ rocking curves are for the (0002)-ZnO peaks. The pressure of an ambient oxygen gas was fixed at 35 mTorr for (a) and the substrate temperature was fixed at 350 °C for (b). The best ZnO film was synthesized at 350 °C and 35 mTorr, with a FWHM value of 1.39° for the θ rocking curve.
and diffuse into the ZnO layers at a sufficiently high substrate temperature, such as 450 °C. Therefore high substrate temperatures are to be avoided for ultrapure ZnO film growth on GaAs substrates, unlike that for ZnO film growth on sapphire substrates. On the other hand, the surface mobility for (Zn, O) elements at the substrate may be too low for high-quality film growth at low temperatures such as 300 °C.

Even though these interpretations require further study, these relations are implicitly indicated by the results in Fig. 2. The best films, with a FWHM value of 1.39°, were synthesized with growth parameters of 350 °C substrate temperature and 35 mTorr ambient oxygen gas pressure.

Film surface morphologies were studied using atomic force microscopy (AFM) measurements. AFM measurements were performed in air using a model Nanoscope IIIa SPM manufactured by Digital Instruments, Inc. All samples were scanned over areas ranging from 500×500 nm² to 10×10 μm², and at several different locations on the film surface. ZnO films synthesized in an ambient pressure of 20 mTorr oxygen showed a few particulates, with a density of about 1/μm². It was very difficult to find any particulates in ZnO films grown at ambient gas pressures higher than 20 mTorr, which is a consistent result with those for ZnSe film growth. For growth at 350 °C, with increased ambient gas pressure, the surface became smoother, and the microcrystalline (grain) size became smaller.

Surface morphologies vary significantly among ZnO films grown at different temperatures. Figure 3 shows AFM images from those samples grown at different temperatures but at the same ambient pressure, 35 mTorr of oxygen. They show the effect of growth temperature on film morphology. In particular, surface roughness and grain size are much different. The ZnO films grown at 350 °C show an average roughness of 9.4 nm (root-mean-square) within the scanned area, 1×1 μm². Their grain sizes are the smallest, though the samples at 450 °C did have the smoothest surfaces. The grain sizes in all samples ranged from 40 to 300 nm. The
growth modes are probably different at different temperatures due to the influence of the substrate.

The optical properties of ZnO films were characterized at 20 K by photoluminescence (PL) spectroscopy. They were excited with an argon ion laser operating at 351.1 nm wavelength. As shown in Fig. 4, the PL spectra for a ZnO film grown at 350 °C and 35 mTorr oxygen are characterized by significant spectral peaks located at 3.360, 3.356, 3.321, 3.321 eV, and 3.217 eV. The dominant and narrow peaks located at 3.360 and 3.356 eV are the near band-edge (NBE) emissions of excitons bound to donors (DX), which are consistent with the results reported previously.12,13 The FWHM values of each peak are less than 5 meV. The peaks located at 3.288 eV might be identified as an LO phonon replica.13 Other interesting peaks at 3.217 eV may be due to donor–acceptor pair transitions. For undoped ZnO films it is reasonable that the intensity of the DX NBE emission is stronger than that of the AX NBE emission because they are intrinsic n type due to the presence of oxygen deficiencies. On the other hand, there are additional broad peaks located at about 2.8 and 2.1 eV (not shown) which could conceivably be deep level emissions. The intensity of the peaks located at about 2.8 eV is very weak, and the intensity of the peaks at about 2.1 eV are comparable to the intensity of the DX NBE emission. For these ZnO films grown by PLD it was very difficult to detect any spectral peak contributions located at about 2.4 eV, the so-called green band;2 however, such contributions are sometimes (but not always) very strong for ZnO grown by the hydrothermal growth technique. The presence of the green band might possibly be correlated with deep levels created by hydrogen atoms. Such a correlation would explain why ZnO grown by PLD without use of hydrogen shows very weak peaks in the green band. These results indicate that further study is required to fully understand the optical properties of ZnO. NBE emissions from DX centers located at 3.360 and 3.356 eV can be separated into at least two peaks for each emission, as shown in Fig. 4.12 The peak at 3.365 eV might be the A band free exciton emission.12

Based on the fact that FWHM values from x-ray θ-2θ rocking curves are greater than 1°, it is rather surprising that the ZnO films in this study show very strong excitonic peaks in their PL spectra. The quality of the crystalline structure of a ZnO film on GaAs is less than, and not even comparable with, that of GaN on sapphire; however, a ZnO film shows very narrow and very strong excitonic peaks, being comparable with those observed for GaN. These particular features of ZnO may be the evidence that a ZnO film possesses high light radiation efficiency.

IV. CONCLUSION

ZnO films were synthesized on GaAs by PLD. Results from this work demonstrate that the PLD technique is a good method for growth of ZnO films. The grain size and the surface morphology of a ZnO film could be controlled by proper adjustment of the oxygen gas ambient pressure and the substrate temperature during growth. It is remarkable that ZnO films show very strong excitonic emission peaks, although the FWHM values of the x-ray θ-2θ rocking curves for their (0002)-ZnO peaks are much larger than 1°. A reasonable interpretation of the XRD and PL results supports the possibility that ZnO has a radiation efficiency value at least comparable to that of ZnSe or GaN. These properties make ZnO a plausible candidate material for short-wavelength visible or UV LEDs.

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