Structural and optical properties of indium-rich InGaN islands

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Indium-rich In\textsubscript{x}Ga\textsubscript{1-x}N islands (x \geq 0.87) grown by metalorganic chemical vapor deposition at growth temperatures ranging from 550-750 °C were investigated. With the increasing growth temperature, the InGaN dot size increases while the dot density decreases due to the enhanced surface migration of In/Ga adatoms at elevated temperatures. In addition, the composition of InGaN islands was also found to be varied with the growth temperature. At lower growth temperatures, a higher Ga content can be achieved due to the relatively lower migration ability of Ga adatoms. At higher growth temperatures, the deposited InGaN material tend to decompose into In-rich islands and a thin Ga-rich layer. Photoluminescence investigations show that these In-rich islands exhibit a near-infrared emission band. For samples grown at higher temperatures, a visible emission band was also observed. The visible emission band may arise from defect bands introduced during the formation of Ga-rich InGaN layer.

1 Introduction

Ternary InGaN alloys have been an important active material for nitride based optoelectronic devices [1], particularly for commercial light-emitting diodes (LEDs) and laser diodes operating in the blue-green and ultraviolet spectral ranges [2]. Recently, it has been shown that the energy gap of InN is around 0.7 eV [3, 4], rather than the previously accepted value of 1.9 eV. This finding makes In-rich InGaN alloys become a promising material for applications in the near infrared (NIR) spectral range.

In the last decade, In-rich nanostructures formed by local compositional fluctuations in InGaN alloys have been intensively investigated, because they were believed to act as “quantum dots” for localizing carriers and were considered to play a key role in the high radiative efficiency of InGaN-based LEDs [5, 6]. InGaN quantum dots formed via self-assembled island growth have also attracted much attention in the past few years [7–10]. However, the In\textsubscript{x}Ga\textsubscript{1-x}N dots reported to date are still in the Ga-rich side (x \leq 0.5), with typical emission wavelength in the blue-green range. On the In-rich side (0.5 < x < 1), the study of In\textsubscript{x}Ga\textsubscript{1-x}N dots with NIR emissions is still absent.

Previously, we have demonstrated that InN nanodots can be formed on GaN surface during the initial stage of heteroepitaxial growth using MOCVD. This kind of InN nanodots exhibit intense photoluminescence (PL) emissions in the NIR range and show pronounced quantum size effects [11, 12]. In this study, we further investigate In-rich In\textsubscript{x}Ga\textsubscript{1-x}N islands (with x > 0.87) grown by MOCVD at various temperatures. Surface morphologies, alloy compositions and PL emission properties of these In-rich In\textsubscript{x}Ga\textsubscript{1-x}N islands have been studied by atomic force microscopy (AFM), x-ray diffraction (XRD), PL measurements, near-field scanning optical microscopy (NSOM). The influences of growth temperature on island formation, alloy decomposition and their impacts on the optical properties of In-rich InGaN islands are discussed.

2 Experimental

The uncapped InGaN islands samples were grown on Sapphire (0001) by MOCVD using trimethylgallium (TMG), trimethylindium (TMI), and NH\textsubscript{3} as source materials. After nitridation of the substrate at 1120 °C, a thin GaN nucleation layer was first grown at 520 °C, followed by a 2-µm-thick undoped GaN buffer layer grown at 1120 °C. After the growth of the GaN
buffer layer, the substrate temperature was decreased to 550-750 °C to grow InGaN islands using modulated precursor injection schemes. The precursor flow rates were 1, 150 and 18,000 SCCM (SCCM denotes cubic centimeter per minute at STP) for the TMGa, TMIn and NH3, respectively. The details of the gas flow sequence have been reported in [12]. The only difference is that a 1 SCCM TMGa flow was supplied concurrently with the TMIn flow to grow InGaN alloys. During TMGa and TMIn flow periods, the NH3 background flow rate was controlled at 10,000 SCCM. PL measurements were carried out at $T = 10$ K using the 325-nm line of a He-Cd laser as an excitation source. The PL signals were analyzed by a 0.5 m monochromator and detected by either a photomultiplier tube or a cooled extended InGaAs detector (with a cutoff wavelength at 2.05 µm) for the visible and NIR spectral ranges, respectively. For NSOM measurements, a fiber tip was mounted on the tuning fork of an AFM system for simultaneous measurement of the surface morphology and emission signal using the illumination-collection mode. The tip-sample distance was kept at ~30 nm controlled by the shear-force feedback mechanism.

### 3 Results and discussion

Figure 1(a) shows the surface morphologies of InGaN islands grown at different growth temperatures ranging from $T_g = 600-725$ °C. With the increasing $T_g$, the average height and diameter of the InGaN islands increased from 24 to 114 nm and 75 to 410 nm, respectively; whereas the dot density decreased from $4.0 \times 10^{10}$ to $2.2 \times 10^7$ cm$^{-2}$. The dot density as function of $T_g$ is depicted in Fig. 1(b). For $T_g = 600-700$ °C, the decreasing dot density with the increasing $T_g$ can be attributed to the enhanced migration length of adatoms, yielding much larger and less dense islands at higher growth temperatures [12]. A rapid decrease in dot density for $T_g > 700$ °C can be seen and dropped eventually to zero at 750 °C. This can be attributed to the desorption of metallic In from the growing surface. We have also prepared a series of samples with InN islands grown at different $T_g$ under similar growth conditions. The density of InN islands as a function of $T_g$ was also shown in Fig. 1(b). A similar dependence of dot density on $T_g$ for both InGaN and InN islands can be seen, implying that the nucleation of InGaN islands is governed by the surface migration of In adatoms, rather than Ga or both.

Figure 2 X-ray diffraction curves (a) and the corresponding In compositions $x$ (b) of the InGaN islands grown at different temperatures.

Figure 2(a) shows XRD $\theta - 2\theta$ scans of the InGaN dot samples near the InN(0002) and GaN(0002) diffraction peaks. A diffraction curve of InN film grown on GaN was also included. For the InGaN samples, a broad diffraction peak corresponding to ternary InGaN islands was found to shift gradually toward the InN(0002) as $T_g$ was increased from 550 to 725 °C. For the sample grown at 750 °C, the InGaN peak disappeared, due to the complete desorption of In from the growing surface. The corresponding In content ($x$) of the In$_x$Ga$_{1-x}$N islands estimated using the Vegard’s law is depicted in Fig. 2(b). The In content was found to increase from 0.87 to 0.99 as $T_g$ was increased from 550 to 725 °C. This increasing trend is also a consequence of the
lower diffusion ability of Ga adatoms. At lower \( T_g \), the growth is under the kinetic-dominant condition so Ga incorporation at island sites could be more dominant. With the increasing \( T_g \), the growth conditions become closer to the thermodynamic conditions, which allows the Ga adatoms to select thermodynamically more stable sites and hence hinders the incorporation of Ga adatoms into the In-rich islands. The above mentioned mechanism suggests that the formation of In-rich InGaN islands is very likely to arise from phase separation, rather than SK growth. As can be seen from the case of \( T_g = 725^\circ \text{C} \), the \( \text{In}_x\text{Ga}_{1-x}\text{N} \) islands become highly In-rich, with \( x \) up to 0.99. Therefore, it can be inferred that most of the deposited Ga adatoms are very likely to be distributed among these In-rich islands, forming a thin Ga-rich layer.

To further confirm this assertion, we have also grown a sample with InGaN islands formed on AlGaN/sapphire (0001) at 725 \(^\circ\text{C}\). The sample was then investigated by Auger electron spectroscopy (AES) to distinguish the surface composition of In, Ga and Al. Fig. 3(a) and 3(b) are the AES results of surface composition measured on InGaN islands and on flat region without islands, respectively. As shown in Fig. 3(c), after 3 min \( \text{Ar}^+ \) sputtering (with the sputtering rate of 3.5 nm/min) on flat region without islands, the indium signal disappeared, while the Al signal emerged, indicating that a thin Ga-rich InGaN layer was formed among these In-rich islands.

![Figure 3 AES of InGaN islands grown on AlGaN buffer layer. Surface composition on dot (a), on flat region among islands (b) and on flat region after 3 min. \( \text{Ar}^+ \) sputtering (c).](image)

We would like to point out that the deduced In content from Fig. 2(a) does not follow the prediction of equilibrium solubility of GaN in InN based on thermodynamic considerations [13], where the Ga content is expected to increase with the growth temperature. For example, in Ref. [6], the solubility of GaN in InN is predicted to be 3\% at 550 \(^\circ\text{C}\) and 6\% at 725 \(^\circ\text{C}\), while our result shows 13\% at 550 \(^\circ\text{C}\) and about 1\% at 725 \(^\circ\text{C}\). This implies that the incorporation of Ga into InN during the growth of InGaN islands is affected by surface kinetics of In and Ga adatoms rather than equilibrium solubility.

PL measurements of these In-rich InGaN islands in both the visible and the NIR ranges are shown in Fig. 4(a) and 4(b), respectively. No PL signal was observed for the 550 \(^\circ\text{C}\) grown sample. Samples grown at \( T_g = 600-725^\circ \text{C} \) exhibit a NIR emission band in the range of 0.77-0.92 eV. The redshift in PL peak energy with the increasing \( T_g \) agrees fairly well with the corresponding In content determined by XRD. Therefore, we identify the NIR band as the emission from In-rich InGaN islands. For the sample grown at 750 \(^\circ\text{C}\), the NIR band disappeared due to the absence of In-rich islands, which is also consistent with its surface morphology and XRD data. Apart from the NIR band, we also observed a visible emission band in the range of 2.0-2.4 eV for samples grown at \( T_g \geq 650^\circ \text{C} \). Significantly, the visible emission is even stronger for the sample grown at \( T_g = 750^\circ \text{C} \), in spite of the absence of In-rich islands, implying that the visible band is not originated from the InGaN islands.

![Figure 4 Photoluminescence spectra of InGaN grown at different temperatures from 550 to 750 \(^\circ\text{C}\) in (a) the NIR and (b) the visible spectral ranges.](image)

To further clarify this point, we have performed NSOM measurements at room temperature to obtain both the surface morphology and the spatial distribution of visible emissions. Figure 5(a) shows such a NSOM image of the 725 \(^\circ\text{C}\) sample mapping at its visible emission peak. The corresponding surface morphology is shown in Fig. 5(b). It can be clearly seen that these two images are nearly complementary, with dark spots in the NSOM image corresponding very well with the In-rich islands revealed in the AFM image. This confirms that the visible emission is indeed originated from the “flat region” outside these In-rich islands. As can be seen from Fig. 5(a), the visible emission is mainly contributed from some bright spots located among these In-rich islands. As mentioned above, the deposited Ga atoms during the growth of InGaN islands may form a thin Ga-rich layer among these In-rich islands, due to the considerable shorter migration length for Ga adatoms. The visible emission band may be originated from...
defect centers induced during the formation of the Ga-rich layer.

Figure 5 (a) NSOM image of the 725 °C grown sample mapping at its visible emission peak at room temperature. (b) The corresponding AFM image.

4 Conclusion In summary, the surface morphologies, alloy compositions and PL properties of In-rich InGaN islands grown by MOCVD at $T_g = 550$-750 °C have been investigated. The nucleation of InGaN islands was found to be governed by the surface migration of In adatoms. In particular, we found that the incorporation of Ga into InN during the growth of InGaN islands is governed by adatom migration capability, which tends to decompose into In-rich islands and a thin Ga-rich layer at higher growth temperatures. In-rich islands exhibit PL emission in the NIR range, while the formation of a thin Ga-rich layer is likely to be responsible for the observed visible emission band.

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