Prevention of In droplets formation by HCl addition during metal organic vapor phase epitaxy of InN

Sang Won Kang, Hyun Jong Park, Yong Sun Won, Olga Kryliouk, Tim Anderson, Dmitry Khokhlov, and Timur Burbaev

Department of Chemical Engineering, University of Florida, Gainesville, Florida 32611

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The low decomposition temperature of InN and relatively high thermal stability of NH3 necessitate the use of a high NH3/TMIn ratio to prevent In droplet formation on the surface. This work shows that the addition of Cl in the form of HCl (Cl/In molar ratio range of 0.3–1.4) to the growth chamber allows the growth of high quality InN films without the formation of a second In phase at a very low value of the N/In ratio (2500). Photoluminescence spectra in the temperature range of 144 to 4.5 K showed a broad spectral band with a cutoff energy close to the reported minimum of the InN band gap energy (0.65 eV). © 2007 American Institute of Physics.

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Until recently, interest in indium nitride (InN) was mainly derived from its role as a dilute component for enhancing emission efficiency of GaN and AlN active layers in optoelectronic devices. Although pure InN is much less studied than GaN or AlN, significant differences in the measured optical properties have generated intense interest. Specifically, the band gap energy of wurtzite InN crystals, although still a subject of research, has been reported in the range of 0.65 to 0.9 eV. These values are considerably lower than the previously accepted value of 1.9 eV, suggesting applications for InN such as solar cells and infrared laser diodes.

The growth of InN films must overcome several challenges. The high N2 equilibrium partial pressure over InN (Refs. 11 and 12) requires either a low growth temperature or high and reactive nitrogen over pressure. For deposition by metal organic vapor phase epitaxy (MOVPE), this translates into a growth temperature typically <650 °C and high N/In ratio (e.g., NH3 to TMIn ratio >104 at 650 °C) (Ref. 9) to avoid In liquid deposition during growth. The high N/In ratio is required since the homogeneous decomposition rate of NH3 is low at 650 °C. At this condition it is estimated that less than 0.01 mol % of the In atom reacts with the NH3 source, rendering the process extremely inefficient in NH3 usage. The formation of liquid In droplets during growth is obviously undesirable because of its impact on the surface morphology and because it fixes the chemical potential and thus the intrinsic equilibrium point defect structure of the grown InN. The growth of InN for device applications will require control of point defect concentrations (e.g., p-type doping, optical properties) and thus the ability to control the film N/In ratio, and specifically to increase it above that typically produced by MOVPE. Increasing the NH3 partial pressure not only increases the reactive nitrogen but also decreases the In through reaction with hydrogen produced by ammonia decomposition.13 This then lowers the growth rate. In addition to growing at high ammonia partial pressure, the generation of reactive nitrogen radicals by both plasma-assisted15 and ArF excimer laser assisted MOVPE (Ref. 17) has been demonstrated to avoid In droplet formation.

In this letter a simple approach to adjust the stoichiometry of InN is studied. The intentional addition of HCl to the growth chamber should form volatile indium chlorides [InCl (Tsub.=450 °C) and InCl3 (Tsublimation=500 °C)] to lower the In chemical potential. A series of films was grown at various HCl inlet partial pressures to test the feasibility of this approach in controlling In droplet formation and subsequently InN stoichiometry.

A simple equilibrium analysis was first performed to better understand the effect of HCl addition on the deposition chemistry. Kumagai et al. reported equilibrium calculations of hydride VPE of InN using either InCl or InCl3 as the precursor. It is not surprising that they suggested that the higher Cl/In precursor, InCl3, is preferred for deposition of InN since the thermodynamic driving force for the formation of InN is higher.18 Hydrogen, as derived from either decomposition of NH3 or the carrier gas, decreases the driving force. Their results agree with the experimental observation that the growth of InN is higher with the use of InCl3 versus InCl and when substituting inert N2 for H2 carrier. This analysis, however, only studied two values of the Cl/In inlet ratio (i.e., 1 and 3, depending on the In species used).

At typical InN deposition temperatures, the dominant Cl, species is InCl. The effect of adding HCl to the reactor can be seen by comparing the Gibbs energy of the etching reactions for liquid In and solid InN to produce InCl and H2 in addition to N2 in the InN case. The standard ΔG° values for these two reactions were calculated using the ThermoCalc database SUB94 (Ref. 20) and plotted as a function of temperature in Fig. 1. It is clear that HCl attack of pure indium metal is preferred over reaction with InN and the large negative ΔG° at typical growth temperature should yield a significant reaction extent. It is also noted that ΔG° for etching InN is zero at 900 K, suggesting that at higher growth temperature, the addition of HCl will produce significant increase in InN growth rate and possibly no growth.

The hypothesis that the addition of Cl to the deposition system can prevent deposition of In droplets during MOVPE of InN was tested in a reactor that has both MOVPE and hydride VPE capability. By adjusting the HCl/TMIn inlet molar ratio to values of 1 and 3, hydride VPE using either
InCl or InCl$_3$ was operational. A more complete description of this reactor is given elsewhere.\textsuperscript{23} InN was grown using TMIn (TMIn solution, Epichem), NH$_3$ (anhydrous grade 5, Matheson-Trigas), and HCl (10% HCl, 90% N$_2$, Air Products) reactants on c-Al$_2$O$_3$ and GaN/c-Al$_2$O$_3$ (5 µm thick GaN grown by MOVPE) substrates. For all films grown in this study, the operating pressure was 760 Torr and the growth temperature was 550 °C, while the NH$_3$/TMIn ratio was 2500. The total N$_2$ carrier gas flow rate was 4 slm (standard liters per minute) while the NH$_3$ and TMIn flows were 1750 and 0.7 SCCM (SCCM denotes cubic centimeter per minute at STP), respectively. The HCl/TMIn inlet molar ratio was varied in the range of 0–5 by changing the 10% HCl in N$_2$ gas flow rates from 0 to 35 SCCM.

Preliminary experiments indicated that MOVPE growth (i.e., no HCl feed) under these conditions gave two-phase In–InN growth. Two-phase deposition at this relatively low NH$_3$/TMIn ratio is consistent with that previously reported.\textsuperscript{22} The x-ray diffraction pattern of a film grown on GaN/c-Al$_2$O$_3$ without HCl is shown in Fig. 2 bottom). The pattern clearly shows the presence of InN (002) and a strong In (101) peak. In contrast, the pattern for a film grown with a flow of 0.5 SCCM of HCl (HCl/TMIn ratio=1.4) while maintaining the same NH$_3$/TMIn ratio of 2500 shows no evidence for the formation of a pure In phase, as shown in Fig. 2 top spectrum). Some reduction in InN film growth rate ~1 to ~0.8 µm/h, however, was observed.

The surface morphology of the InN films grown at various values of HCl/TMIn molar ratio was examined by scanning electron microscopy (SEM) and atomic force microscopy (AFM), while the composition was determined by Auger electron spectroscopy (AES) and energydispersion spectroscopy (EDS). Only the GaN/Al$_2$O$_3$ substrates were examined in this study. Islands of pure solid In, presumably formed by solidification upon cooling of In droplets nucleated during growth, are clearly observed in the SEM image shown in Fig. 3(a) for the sample grown at the relatively low NH$_3$/TMIn ratio (2500) of this study. An EDS analysis of one of the islands detected only In [see insert in Fig. 3(a)]. As shown in Fig. 3(a), the In surface feature size was variable (3–5 µm) for growth without HCl. This size variation suggests that during growth In droplet nucleation occurred at different times or they partially coalesced.

The SEM micrograph shown in Fig. 3(a) is contrasted to that shown in 3(b) for which HCl was added during film growth (HCl/TMIn molar ratio=1). In this latter film, the InN is polycrystalline and In droplet formation was not evident. The rms roughness of the surface of the InN grown at HCl/TMIn molar ratio=1 is 91.9 nm, as determined by AFM [Fig. 3(d)]. It is noted that In droplets did not nucleate on any films grown when HCl was added in the range of 0.3 ≈ HCl/TMIn < 5. The observed surface morphologies of films grown with a HCl/TMIn ratio in the range of 0.3–3.0 were similar to that shown in Fig. 3(b). As shown in the micrograph in Fig. 3(c), the surface was not completely covered at high HCl/TMIn ratio=4.0, apparently because of the enhanced HCl etching of InN. Interestingly, for higher HCl/TMIn molar ratio >5, a sparse density of microrods was observed. AES did not detect chlorine to be within the detection limit (~0.1 at.%) in the InN film even at the highest HCl/TMIn ratio examined (5.0).

Figure 4(a) shows photoluminescence (PL) spectra collected at three measurement temperatures for an InN film grown on GaN/sapphire. The GaN film thickness was 5 µm.
and the 1 μm InN film was grown at the base conditions (N/In=2500, \( T=550 ^\circ C \)) and \( \text{HCl/TMIn}=1 \). The PL occurs in a wide spectral band from 0.75 to 1.2 eV with a maximal PL intensity at about 1 eV. The spectral position of the PL maximum does not shift with temperature, while the PL intensity grows by a factor of about 1.5 upon lowering the measurement temperature from 144 to 4.5 K. Although not shown in this figure, there is little change in the PL spectra for samples grown at a HCl/TMIn ratio in the range of 0.3–5.0.

One possible explanation for the wide energy band of the PL spectra is the formation of a graded \( \text{In}_{1-x} \text{Ga}_x \text{N} \) interfacial region emanating from the InN/GaN interface. In this case, contribution of layers with different compositions would provide PL at different energies, thus forming a wide PL band.

To determine if the GaN buffer layer affects the PL, films were grown on two different substrates, one with a GaN buffer layer and another with an InN buffer. Figure 4(b) compares room temperature PL spectra for two samples grown on c sapphire at the base conditions and HCl/TMIn ratio =1.4 with either a GaN or a low-temperature InN buffer layer. The PL maximum in the GaN-free sample is strongly shifted to lower energy consistent with GaN incorporation in the InN film playing a significant role in shaping the PL spectra. The process of incorporating Ga into the InN could result from bulk diffusion or through initial etching of the GaN by HCl and reincorporation in the growing InN film. It is important to note that the red cutoff energy of the PL band is close to the lowest reported gap value of InN (i.e., 0.65 eV).

In conclusion, the addition of Cl to the deposition zone was shown to be effective in reducing the In chemical potential. Simply adding a threshold level of HCl during growth led to the disappearance of In droplets that occurred at otherwise identical conditions. As suggested by a simple thermodynamic analysis and supported by experimental results, InN can be grown without the inclusion of a second phase at N/In ratio as low as 2500 by adding HCl in the HCl/TMIn molar ratio range of 0.3–1.4 without chlorine incorporation. PL spectra of InN films show emission over a wide energy range with a lower edge close to the lowest reported value of 0.65 eV. Comparison of room temperature PL spectra for films grown with and without a GaN buffer layer on c sapphire gives results that are consistent with the formation of a \( \text{Ga}_x \text{In}_{1-x} \text{N} \) interfacial layer on the GaN buffer layer.

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