

∂ Open access • Journal Article • DOI:10.1088/0022-3727/43/35/354003

Microstructural origins of localization in InGaN quantum wells — Source link []

Rachel A. Oliver, S.E. Bennett, Tongtong Zhu, D.J. Beesley ...+4 more authors

Institutions: University of Cambridge, University of Oxford

Published on: 08 Sep 2010 - Journal of Physics D (IOP Publishing)

Topics: Quantum well

Related papers:

- Origin of defect-insensitive emission probability in In-containing (AI,In,Ga)N alloy semiconductors.
- Electron-beam-induced strain within InGaN quantum wells: False indium "cluster" detection in the transmission electron microscope
- Three-dimensional atom probe studies of an InxGa1-xN/GaN multiple quantum well structure: Assessment of
 possible indium clustering
- Role of self-formed InGaN quantum dots for exciton localization in the purple laser diode emitting at 420 nm
- Optical and microstructural studies of InGaN/GaN single-quantum-well structures





Microstructural origins of localization in InGaN quantum wells

R A Oliver, S E Bennett, T Zhu, D J Beesley, M J Kappers, D W Saxey, A

Cerezo, C J Humphreys

▶ To cite this version:

R A Oliver, S E Bennett, T Zhu, D J Beesley, M J Kappers, et al.. Microstructural origins of localization in InGaN quantum wells. Journal of Physics D: Applied Physics, IOP Publishing, 2010, 43 (35), pp.354003. 10.1088/0022-3727/43/35/354003. hal-00569690

HAL Id: hal-00569690 https://hal.archives-ouvertes.fr/hal-00569690

Submitted on 25 Feb 2011

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Microstructural origins of localisation in InGaN quantum wells

R.A. Oliver¹, S.E. Bennett¹, T. Zhu¹, D.J. Beesley¹, M.J. Kappers¹, D.W. Saxey², A. Cerezo², C.J. Humphreys¹

1 Department of Materials Science and Metallurgy, University of Cambridge, Pembroke Street, Cambridge, CB2 3QZ, UK

2 Department of Materials, University of Oxford, Parks Road, Oxford, OX1 3PH, UK

Abstract

The startling success of GaN-based light emitting diodes despite the high density of dislocations found in typical heteroepitaxial material has been attributed to localisation of carriers at non-uniformities in the quantum wells which form the active region of such devices. Here, we review the different possible structures within the quantum wells which could act as localisation sites, at length scales ranging from the atomic to the tens of nanometre range. In some quantum wells several localisation mechanisms could be operational, but the challenge remains to optimise the quantum wells' structure to achieve improved quantum efficiencies, particularly at high excitation powers.

Keywords

GaN; InGaN; quantum wells; localisation; transmission electron microscopy; atom probe tomography; spinodal decomposition

PACS codes

68.37.-d, 68.65.Fg, 81.07.St, 78.55.Cr, 81.05.Ea III-V

<u>1. Introduction</u>

Light emitting diodes (LEDs) based on gallium nitride (GaN) are increasingly ubiquitous, in everyday objects such as televisions and mobile phones. However, their most crucial component, the active region of the device from which the light is actually emitted, remains poorly understood, with its very structure a matter for debate. In this paper, we seek to review current understanding of the micro- and nano-scale structure of the quantum wells (QWs) which form the active region of almost all LEDs, and to delineate the potential impact of that structure on device efficiency. In so doing, we will have to address controversies over the best ways to characterise such structures.

The challenges in understanding the behaviour of GaN-based LEDs arise from the fact that most such devices are grown heteroepitaxially on substrates such as sapphire, silicon carbide or silicon. The lattice parameter mismatch between the GaN and the substrate results in the presence of a high density of threading dislocations (more than 10^{10} cm⁻³ in early devices [1]). It should be noted that although these threading dislocations arise as a consequence of the lattice parameter mismatch, they do not necessarily relieve the misfit strain between the epitaxial layer and the sample, and their mechanism of formation is a subject of ongoing debate [2, 3, 4]. Irrespective of their origin, however, the success of GaN-based LEDs is rather surprising in the context of such high defect densities, since in other, more conventional optoelectronic semiconductors, such as GaAs, defect densities a million times lower would be expected to lead to excessive non-radiative recombination, generation of heat and speedy device failure [5].

The suggestion thus arises that in GaN-based LEDs some feature of the micro- or nanostructure is preventing carriers from diffusing to dislocation cores and recombining nonradiatively. Here we will focus on the highly efficient GaN-based LEDs which emit visible light and employ one or more indium gallium nitride (InGaN) QWs in their active region, one purpose of which is to provide a narrow spatial region with reduced bandgap, in which both electrons and holes are trapped, increasing the probability that they will meet and recombine. Hence, it is the small-scale structure of the InGaN QW, and its potential role in localising the carriers, preventing their diffusion to dislocations, which will most concern us. The reader should be aware that this review focuses solely on polar structures grown on the (0001) plane of GaN. Assessment of the impact of the small scale structure non- and semi-polar QWs is an interesting topic, but has not yet received much attention from researchers.

Localisation in InGaN QWs has been postulated to occur on a range of different length scales - ranging from the single or few atom scale, through the few nanometre scale to structures with extents up to tens of nanometres. We will review each of these different length scales. However, it should be noted that postulated mechanisms at the single or few atom level in InGaN should be relevant to all indium-containing nitride QWs, regardless of their broader microstructure. Mechanisms of localisation at this near-atomic level will be assessed briefly in Section 2. Section 3 will then assess postulated microstructures which might lead to localisation at the few nanometre scale such as local variations in composition or QW thickness. Lastly, Section 4 will cover observations of microstructural changes on longer length scales (tens of nanometres to a few hundred nanometres) which could alter carrier localisation or otherwise prevent carrier diffusion to dislocation cores. It should be noted that here we are using the term "microstructure" to indicate all aspects of small-scale structure which may be assessed using a variety of microscopes, not just structures on the micron scale.

2. Localisation at the atomic scale

Much of the discussion about localisation in InGaN QWs has been driven by considerations of experimental data - data concerning either QW microstructure or optical properties. However, the driving force for consideration of the potential role of single atoms or few-atom structures comes not from experiment, but from elegant theoretical studies employing large supercell empirical pseudopotential calculations. Bellaiche *et al.* [6] used this approach to suggest that in InGaN strong localisation of the hole around indium sites may occur even in the case of an entirely random alloy of InN and GaN. Whilst Bellaiche *et al.* [6] modelled zincblende InGaN, rather than the wurtzite structure which is more commonly grown, Wang [7] soon performed similar calculations for wurtzite material, and predicted hole localisation on chains of atoms containing two adjacent indium atoms (i.e. -In-N-In- chains).

Whilst these theoretical contributions are undoubtedly thought-provoking, some comparisons of their predictions with experiment may reduce our confidence in them. For example, Bellaiche *et al.* [6] predict a bandgap bowing parameter in excess of 5 eV for the GaN-InN alloy system, and suggest that this very large value is a consequence of the localisation of holes at Indium sites. However, whilst this calculated value was in reasonably good agreement with experimental values available at the time, more recently a reassessment of the band gap of InN [8], and subsequent new measurements of the bowing parameter, suggest much smaller values, typically around 1.4 eV [9].

Wang's more extensive study [7] aimed to predict how the photoluminescence spectra of InGaN QWs might vary with the indium content of the QW as a consequence of the calculated localisation phenomena. The theory predicted a decrease in the observed linewidth as the indium content increased from zero towards 50%. However, experimental observations yield an opposite trend [10]. For example green-emitting QWs typically contain a higher proportion of indium and exhibit a larger photoluminescence linewidth than similar blue-emitting structures. This discrepancy suggests that whilst Wang's theory may be relevant to the localisation process in InGaN QWs, it does not tell the whole story.

Nonetheless, there is some experimental data which supports the suggestion that localisation of the hole occurs on an atomic or a near-atomic scale. Most notably, Chichibu *et al* [11] showed that positron annihilation spectroscopy provides evidence for localisation of holes on a very small length scale, and suggested that this localisation arose at -In-N-In- chains. Additionally, Graham *et al.* [12] compared resonantly-excited photoluminescence spectra recorded at low temperature from InGaN/GaN multiple quantum wells (MQWs) with the results of a theoretical model which included the deformation potential coupling of carriers to an effectively continuous distribution of longitudinal acoustic phonon modes. Based on this analysis, they found an in-plane localisation length scale of about 2.5 Å, which is compatible with the localisation of holes on -In-N-In- chains [12].

In discussing the concept of localisation at the near-atomic scale, most authors [6, 7, 11] suggest that if the hole is localised at an -In-N-In- chain, the electron will also be localised due to the formation of a bound excitonic state. However, some experimental analysis by Kalliakos *et al.* [13] contradicts this suggestion. Kalliakos *et al.* [13] studied low-temperature photoluminescence spectra from InGaN/GaN QWs in detail, comparing the intensity of the observed phonon replicas and the main emission peak, and calculating the Huang-Rhys factor, *S.* They were unable to fit the resulting data to an excitonic model of the electron and hole in the InGaN, and instead suggested that a model similar to a donor-

acceptor pair (DAP) picture was more suitable [13]. This would suggest that the electron and the hole are localised separately, at spatially distinct locations in the quantum well.

Hence, a picture of localisation in InGaN quantum wells involving holes trapped at randomly formed -In-N-In- chains does not appear to account for all the experimental observations, although there is certainly some evidence that near-atomic scale localisation is a contributory factor. However, the question of the localisation of the electron is apparently left outstanding. Kalliakos *et al.* [13] suggest that the electron and hole are localised at spatially separate locations, but the theoretical studies do not suggest that the electron would be localised at -In-N-In- chains unless it were bound to a localised hole. Hence, the nature of electron localisation sites remains mysterious, and provides us an impetus to turn to microstructural characterisation of InGaN QWs, to look for non-uniformities in the quantum well at above the single atom level which might solve this puzzle.

3. Localisation at the nanometre scale

In considering possible microstructural origins of localisation at the nanometre scale, two main candidates emerge: non-random variations in the composition of the InGaN QW, within the plane of the well, often referred to as indium clustering, and variations in the thickness of the InGaN QW. We will address each of these in turn.

3.1 Indium clustering

The formation of non-random "clusters" of indium in InGaN quantum wells was suggested by Narukawa *et al.* [14], not long after the first nitride-based blue-violet laser diodes (LDs) were introduced [15]. In transmission electron microscopy (TEM) images of InGaN QWs in such an LD structure, small (2 - 5 nm) regions of strain contrast were observed. These features were interpreted as being indium-rich regions which had self-assembled in the quantum well, possibly by spinodal decomposition. Energy dispersive X-ray analysis also suggested an increased indium content in these dot-like regions. Subsequent studies by Gerthsen *et al.* [16] using digital analysis of the lattice fringe spacings in high resolution TEM images suggested that the indium content in these regions (given here as the value of x in $In_xGa_{1-x}N$) could be greater than 80%. Such regions of increased indium content – and hence reduced bandgap – would be expected to localise carriers in the quantum well, and indeed theoretical calculations by Kent and Zunger [17] (again using the large supercell empirical pseudopotential method) indicate that both electrons and holes would be localised at these types of non-random compositional fluctuations.

Hence, the idea that nanoscale indium-rich regions localised both the electron and hole in InGaN quantum wells, preventing carrier diffusion to dislocations, appeared to solve the mystery surrounding GaN-based LEDs. Gradually, however, problems with this hypothesis started to emerge. Firstly, it was not clear how these regions of very high indium content had formed. Narukawa *et al.* [14] initially suggested that a notionally barrierless phase separation mechanism called spinodal decomposition was responsible, basing this suggestion on a theoretical phase diagram calculated by Ho and Stringfellow [18] for the InN-GaN alloy system in the zincblende structure. This diagram does suggest that for typical InGaN growth temperatures of 700 to 800 °C, alloys containing more than about 20% indium would be within the spinode. These indium levels are found in blue- and green-emitting quantum wells, so the spinodal decomposition concept initially appears relevant.

However, the phase diagram Ho and Stringfellow [18] calculated is a *chemical* spinode, which does not take into account the coherency strains associated with the formation of interfaces between indium- and gallium-rich regions. The *coherent* spinode, influenced by

these strains, would be expected to be shifted to lower temperatures [19], making spinodal decomposition at relevant growth temperatures less likely. Additionally, the Ho and Stringfellow phase diagram [18] is for relaxed, bulk-like InGaN, whereas InGaN in QWs is biaxially strained. Calculations by Karpov [20] for the phase diagram of biaxially strained InGaN suggested a spinode shifted to much lower temperatures and higher indium contents than those predicted by Ho and Stringfellow. The Ho and Stringfellow [18] phase diagram and the Karpov [20] phase diagram are displayed in Figure 1 for comparison. (More recent work by Liu and Zunger also supports the suggestion that spinodal decomposition may be suppressed by epitaxial strain [21]). For the Karpov phase diagram, even QW material in green LEDs (which might be grown at 700 °C, and contain about 25% indium) would be outside not only the region within which the barrierless spinodal decomposition process occurs, but outside the broader region for which phase separation may occur by the slower nucleation and growth mechanism. In fact, according to Karpov's calculations [20], InGaN in QWs in blue, green and UV LEDs should be a random alloy. Some experimental support for the suppression of decomposition by epitaxial strain is provided by the work of Rao et al. [22] who studied InGaN epilayers more than 200 nm thick in TEM, and observed phase separation resulting in indium rich regions on a scale of 10 - 20 nm in relaxed regions of their films. However, regions of the films close to the InGaN/GaN interface, which were coherent with the substrate, did not show any evidence of phase separation. (Phase separation in thick, relaxed InGaN epilayers has been described in various publications (for example ref. [23]). However, we will not review this phenomenon extensively here since such thick InGaN layers have little relevance to the active region of most current LEDs).

A significant concern about the TEM methods used to detect apparently indium-rich clusters in InGaN QWs was raised by Smeeton *et al.* [24, 25] and O'Neill *et al.* [26]. Both sets of authors observed that in TEM images of InGaN QWs which had not been exposed to the electron beam prior to the image being recorded, the QWs appeared rather uniform. However, continued exposure to the electron beam led to the appearance of locally strained regions very similar to those observed by Narukawa *et al.* [14]. An example of this effect is shown in Figure 2, which shows a comparison between high resolution TEM images of an In_{0.22}Ga_{0.78}N QW before (Figure 2(a)) and after (Figure 2(b)) exposure to a 400 kV electron beam for several minutes. Following exposure, strong contrast fluctuations are observed in the InGaN. Smeeton *et al.* [27] went on to demonstrate that this phenomenon of electron beam damage causing the formation of apparent indium-rich regions in the QW applied to material grown by both of the most common nitride growth techniques (molecular beam epitaxy (MBE) and metalorganic vapour phase epitaxy (MOVPE)). Additionally, the phenomenon was observed even when the thin-foil TEM specimen was prepared solely by mechanical polishing and etching, without exposure to potentially damaging ion beams [27].

The possibility that indium-rich regions observed in QWs in TEM were an imaging artefact rather than an as-grown feature of the material, motivated attempts to study the microstructure of nitride LEDs using alternative techniques. Atom probe tomography (APT) appeared particularly promising since it provides three-dimensional information about a material's composition at a sub-nanometre scale [28, 29]. For APT a needle-shaped sample with an apex radius of less than 100 nm is held at high voltage, so that the electric field at the apex is almost sufficient to ionize and field-evaporate atoms from the surface. In the analysis of semiconductors, field-evaporation is triggered by pulses of thermal energy provided by a laser. Evaporated ions are projected by the radial electric field on to a position sensitive detector and their chemical identities are determined by time-of-flight mass spectrometry. A three-dimensional atom map is then reconstructed from the data [30].

Having first developed a technique for the preparation of appropriate needle-shaped samples from GaN/sapphire wafers in the focused ion beam microscope (FIB) [31], Galtrey et al. [32] were able to record the first APT data from a nitride semiconductor. The initial data set recorded from a blue-emitting MQW sample, showed a region of about 20 nm × 20 nm in lateral extent, containing four quantum wells, one of which had been damaged in the sample preparation process. Following improvements to the sample preparation technique and employing the local electrode atom probe (LEAP) which has an enhanced field of view, significantly larger regions of both blue- and green-emitting MQW samples were analysed. Figure 3 shows an example reconstruction of an APT data set recorded from a green-emitting InGaN QW sample, in which three QWs can be seen across a region of lateral extent ca. 100 nm [33]. For all the available data sets the distribution of indium in the quantum well was compared to that expected for a random alloy, but no statistically significant deviations from randomness were observed. APT has frequently been used in other materials systems to detect compositional variations at the nanometer scale, and even the rather slight fluctuations which may be present at the onset of spinodal decomposition have been detected [34, 35, 36]. Hence, at least for the MQW samples that have been studied thus far by APT, the existence of non-random nanometer-scale regions with indium content in excess of 80%, such as those suggested by Gerthsen et al. based on TEM data [16], appears extremely unlikely, and there is not even any evidence to support the existence of more subtle non-random indium content fluctuations at the few nanometer scale. This suggests that such indium clusters cannot be necessary for bright luminescence from InGaN QWs. The available data does not however imply that such clusters can *never* be present in QW samples, nor that such clusters, if present, would not be effective in preventing carrier diffusion to dislocation sites.

Since Narukawa et al. [14] initially used TEM to attempt to image indium clustering in InGaN QWs, the application of TEM to these materials has been substantially refined, perhaps most notably by Bartel et al. [37], who have made every effort to optimize the preparation of their thin foil TEM samples and to minimize the extent of electron-beam damage by using short exposure times. They have also very carefully assessed the impact of noise in the TEM in defining detection limits for indium concentration fluctuations. Their resulting high-resolution images appear qualitatively rather different to those seen in earlier studies, with the quantum wells looking rather uniform to the naked eye with an absence of blotchy strain contrast. However, digital analysis of lattice parameter variations within the quantum well can pick up subtleties that the eye would miss. Bartel et al. [37] used this approach to perform a statistical comparison of their data with that expected if the QWs were a random alloy (similar to the approach used to analyse APT data). The analysis here is somewhat hampered by the difficulty of quantifying the exact thickness of the TEM foils used, so that in some cases the data had to be compared to the results expected for a random alloy for various thicknesses of TEM foil and then suggestions made as to which thickness was most realistic. However, for samples where it was possible to measure the foil thickness, Bartel et al. [37] reported deviations from randomness for QWs with indium contents of 25% or more. The blue- and green-emitting samples studied by Galtrey et al. [33] had indium contents of 18% and 25% respectively so that there is some cross-over in the compositions of the quantum wells found to be random by APT and those found to exhibit some deviation from randomness in the TEM studies by Bartel et al. [37]. This discrepancy has been the source of ongoing controversy.

In considering the differences between the two data sets, however, it should be noted that samples of the same average composition grown by different methods can differ either subtly

or sometimes substantially, as Bartel *et al.* [37] note in their discussion of X-ray diffraction data pertaining to phase separation in InGaN. Additionally, whilst Bartel *et al.* [37] do not directly quantify the local indium contents associated with the fluctuations they observe in QWs, it seems unlikely that the subtle variations seen in their data relate to areas a few nanometers in lateral extent containing up to 80% indium. Related studies on thick InGaN epilayers suggest deviations in indium content of around $\pm 10\%$ on length scales a few nanometers [37]. The impact of such variations on electron and hole localization has not yet been thoroughly modeled.

Overall, state-of-the art experimental data suggest that InGaN quantum wells of the compositions typically found in blue- and green-emitting LEDs do not contain gross indium clusters with an indium content of around 80%. Instead, the InGaN in QWs is either a random alloy, or in some samples may contain subtle non-random variations in composition. If InGaN is a random alloy, localization of holes may still occur at randomly formed –In-N-In- chains. The impact of slight non-random indium content fluctuations on localization has not yet been quantified.

3.2 Monolayer well-width fluctuations

The concept of indium clustering has been quite extensively studied. The possibility of slight (1 or 2 monolayer) fluctuations in the QW's width has received far less attention. Such monolayer well-width fluctuations can localise carriers in GaAs-based QWs, with quantum dot-like optical behaviour being observed in such samples at low temperatures [38]. This is due to the reduction of the impact of quantum confinement on the electron and hole energy levels as the QW width increases. This change in potential is fairly slight and only a little thermal energy is needed for the carriers to escape from the local minima. However, in

InGaN QWs, an additional effect alters the local potential landscape: the high piezoelectric constants of the wurtzite nitrides lead to significant band-bending in strained quantum wells, and as the well thickness increases, so does the impact of this band-bending. For example, for a 3.3 nm thick QW containing 25% indium, Graham *et al.* [39] calculated that an increase in thickness of just 1 monolayer should decrease the transition energy by 58 meV. This suggests that monolayer well-width fluctuations in InGaN could localise carriers at room temperature.

Both TEM and APT have been used to observe such monolayer width fluctuations in InGaN QWs. Scanning transmission electron microscopy high angle annular dark field (STEM-HAADF) images show that the lower interface of an InGaN/GaN quantum well (i.e. the interface where InGaN was grown on GaN) appears abrupt, whereas the upper interface (the interface where GaN was grown on InGaN) appears rough [39]. APT similarly reveals a difference in roughness between the upper and lower QW interfaces [33], with the upper interface being characterised by one or two monolayer high islands a few nanometres in lateral extent atop the InGaN QW surface. Figure 4 shows three-dimensional isoconcentration surfaces derived from an experimental APT data set from a green-emitting InGaN QW sample, illustrating the structure of both the rough upper (Figure 4(a)) and smoother lower (Figure 4(b)) interfaces of well [33]. Detailed atomic force microscopy (AFM) studies of thin InGaN QWs grow via the nucleation, growth and coalescence of two-dimensional islands (the Frank- van-der-Merwe growth mode), providing a mechanism for the formation of the observed well-width fluctuations [33].

The impact of such monolayer well-width fluctuations on carrier localisation has not been studied in detail theoretically, and may not be easily addressed using currently available large supercell empirical pseudopotential models, most of which are for zincblende-structure InGaN and which do not include quantum well interfaces or the effects of strain. However, this area appears ripe for further theoretical exploration to guide potential experimental optimisation of QW interface structure.

4. The impact of broader scale microstructure

When considering the potential impact of broader-scale microstructure (that is microstructure on the tens of nanometre scale, rather than the few nanometre scale) on luminescence from QWs, it is perhaps less appropriate to use the term "localisation" than when describing smaller-scale structure. Whilst many of the broader scale structures which have been observed in InGaN QWs may act to keep carriers away from dislocation sites, they do not do so by trapping the carrier in a small, nanometre-scale area. Instead, they provide a region of increased bandgap around the dislocation, providing a barrier to carrier diffusion to the dislocation, but not necessarily preventing the carriers from moving around freely in other regions of the QW. Since low temperature optical measurements [12], and positron annihilation spectroscopy [11], have suggested very short localisation length scales in InGaN, such "dislocation screening" is unlikely to be the only mechanism influencing carrier diffusion in InGaN QWs, However, reports of very high internal quantum efficiencies in quantum wells in which microstructural characterisation suggests that dislocation screening is occurring [40, 41] suggest that it could be an important contributory factor.

The microstructures which have been suggested to lead to dislocation screening are diverse. Hangleiter *et al.* [40] suggested that the formation of V-pits around dislocations during MQW growth leads to the formation of thinner regions of QW on the V-pit walls, hence providing a region of higher potential around the dislocation. Grandjean et al. [42], on the other hand, suggested that step-pinning by dislocations leads to locally thinner QWs around dislocations even in the absence of notable V-pits, again providing a barrier to carrier diffusion to dislocation cores. Van der Laak et al. [43] noted the presence of gross fluctuations in QW width in commercial LEDs on a lateral length scale of tens of nanometres. (By gross here, we imply fluctuations in the QW width of several monolayers, rather than the one to two monolayer fluctuations discussed in the previous section. In fact, the QW width was observed to drop to nothing, leaving a gap in the QW, filled with GaN, at several locations). They showed that such "gappy" QWs consist of a network of interlinking InGaN strips separated by troughs filled with GaN. The majority of threading dislocations were observed to pass through the gaps in the QW or through the very edges of the InGaN strips where the QW thickness and indium content were both found to be lower [43]. Hence, the QW morphology again provides a local potential barrier around the dislocation core. We will discuss this last case in a little more detail to illustrate the ways in which multiple microstructural characterisation techniques can be brought together to provide an enhanced understanding of this type of quantum structure.

Van der Laak *et al.* [43] suggested that the network structures they observed in InGaN QWs arose as a consequence of a two-temperature MOVPE growth method, in which the InGaN QW is grown at a lower temperature than the GaN barrier, and is left exposed to a flow of NH₃ and N₂ during the temperature ramp which is thus required. Such two-temperature growth has been observed to provide high internal quantum efficiencies in both green- and near-ultra-violet-emitting QWs [41, 44]. Similar gross well-width fluctuations were also observed in QWs which had been annealed at their growth temperature for durations

comparable with those required for the temperature ramp [43]. AFM on thin InGaN epilayers intended to mimic these QW growth conditions revealed the existence of the network of interlinking InGaN strips, which were later also observed in QW samples using STEM-HAADF [45]. The location of the dislocations relative to the InGaN strips was originally assessed using dark-field TEM on epilayer samples, which showed that $90 \pm 10\%$ of the TDs intersected the QW plane between or close to the edges of the InGaN strips [43]. We have recently confirmed this observation using cathodoluminescence (CL) imaging in the scanning electron microscope (SEM). Figure 5(a) shows an AFM image of an annealed InGaN epilayer, showing the network morphology. A rather similar pattern is seen in a panchromatic cathodoluminescence image of a QW grown under similar conditions (Figure 5(b)). The InGaN strips appear bright in a CL image because of the greater probability of radiative recombination for carriers trapped in the QW. The location of dislocations in the QW can be identified from the position of surface pits seen in an SEM image which was recorded simultaneously with the CL data (Figure 5(c)). However, the density of surface pits measured by SEM is less than that measured by AFM, suggesting that some pits may have been missed in this analysis. Nonetheless, it is useful to note that comparison of multiple SEM and CL images suggests that $92 \pm 7\%$ of the dislocations run between the InGaN strips or at their very edges, consistent with the result of the TEM-based analysis. The positions of the visible surface pits in Figure 5(c) are shown as white rings on a copy of the panchromatic CL image in Figure 5(d) to facilitate comparison of the two data sets.

Even where a dislocation runs through the edge of a strip it will still be effectively screened from the carriers since the QW is narrower at the strip edge, and also has a lower indium content. The variations in indium content across the strip were initially assessed for epilayer samples using energy dispersive X-ray spectroscopy in STEM [45]. More recently, however, we have used APT to assess the variations in composition within a green MQW sample grown using the two-temperature method. The photoluminescence from this MQW has previously been analysed in some detail, and for its wavelength of emission (552 nm), it exhibits a high internal quantum efficiency of 12% [44]. The APT data is illustrated in Figure 6(a-d). Although the APT field-of-view is fairly small compared to the width of the InGaN strips typically observed in these samples, gaps in several of the InGaN QWs can nonetheless be observed, consistent with previous investigations by TEM [44]. Figure 6(e) shows the variation in indium content from the body of an InGaN strip, towards its edge, which confirms the earlier observation that the edges of the strips are less indium rich than their centres. Comparable analyses across several QWs in different APT images reveal similar trends.

Hence, the combination of TEM, AFM, APT, SEM and CL gives us a rather complete picture of QWs with gross variations in width. With the type of understanding garnered from such microstructural characterisation we can hope to optimise such morphologies in the future to achieve improved device performance.

5. Concluding remarks

We have noted that the structure of InGaN QWs at three different length scales may help prevent the diffusion of carriers to dislocation cores: at the near-atomic level, there is evidence that randomly formed –In-N-In- chains may act to localise holes. At the few nanometre scale, the concept of indium clustering remains controversial, and for blue- and green-emitting quantum wells with indium contents up to about 25% there is little or no reliable evidence for the regions of very high indium content (70 or 80% or even more) that were once thought to be key to the operation of GaN-based LEDs. However, TEM, AFM

and APT all provide evidence for monolayer well-width fluctuations a few nanometres in lateral extent. Lastly, various broader scale morphologies (at a length scale of tens of nanometres) may provide an increased band gap in the regions around threading dislocations, erecting an additional barrier to carrier diffusion to the dislocation core. With various plausible explanations thus on offer for the surprisingly high quantum efficiencies achievable in nitride LEDs despite the high density of defects, it is not necessary that there exists one specific structure common to all LEDs which is the sole key to good performance. (Although if holes are localised at –In-N-In- chains, this will be relevant to all QWs containing indium). Equally, more than one of the mechanisms discussed here may operate within a single LED. (For example, one might envisage holes being localised at –In-N-In- chains, and electrons at monolayer well width fluctuations. At room temperature, where carriers may be able to hop between these localised sites, some broader morphology providing enhanced dislocation screening might also come into play).

The enhanced understanding of QW structure provided by the application of multiple microscopy techniques may, when coupled with appropriate modelling, provide us with sufficient understanding of these various mechanisms that we can design the QW microstructure to achieve optimal efficiencies. This approach may be crucial to the future success of LEDs particularly in the demanding arena of solid state lighting, where industry insiders identify increasing internal quantum efficiency as a key outstanding challenge [46], and where the requirement for low cost solutions probably precludes the use of expensive bulk GaN substrates with very low defect densities. Additionally, another important unanswered question concerns the impact of the various QW microstructures on LED "droop". (This phenomenon involves a significant reduction in LED efficiency as the drive current is increased). The effectiveness of well width fluctuations – for example - in

localising carriers may be rather reliant on the presence of internal electric fields. At higher drive currents, these fields are screened by the large density of carriers present in the material, and hence localisation may become less effective, contributing to the efficiency decrease. Our developing understanding of QW microstructure must address these issues if it is to have significant impact on the development of LEDs for real-world applications.

Acknowledgements

This work was supported in part by the EPSRC. RAO would like to acknowledge funding from the Royal Society. SEB would like to acknowledge funding from Sharp Laboratories of Europe and Pembroke College, Cambridge. The authors would like to thank Prof. George Smith of Oxford University for useful discussions.

Figure Legends

Figure 1: Theoretical phase diagrams for the GaN-InN alloy system: (a) Phase diagram calculated by Ho and Stringfellow for bulk InGaN. Note that the temperature scale is in °C. (Reprinted with permission from [18]. Copyright 1996, American Institute of Physics.) (b) Phase diagram calculated by Karpov including the effects of biaxial strains present in epitaxial growth. Note that the temperature scale is in K. (Reprinted with permission from [20]).

Figure 2: High resolution TEM images of an In_{0.22}Ga_{0.78}N QW: (a) Image recorded within 20 s of first exposing this part of the QW to an electron beam. The InGaN appears fairly uniform. (b) Another image of the same area after a few minutes of exposure to a 400 kV electron beam. Strong fluctuations are now observed in the InGaN. (Reprinted with permission from [24]. Copyright [2003], American Institute of Physics.)

Figure 3: A reconstruction of APT data showing three InGaN QW layers in a greenemitting sample. Indium atoms are shown as black, gallium atoms as gray. The other atoms are omitted for clarity. (Reprinted with permission from [33]. Copyright [2008], American Institute of Physics.)

Figure 4: The (a) upper and (b) lower interfaces of a typical InGaN QW displayed as three-dimensional isoconcentration surfaces derived from the experimental APT data from a green-emitting sample. (Reprinted with permission from [33]. Copyright [2008], American Institute of Physics.)

Figure 5: (a) An annealed InGaN epilayer showing a network of interlinked InGaN strips, with troughs between them. (b) A panchromatic CL image of an InGaN QW, grown and annealed under the same conditions as the epilayer, showing bright emission from the InGaN strips. (c) A SEM image recorded simultaneously with the CL image in B. Pits in the surface indicate the location of dislocations. (d). The positions of the dislocations observed in (c) are here superimposed on the CL image (b) as white rings. Figure 6: APT investigations of a high efficiency green-emitting quantum well: (a) Overview of the atom probe data set viewed from the side, using a x = 6% isoconcentration surface to highlight the quantum wells. Gaps in the quantum wells may be observed. (b) - (d) Three individual quantum well isoconcentration surfaces viewed from above. Gaps are evident in (b) and (c). (e) The variation in indium content moving towards the edge of an InGaN strip. As was previously seen in TEM, the InGaN strips have a lower indium composition towards their edges (i.e. in the area adjacent to the gap). **Figures**

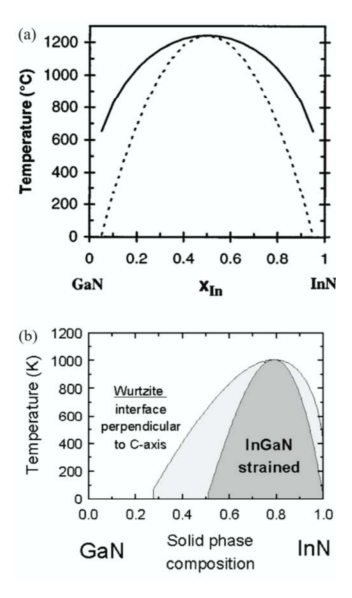


Figure 1

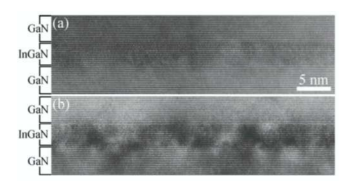


Figure 2

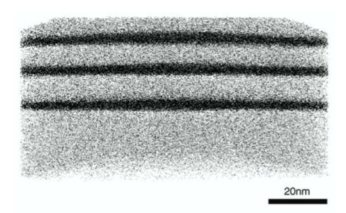


Figure 3

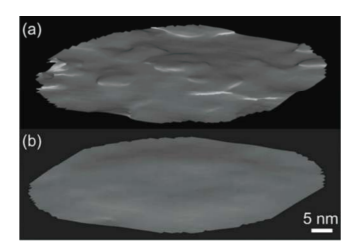


Figure 4

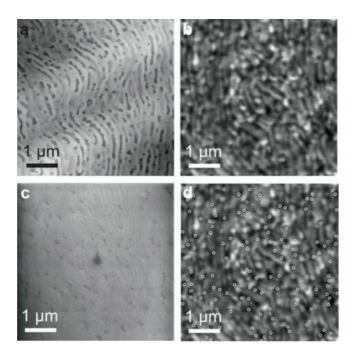
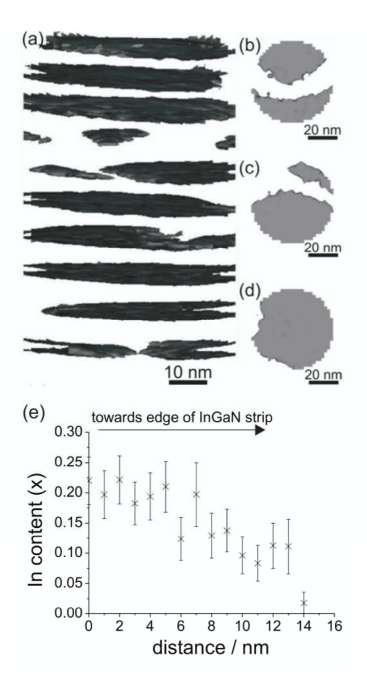


Figure 5





References

- ¹ Nakamura S 1998 Science 281
- ² Ning X J, Chien F R, Pirouz P, Yang J W and Asif Khan M 1996 J. Mater. Res. 11 580
- ³ Oliver R A, Kappers M J and Humphreys C J Appl. Phys. Lett. 89 011914
- ⁴ Moram M A, Oliver R A, Kappers M J and Humphreys C J 2009 Adv. Mat. 21 3941
- ⁵ Herzog A H, Keune D L and Craford M G 1972 J. Appl. Phys. 43 600
- ⁶ Bellaiche L, Mattila T; Wang L W, Wei S H and Zunger A 1999 Appl. Phys. Lett. 74 1842
- ⁷ Wang L W 2001 *Phys. Rev. B* **63** 245107

⁸ Wu J, Walukiewicz W, Yu K M, Ager J W, Haller E E, Lu H, Schaff W J, Saito Y and Nanishi Y 2002 Appl. Phys. Lett. 80 3967

⁹ Wu J 2009 J. Appl. Phys. **106** 011101

¹⁰ Shan W, Walukiewicz W, Haller E E, Little B D, Song J J, McCluskey M D, Johnson N M, Feng Z C, Schurman M and Stall R A 1998 J. Appl. Phys. 84, 4452.

¹¹ Chichibu S F, Uedono A, Onuma T, Haskell B A, Chakraborty A, Koyama T, Fini P T, Keller S, Denbaars S P, Speck J S, Mishra U K, Nakamura S, Yamaguchi S, Kamiyama S, Amano H, Akasaki I, Han J and Sota T 2006 Nature Materials 5 810

¹² Graham D M, Dawson P, Godfrey M J, Kappers M J and Humphreys C J 2006 Appl. Phys. Lett. 89 211901 ¹³ Kalliakos S, Zhang B, Taliercio T, Lefebvre P, Gil B, Grandjean N, Damilano B and Massies J 2002 Appl.

Phys. Lett. 80 428

¹⁴ Narukawa Y, Kawakami Y, Funato M, Fujita S, Fujita S and Nakamura S 1997 Appl. Phys. Lett. 70 981 ¹⁵ Nakamura S, Senoh M, Nagahama S, Iwasa N, Yamada T, Matsushita T, Kiyoku H and Sugimoto Y 1996

Jpn. J. Appl. Phys. **35** L74 ¹⁶ Gerthsen D, Hahn E, Neubauer B, Rosenauer A, Schön, Heuken M and Rizzi A 2000 phys. stat. sol. (a) **177** 145

¹⁷ Kent P R C and Zunger A 2001 Appl. Phys. Lett. 79 1977

¹⁸ Ho I-h and Stringfellow G B 1996 Appl. Phys. Lett. 69 2701

¹⁹ Porter D A and Easterling K E 1981 Phase Transformations in Metals and Alloys 2nd Edition (London: Chapman and Hall) p313

²⁰ Karpov S Y 1998 MRS Internet Journal of Nitride Semiconductor Research **3** 16

²¹ Liu J Z and Zunger A 2008 Phys. Rev. B 77 205201

²² Rao M, Kim D and Mahajan S 2004 Appl. Phys. Lett. 85 1961

²³ Westmeyer A N and Mahajan S 2001 Appl. Phys. Lett. 79 2710

- ²⁴ Smeeton T M, Kappers M J, Barnard J S, Vickers M E and Humphreys C J 2003*phys. stat. sol.* (c) 7 297
- ²⁵ Smeeton T M, Kappers M J, Barnard J S, Vickers M E and Humphreys C J 2003 Appl. Phys. Lett. 83 5419
- ²⁶ O'Neill J P, Ross I M, Cullis A G, Wang T and Parbrook P J 2003 Appl. Phys. Lett. 83 1965
- ²⁷ Smeeton T M, Humphreys C J, Barnard J S and Kappers M J 2006 J. Mater. Sci. 41 2729

²⁸ Cerezo A, Godfrey T J and Smith G D W 1988 Rev. Sci. Inst. 59 862

²⁹ Miller M K, Cerezo A, Hetherington M G and Smith G D W 1996 Atom probe field ion microscopy (Oxford: Oxford University Press)

³⁰ Kelly T F and Miller M K 2007 Rev. Sci. Instrum. 78 031101

³¹ Galtrey M J, Oliver R A, Kappers M J, Humphreys C J, Stokes D J, Clifton P H and Cerezo A 2007 Springer Proceedings in Physics 120 161

³² Galtrey M J, Oliver R A, Kappers M J, Humphreys C J, Stokes D J, Clifton P H and Cerezo A 2007 Appl. Phys. Lett. 90 061903

³³ Galtrey M J, Oliver R A, Kappers M J, Humphreys C J, Clifton P, Larson D, Saxey D, Cerezo A 2008 J.

Appl. Phys. **104** 013524 ³⁴ Miller M K, Hyde J M, Hetherington M G, Cerezo A, Smith G D W and Elliott C M 1995 *Acta Metall. Mater.*

43 3385 ³⁵ Hyde J M, Miller M K, Hetherington M G, Cerezo A, Smith G D W and Elliott C M 1995 *Acta Metall.Mater*. **43** 3403

³⁶ Hyde J M, Miller M K, Hetherington M G, Cerezo A, Smith G D W and Elliott C M 1995 Acta Metall. Mater. **43** 3415

³⁷ Bartel T P, Specht P, Ho J C and Kisielowski C 2007 Phil. Mag. 87 1983

³⁸ Phillips R T, Steffan A G, Newton S R, Reinecke T L, Kotlyar R 2003 phys. stat. sol. b 238 601

³⁹ Graham D M, Soltani-Vala A, Dawson P, Godfrey M J, Smeeton T M, Barnard J S, Kappers M J, Humphreys C J and Thrush E J 2005 J. Appl. Phys. 97 103508

⁴⁰ Hangleiter A, Hitzel F, Netzel C, Fuhrmann D, Rossow U, Ade G, and Hinze P 2005 Phys. Rev. Lett. 95 127402

⁴³ van der Laak N K, Oliver R A, Kappers M J and Humphreys C J 2007 *Appl. Phys. Lett.* **90** 121911.
⁴⁴ Costa P M F J, Datta R, Kappers M J, Vickers M E, Humphreys C J, Graham D M, Dawson P, Godfrey M J, Thrush E J and Mullins J T 2006 *phys. stat. sol. (a)* **203** 1729
⁴⁵ van der Laak N K, Oliver R A, Kappers M J and Humphreys C J 2007 *J. Appl. Phys.* **102** 013513.
⁴⁶ Krames M R, Shchekin O B, Mueller-Mach R, Mueller G O, Zhou L, Harbers G and Craford M G 2007
⁴⁷ Lower Laboratory **1** 160

Journal of Display Technology **3** 160

 ⁴¹ Galtrey M J, Oliver R A, Kappers M J, McAleese C M, Zhu D, Humphreys C J, Clifton P H, Larson D J and Cerezo A 2008 *Appl. Phys. Lett.* **92** 041904
 ⁴² Grandjean N, Feltin E, Butté R, Carlin J-F, Sonderegger S, Deveaud B and Ganière J-D 2007 *Phil. Mag.* **87**