Absence of quantum-confined Stark effect in GaN quantum disks embedded in (Al,Ga)N nanowires grown by molecular beam epitaxy

C. Sinito,*†‡ P. Corfdir,†‡ C. Pfüller,† G. Gao,†‡ J. Bartolomé Vílchez,†‡ S. Kölling,⊥ A. Rodil Doblado,⊥‡ J. K. Zettler,†‡ T. Flissikowski,† P. Koenraad,⊥ H. T. Grahn,† L. Geelhaar,† S. Fernández-Garrido,†△ and O. Brandt*†

†Paul-Drude-Institut für Festkörperphysik, Leibniz-Institut im Forschungsverbund Berlin e.V., Hausvogteiplatz 5–7, 10117 Berlin, Germany
‡Present address: Attolight AG, EPFL Innovation Park, Bât. D, 1015 Lausanne, Switzerland
¶Present address: ABB Corporate Research, 5405 Baden-Dättwil, Switzerland
§Present address: Materials Science and Nanoengineering Department, Rice University, Houston, Texas 77005, United States of America
∥Present address: Departamento de Física de Materiales, Universidad Complutense de Madrid, Madrid 28040, Spain
⊥Department of Applied Physics, TU Eindhoven, 5612 AZ Eindhoven, The Netherlands
#Present address: Biomedical Neuroengineering Research Group (nBio), Bioengineering Institute, Universidad Miguel Hernández, Elche 03202, Spain
@Present address: LayTec AG, Seesener Str. 10–13, 10709 Berlin, Germany
△Present address: Grupo de electrónica y semiconductores, Dpto. Fisica Aplicada, Universidad Autónoma de Madrid, C/ Francisco Tomás y Valiente 7, 28049 Madrid, Spain

E-mail: chiara.sinito@attolight.com; brandt@pdi-berlin.de

Abstract

Several of the key issues of planar (Al,Ga)N-based deep-ultraviolet light emitting diodes could potentially be overcome by utilizing nanowire heterostructures, exhibiting high structural perfection and improved light extraction. Here, we study the spontaneous emission of GaN/(Al,Ga)N nanowire ensembles grown on Si(111) by plasma-assisted molecular beam epitaxy. The nanowires contain single GaN quantum disks embedded in long (Al,Ga)N nanowire segments essential for efficient light extraction. These quantum disks are found to exhibit intense emission at unexpectedly high energies, namely, significantly above the GaN bandgap, and almost independent of the disk thickness. An in-depth investigation of the actual structure and composition of the nanowires reveals a spontaneously formed Al gradient both along and across the nanowire, resulting in a complex core/shell structure with an Al deficient core and an Al rich shell with continuously varying Al content along the entire length of the (Al,Ga)N segment. This compositional change along the nanowire growth axis induces a polarization doping of the shell that results in a degenerate electron gas in the disk, thus screening the built-in electric fields. The high carrier density not only results in the unexpectedly high transition energies, but also in radiative lifetimes depending only weakly on temperature, leading to a comparatively high internal quantum efficiency of the GaN quantum disks up to room temperature.
(Al,Ga)N is the material of choice for the fabrication of deep ultraviolet (UV) solid-state emitters with a wide range of applications in biological, medical, and communication areas. However, planar (Al,Ga)N-based heterostructures suffer from a number of problems, among them high densities of threading dislocations, low dopant activation and carrier injection, poor light extraction, and strong internal electrostatic fields in quantum wells. As a result, the currently existing (Al,Ga)N-based UV light emitting diodes (LEDs) still exhibit comparatively low external quantum efficiencies, and their performance degrades with decreasing wavelength.

(Al,Ga)N-based heterostructures in the form of three-dimensional nanostructures have the potential to resolve several of these issues challenging conventional planar devices. Consequently, the fabrication of such nanostructures is currently the subject of worldwide research activities, utilizing a wide range of techniques ranging from basic top-down and bottom-up approaches to hybrid schemes combining top-down processes and subsequent regrowth. A particularly active field of research is the synthesis of GaN/(Al,Ga)N nanowires by molecular beam epitaxy (MBE). These nanowires tend to form on a variety of structurally and chemically dissimilar substrates, invariably yielding material of high structural perfection, since dislocations do not propagate along the nanowire growth axis. In addition, the efficient elastic strain relaxation at the nanowire sidewalls facilitates the fabrication of GaN/(Al,Ga)N heterostructures with coherent interfaces. Moreover, the elastic strain relief leads to reduced polarization fields as compared to planar layers, which is expected to result in higher internal quantum efficiencies. Finally, the high geometric aspect ratio of nanowires offers superior light extraction.

In most previous works utilizing MBE, the structures under investigation consisted of axial GaN/(Al,Ga)N nanowires containing an active region formed by GaN or (Al,Ga)N quantum disks clad by comparatively thin (Al,Ga)N or AlN barriers. Ultimately, a nanowire heterostructure serving as efficient deep UV emitter should be free of GaN segments to avoid any reabsorption of the light emitted by the quantum disk. The same time, a GaN quantum disk embedded in a pure (Al,Ga)N nanowire or a long (Al,Ga)N segment will exhibit a significantly larger internal field than the same disk in a GaN nanowire with short (Al,Ga)N segments. Since the radiative recombination rate of charge carriers confined in the disk critically depends on the magnitude of the internal electrostatic field, it is of high relevance for future applications to investigate the spontaneous emission of GaN quantum disks embedded in (Al,Ga)N nanowires.

In this Letter, we examine the carrier recombination of GaN/(Al,Ga)N nanowires synthesized by plasma-assisted MBE (PAMBE) on Si(111) substrates. The nanowires contain a single GaN quantum disk embedded in an 0.8 µm long (Al,Ga)N segment. The disks have a thickness ranging from 1.8 to 6.3 nm and emit intense luminescence at an energy invariably above the bandgap of bulk GaN, suggesting that the quantum-confined Stark effect in the disk is essentially absent. A detailed investigation of the composition of the (Al,Ga)N segment reveals a core/shell microstructure formed spontaneously during growth, with a continuously varying Al content both across the nanowire diameter and along the growth direction. The latter induces a high electron density in the shell via polarization doping, leaving the quantum disk in a degenerately doped state with screened internal electrostatic fields thus resulting in the high transition energies regardless of the disk thickness. A quantitative analysis of the transition energies and the temperature dependence of the radiative lifetime allows us to extract not only the mean value of the electron density in the disks, but also its distribution. Further consequences of the degenerate doping level are short radiative lifetime and a comparatively high internal quantum efficiency up to room temperature.

Figure 1(a) shows the nominal structure of our nanowires, which consist of a 750-nm-
Figure 1: Spectrally and spatially resolved luminescence of the GaN/(Al,Ga)N nanowires under investigation at room temperature. (a) Schematic representation of the nominal structure of the nanowires (not to scale). (b) PL spectra of the four GaN/(Al,Ga)N nanowire ensembles compared to the PL spectrum of a GaN nanowire ensemble. The spectra are normalized and vertically shifted for clarity. The numbers indicate the thickness of the GaN quantum disk for the samples containing one. (c) and (d) Monochromatic CL maps at 3.85 eV and 3.55 eV superimposed to bird’s eye view secondary electron micrographs of the GaN/(Al,Ga)N nanowire ensemble with a 6.3 nm-thick GaN disk.

long GaN base grown on Si(111), followed by an 800-nm-long (Al,Ga)N segment and a GaN quantum disk embedded between the (Al,Ga)N segment and a 35-nm-long (Al,Ga)N cap. The samples under investigation differ in the thickness of the disk, with nominal values of 1.8, 3.3, and 6.3 nm. Both the (Al,Ga)N segment and the (Al,Ga)N cap have a nominal Al content of 0.3. For comparison, we also include a sample nominally identical to those above, but without an embedded GaN quantum disk.

Figure 1(b) depicts room temperature photoluminescence (PL) spectra of the four nanowire ensembles under continuous-wave excitation at 244 nm together with that of a bare GaN nanowire ensemble under continuous-wave excitation at 325 nm. The PL spectrum of the GaN nanowires shows a line at 3.412 eV originating from the recombination of free A excitons in strain-free GaN. The PL spectrum of the bare (Al,Ga)N nanowires exhibits a single band at 3.80 eV. A band at essentially the same spectral position (3.85 eV) is also observed in the PL spectra of the (Al,Ga)N nanowires with an embedded GaN disk, and we thus attribute this high-energy band to emission from the (Al,Ga)N segment. Consequently, the low-energy band in the spectra is associated with the GaN quantum disks. In all cases, the corresponding transition energy is notably higher than the bandgap energy of GaN and depends only weakly on the thickness of the disk itself. The same is true for the integrated PL intensity of this band. In addition, the PL intensity drops by only a factor of two to three between 10 and 300 K (see Supporting Information) and is thus about two orders of magnitude higher than that of the bare GaN nanowires at 300 K. These results suggest that the GaN quantum disks exhibit a very high internal quantum efficiency up to room temperature. As shown in the Supporting Information, the low thermal quenching is in part due to an efficient transfer of photoexcited carriers from the (Al,Ga)N segment to the quantum disk, but the actual internal quantum efficiency re-
mains significantly higher (about a factor of 10) than that of GaN nanowires of comparable length and density.

For the nominal parameters of our nanowires [see Fig. 1(a)], we would expect very different transition energies for both the (Al,Ga)N segment and the GaN quantum disk. Specifically, near-band edge emission of Al$_{0.3}$Ga$_{0.7}$N should occur above 4 eV,\textsuperscript{19–22} while the energy observed (3.85 eV) is rather characteristic for an Al content of 0.15.\textsuperscript{23} Moreover, the large internal electrostatic fields in GaN/Al$_{0.3}$Ga$_{0.7}$N quantum disks should result in a quantum-confined Stark shift of typically 200–350 meV/nm,\textsuperscript{24} leading to a strong dependence of the transition energies on disk thickness and emission energies well below the band gap of GaN for the two thicker quantum disks.\textsuperscript{15,25,26}

To confirm our assignment of the PL bands in Fig. 1(b), we perform spatially resolved cathodoluminescence (CL) measurements at room temperature on the ensemble with the thickest GaN disk. Figures 1(c) and 1(d) show monochromatic CL maps collected at emission energies of 3.85 and 3.55 eV, respectively, superimposed on the same bird’s eye view secondary electron micrograph of the nanowire ensemble. The maps clearly show the complementary nature of the two emission bands, in that the band at 3.55 eV arises from the top of the nanowires, where the GaN disk is located, while the homogeneous emission at 3.85 eV originates from the region corresponding to the (Al,Ga)N segment.

The high transition energies of the GaN quantum disks could be explained if their were actually much thinner than nominally. In this case, the quantum-confined Stark shift would be outweighed by quantum confinement, resulting in energies larger than the GaN band gap. To test this hypothesis, we perform a detailed structural and chemical analysis of the nanowire ensemble with the 6.3-nm-thick GaN disk, whose transition energy depends most sensitively on the internal electrostatic fields. Figure 2(a)–2(c) show scanning transmission electron microscopy (STEM) images acquired in high-angular annular dark-field (HAADF) mode for three representative single nanowires from this ensemble. The GaN quantum disk is clearly visible in all three micrographs and has a thickness of (6 ± 1) nm, confirming that the actual thickness is consistent with the nominal value. This result rules out the possibility that the high transition energies are caused by strong quantum confinement in thin disks.

The HAADF-STEM images also reveal that the quantum disks are not delimited by flat (0001) planes, but by facets with semi-polar orientation, as most clearly seen in Fig. 2(c). The angles between the growth axis and the direction perpendicular to the semi-polar plane lie in the range of 20–30°, which thus correspond to (10\overline{1}ℓ) planes with ℓ ranging from 5 to 3. The internal electrostatic field in semipolar strained GaN/(Al,Ga)N quantum wells with these orientations is reduced, but only to about 70% of the one in a GaN/(Al,Ga)N[0001] quantum well.\textsuperscript{27} Fields of this strength (≈ 2 MV/cm) would still lead to transition energies well below the band gap of GaN,\textsuperscript{16} and the occurrence of semipolar facets therefore does not explain the discrepancy between the expected and experimentally observed transition energies.

What we would expect from the various shapes of the GaN quantum disks, however, is a significant variation in emission energy and thus a strong broadening of ensemble spectra such as those depicted in Fig. 1(b). Compared to the full width at half maximum (FWHM) of 40 meV for the room temperature PL line of the GaN nanowire ensemble, the PL bands of the GaN quantum disks are certainly broader, but with an FWHM of 120 meV not nearly as much as expected from the variation in thickness and orientation. The surprising uniformity of the emission energy of the GaN disks is also evident in the CL map displayed in Fig. 1(d), where the majority of the nanowires are seen to emit at 3.55 eV.

The contrast obtained by HAADF-STEM is primarily determined by the composition of the material. For several nanowires, we have observed a contrast along the nanowire cross-section as most clearly visible in Fig. 2(a),
Figure 2: Structural properties of the nanowire ensemble with the 6.3-nm-thick GaN disk. (a)–(c) HAADF-STEM images of three representative nanowires. The micrographs were acquired in the \langle11\overline{2}0\rangle zone axis. (d) Radially averaged Al content along the nanowire growth axis of two individual nanowires recorded by EDX line scans in an SEM. Inset: secondary electron micrograph of a representative nanowire with the scale bar having a length of 200 nm. Most dispersed nanowires exhibit a length between 1.0 and 1.4 \mu m, indicating that they break off at their coalescence joints (see sideview in Supporting Information) upon ultrasonic harvesting. The arrow indicates the [000\overline{1}] growth direction. (e) Ga and Al profiles across the nanowire diameter in proximity of the GaN disk extracted from APT data on a single nanowire. (f) Al content across a two-dimensional section extracted from the same data.

namely, a brighter core and a darker shell, indicating the presence of a core/shell structure with an Al-deficient inner core and an Al-rich outer shell which wraps around the core.

To investigate this finding in more detail, we analyze individual nanowires with chemically sensitive techniques. Figure 2(d) shows the (radially averaged) Al content along the growth axis for two representative single nanowires as obtained by energy-dispersive x-ray spectroscopy (EDX) performed in our analytical scanning electron microscope (SEM) (see Methods). The inset illustrates the direction of the line scan on an exemplary nanowire. From the bottom to the top of the nanowire, the average axial Al content of the two nanowires increases from 0.1 to 0.7 up to a length of 400 nm, stays then constant for about 300 nm, and decreases from 0.7 to 0.3 for the top 500 nm of the nanowire. Given that the length of the (Al,Ga)N segment amounts to only 835 nm, the initial increase of the average Al content for the first 400 nm is likely caused by the formation of an AlN shell (with decreas-
ing thickness toward the bottom) wrapping around the GaN base. The average Al content in proximity of the GaN disk is about 0.35, which is in fact close to the nominal value. The presence of this axial gradient in the radially averaged Al content cannot explain the constant, low emission energy observed along the (Al,Ga)N segment [Fig. 1(c)]. To resolve also the radial composition, we utilize atom probe tomography (APT). Figure 2(e) shows the Ga and Al profiles across the diameter of a single nanowire in proximity of the GaN disk extracted from full three-dimensional APT data. Several profiles have been extracted at different heights of the same nanowire (not shown), and they are all consistent with the profiles shown in Fig. 2(e), which demonstrates the presence of an Al-deficient inner core and a complex Al-rich shell with a continuously varying Al content. Figure 2(f) shows the Al content across a two-dimensional section of the same nanowire in proximity of the GaN disk. The inner blue region represents the Al-deficient core, which is surrounded by a thin shell with high Al content and an outer shell with lower Al content. The core appears compressed in the APT image due to well-known image aberration effects. When correcting for the compression, the dimensions of the core would change slightly but the anisotropic shape would remain. However, the exact shape of the core may be affected by the dependence of the evaporation rate of (Al,Ga)N on the Al content, which may potentially result in a change of the curvature of the surface of the specimen during evaporation. Note that the cation concentrations given in Figs. 2(e) and 2(f) correspond to the raw measured data without correcting for the loss of N atoms common in APT, which is why the sum of the Al and Ga content does not equal one. The actual concentrations will thus be slightly higher. Considering this fact, the Al content in the core should be close to the value of 0.15 deduced from the transition energy in the PL and CL experiments shown in Fig. 1.

Combining the results of APT with EDX and assuming that the Al content in the core is constant along the length of the (Al,Ga)N segment as suggested by the uniform emission energy observed in CL [Fig. 1(c)], we deduce that the shell essentially consists of pure AlN at the bottom of the (Al,Ga)N segment [i.e., at about 400–500 nm in Fig. 2(d)]. Subsequently, the shell composition decreases toward the nanowire top to a value of about 0.4 close to the position of the disk at the top. Figure 3(a) visualizes the resulting complex core/shell structure that forms spontaneously during growth despite a constant temperature and constant fluxes, much like reported previously by Allah et al. for 300 nm short (Al,Ga)N nanowire segments grown by PAMBE. The drastic deviation from the intended uniform alloy was ascribed to the different diffusion lengths of Ga and Al on the nanowire sidewalls and the geometric shadowing effect occurring during the growth of dense nanowire ensembles. An axial gradient of the Al content in an (Al,Ga)N segment followed by a constant composition after 200 nm of growth was also reported by Pierret et al. In fact, growth would be expected to approach a steady-state regime—leading to a radial composition profile that does not change anymore along the nanowire axis—once the nanowire diameter and density have become constant. In the present case, Fig. 2(d) clearly shows that we do not reach this steady-state conditions despite the considerable length of the (Al,Ga)N segment. The origin of this extended growth under non-stationary conditions is the one order of magnitude lower nanowire density compared to the ensembles studied in Refs. 31 and 32 for the density, see Ref. 33, the correspondingly weaker influence of shadowing, and the resulting continuous tapering of the nanowires (see the top- and sideview secondary electron micrographs in the Supporting Information).

The complex core/shell structure has important consequences for the electronic and optical properties of the nanowires, which have not been elucidated so far. In particular, due to the spontaneous polarization of the group-III nitrides, the compositional grading of the (Al,Ga)N segment along the nanowire axis induces a bulk polarization charge, which in turn leads to the accumulation of mobile
charge carriers of the opposite sign. This phenomenon, known as polarization doping,\textsuperscript{34} has been exploited for $n$-type as well as $p$-type doping of GaN/(Al,Ga)N heterostructures,\textsuperscript{35} and is the rationale for the nanowire diode design proposed in Refs.\textsuperscript{36} and \textsuperscript{37}.

Obviously, the radial and axial compositional gradients as well as the effects of polarization doping are essential for an understanding of the radiative transitions observed in PL and CL [cf. Figs. 1(b) and 1(d)]. The low energy of the emission band attributed to the (Al,Ga)N segment, and the absence of any signal above 4.1 eV either in the PL [Fig. 1(b)] or CL spectra (not shown here) is a direct consequence of the nanowires' core/shell structure in conjunction with an efficient transfer of photo- or cathodogenerated carriers from the Al-rich shell into the Al-deficient core. Likewise, electrons induced in the shell by polarization doping transfer to the core and from there to the GaN quantum disk, thus screening its internal electrostatic field. This charge accumulation potentially accounts for the absence of the quantum-confined Stark effect demonstrated by the spectra in Fig. 1(b).

For a quantitative determination of the electron density and the resulting transition energy, a full three-dimensional simulation of the complex core/shell structure would be desirable. However, when attempting to simulate such a core/shell nanowire including the
axial and radial gradients in composition using NextNano™, we found that the solution depends sensitively on details of the structure, such as the precise thickness of the shell and the presence or absence of semipolar facets. To illustrate the physical principles, we thus employ simple one-dimensional simulations \(^{38,39}\) as discussed in the following (for further details, see the Methods section).

To estimate the electron density induced by polarization doping, we consider a planar heterostructure with a 6.5-nm-wide GaN well embedded in an (Al,Ga)N layer with a negative Al gradient along the growth axis (the [000\(\bar{1}\)] direction). Figure 3(b) shows the band profile of this heterostructure where the Al content is assumed to vary linearly from 1 to 0.35 from the bottom to the top of the structure, similar to the variation of the Al content in the Al-rich shell of the top segment of our nanowires containing the GaN quantum disk \([\text{cf. Fig. 2(d)}]\). The negative gradient of the Al content along the [000\(\bar{1}\)] direction leaves a three-dimensional slab of positive bound charge, which is compensated by free electrons that diffuse inward from surface donor states. This polarization-induced doping manifests itself by the change of band gap occurring entirely by a variation of the valence band edge, while the conduction band edge remains close to the Fermi level.

The dashed line in the inset of Fig. 3(b) shows the polarization-induced free electron density in a graded (Al,Ga)N segment without GaN disk to be on the order of \(5 \times 10^{17} \text{ cm}^{-3}\). The solid line in the inset shows the electron concentration in a structure with a graded (Al,Ga)N segment and additional GaN quantum disk, corresponding to the sample whose band profile is shown in the main figure. The charge transfer taking place in this structure results in a degenerately doped GaN disk with an electron sheet density \(n_s\) of \(4 \times 10^{12} \text{ cm}^{-2}\).

To quantitatively determine the impact of this high electron density on the transition energies in the GaN quantum disk, we next calculate the electron and hole states in GaN quantum wells with a width between 1 and 7 nm embedded in Al\(_{0.1}\)Ga\(_{0.9}\)N, the approximate composition of the nanowire core. The calculations are done for undoped wells, for which the polarization results in an internal electrostatic field of around 1 MV/cm,\(^{24}\) and for wells with an electron density as determined by the simulations discussed above. We also take into account the compressive strain in the GaN quantum well imposed by a coherent interface with the Al\(_{0.1}\)Ga\(_{0.9}\)N layer. Figure 3(c) shows the transition energies for these two cases together with the experimental peak energies of the low-energy PL bands obtained by PL spectroscopy at 10 K (not shown here). The transition energies for the undoped wells are governed by the quantum confined Stark effect, i.e., they depend linearly on width with a slope given by the internal electrostatic field. As expected, they fall below the GaN band gap (3.504 eV) for a well width exceeding 2.5 nm. In contrast, the quantum wells with an electron density equal to that induced in the disks by polarization doping in the shell exhibit a transition energy above the GaN band gap even for the widest well, in close agreement with the experimental results.

The insets of Fig. 3(c) illustrate that the screening of the internal fields increases the electron-hole overlap in the quantum disk. Whereas electrons and holes are spatially largely separated in the undoped disk (bottom left), they have an appreciable overlap for the disks with a high electron sheet density (right top). This enhanced electron-hole overlap is expected to affect the recombination dynamics of the electron-hole pairs within the GaN disk and consequently the internal quantum efficiency. Even more importantly, carrier recombination in these structures is affected not only quantitatively, but also qualitatively: the electron density in the disk is well above the Mott density,\(^{40,41}\) and recombination is thus no longer excitonic, but occurs between the two-dimensional electron gas and photoexcited holes.

To investigate the consequences of this high electron density for the recombination dynamics in the GaN disk, we perform time-resolved PL measurements on the nanowire
ensemble with the 6.3 nm-thick GaN disk at temperatures between 5 and 300 K. The disks were excited directly using an excitation wavelength of 325 nm, corresponding to an energy below the bandgap of the (Al,Ga)N core. The PL transients presented below are thus not affected by carrier transfer processes.

Figure 4(a) shows a representative streak camera image at 10 K. The emission band of the GaN quantum disk is observed to redshift by 30 meV during the initial 4 ns of the PL decay. For polar GaN/(Al,Ga)N quantum wells, such a temporal redshift may occur as a consequence of a dynamical screening of the internal electrostatic fields induced by a high excitation density.\(^{42,43}\) In the present case, this phenomenon can be safely ruled out, since the photogenerated carrier density \(\Delta n\) inside the disk is at most \(2 \times 10^{10} \text{ cm}^{-2}\), i.e., more than two orders of magnitude smaller than the electron sheet density resulting from polarization doping \((n_s \approx 4 \times 10^{12} \text{ cm}^{-2})\).

Another possible reason for this gradual redshift is the progressive relaxation of photoexcited excitons within a band of localized states, leading to their spectral diffusion.\(^{44,45}\) This phenomenon is most frequently observed in ternary layers and quantum wells, for which compositional fluctuations induce the disorder responsible for exciton localization.\(^{46-48}\) In the present case, compositional fluctuations in the barriers or steps at the GaN/(Al,Ga)N interfaces could potentially induce localization as well.\(^{49}\) However, the transfer of carriers from higher to lower energy states manifests itself in a characteristic temporal delay of the emission at lower energies, which is not observed here. Instead, the rise time of the emission is found to be the same regardless of energy. We also do not observe any other signature of localization effects, such as the typical S shape of the transition energy with temperature (not shown).\(^{50-52}\)

An alternative explanation for the spectral shift suggests itself when considering that we are not dealing with a spatially uniform emitter, but a large ensemble of individual emitters (in the present experiments, we excite about \(10^4\) nanowires). In other words, the temporal redshift may simply result from a spectral superposition of GaN quantum disks emitting at different energies. In fact, for the present nanowire ensembles with a spatially random arrangement, it is not conceivable that the Al gradient in the shell is identical for each nanowire. Differences must occur because of both shadowing effects\(^{51,53}\) and the different diameters of individual nanowires, affecting the diffusion of the group-III adatoms on the nanowire sidewalls. These differences in the Al gradient in turn result in different electron densities induced by polarization doping. The higher the electron density in the quantum disk, the lower the internal electrostatic field due to screening, which translates into higher emission energies and shorter lifetimes. The most heavily doped quantum disks thus dominate the high-energy emission at short times. In more lightly doped disks, the residual field leads to longer lifetimes and lower emission energies, which lead to the long-living and redshifted tail of the emission. The net result of this superposition of an ensemble of disks with different transition energies and lifetimes is the apparent spectral shift of the emission band as seen in Fig. 4(a).

Figure 4(b) shows PL transients spectrally integrated over the entire emission band between 5 and 300 K. The transients are characterized by a continuously changing slope on a semilogarithmic scale and can be adequately described only by the sum of at least three exponentials. This finding is in full agreement with the hypothesis discussed above: the spectrally integrated emission is a superposition of the individual (exponential) transients of nanowires with different electron densities, and thus different lifetimes. Approximating the fast and slow components of the transient at 5 K by single exponentials, we obtain decay times of about 0.4 and 4 ns, respectively. For comparison, planar GaN/(Al,Ga)N quantum wells of similar width and average doping density were found to exhibit a lifetime of 2 ns.\(^{53}\) The maximum electron density in our nanowire ensemble is thus significantly higher than the average value of \(n_s\) determined from the average transition energy.
Figure 4: Recombination dynamics of the nanowire ensemble with a 6.3 nm thick GaN disk. (a) Streak camera image obtained upon pulsed excitation at 325 nm and 10 K. The dashed line highlights the spectral diffusion of the emission band from the GaN quantum disk. (b) Spectrally integrated PL intensity transients obtained from streak camera images acquired at temperatures between 5 and 300 K. (c) Temperature dependence of the radiative lifetime determined from the peak intensity of the transients (solid circles). The dashed lines show the calculated temperature dependence for electron sheet densities in the quantum disk of 0.2, 0.6, and $1.6 \times 10^{12}$ cm$^{-2}$ (from left to right). The solid line represents the radiative lifetime of a nanowire ensemble with an ensemble distribution of the electron volume density as shown in the inset.

For temperatures above 20 K, the decay is seen to accelerate accompanied by a continuously decreasing integrated intensity [cf. Fig. 4(b)], revealing an increasing participation of nonradiative decay channels (see also the comparison of resonant and nonresonant excitation in the Supporting Information). To separate radiative and nonradiative channels, we examine the peak intensity of the transients, which is directly proportional to the radiative decay rate.\textsuperscript{54-56} Figure 4(c) shows the temperature dependence of the radiative lifetime determined in this way. The radiative lifetime starts to increase already at temperatures as low as 10 K, but does not enter the linear temperature dependence characteristic for nondegenerate two-dimensional systems in general and GaN/(Al,Ga)N quantum wells in particular.\textsuperscript{57,58} To quantitatively understand this peculiar temperature dependence, we calculate the spontaneous recombination rate using Fermi’s golden rule for the general case of Fermi-Dirac statistics valid also for the high carrier density in the GaN quantum disks (for details, see the Supporting information).\textsuperscript{59-61} Note that this calculation ignores any residual electric fields in the disk. Consequently, we do not compare the absolute transition rate in the following, but only its temperature dependence.

The three dashed lines in Fig. 4(c) show the inverse radiative rates obtained by our calculations for electron sheet densities of $n_s = 0.2$, 0.6, and $1.6 \times 10^{12}$ cm$^{-2}$. In contrast to the strictly linear dependence of a nondegenerate quantum well, the radiative rate for high electron sheet densities is constant at low temperatures, for which the system becomes degenerate. The higher the electron density, the higher the temperature up to which degeneracy prevails, and thus the larger the temperature range of constant radiative lifetime (for more details, see the Supporting Information).

The theoretical dependence for an electron density of $n_s = 6 \times 10^{11}$ cm$^{-2}$ is closest to the experimental data, but it does not describe them well, particularly at low temperatures. The agreement can be much improved when allowing for a distribution of the electron density as qualitatively discussed above in the context of Figs. 4(a) and 4(b). Quantitatively, we obtain the solid line in Fig. 4(c) by assuming a log-normal distribution of $n_s$ with a mean of $2 \times 10^{12}$ cm$^{-2}$ or a volume
density of $3 \times 10^{18} \text{ cm}^{-3}$ as shown in the inset of Fig. 4(c), not too far from the average value estimated above as a result of polarization doping ($n_s = 4 \times 10^{12} \text{ cm}^{-2}$). This agreement suggests that despite the crudeness of our approach concerning the energy and the lifetime of carriers in the GaN quantum disks in our nanowires, we have reached a fairly coherent understanding of both the static and dynamical electronic properties of these disks. In particular, the modest thermal quenching of the emission from the GaN quantum disks (see Supporting Information) is understood to be a result of the short radiative lifetime due to the high electron density and not a particularly high material quality.

To summarize and conclude, we have shown that, during the synthesis of GaN/(Al,Ga)N nanowire ensembles by molecular beam epitaxy, a complex core/shell structure forms spontaneously, characterized by a continuously varying Al composition both along the growth axis and across the diameter of the nanowires. This phenomenon has been observed in similar form by other groups, which shows that it is not caused by specific growth conditions, but the fundamental growth mechanisms of (Al,Ga)N nanowires by PAMBE. The peculiar structure resulting from this self-assembly process has important consequences for the optoelectronic properties of the nanowires. First of all, the axial Al gradient in the Al-rich shell induces a three-dimensional polarization charge, leading to the accumulation of electrons in the shell. Second, due to the Al deficient core, these electrons in the shell are efficiently transferred to the GaN quantum disk. The resulting high electron sheet density inside the GaN disk screens the internal electrostatic field, effectively canceling the quantum-confined Stark effect, and manifesting itself in both transition energies almost independent of thickness and short radiative lifetimes up to room temperature that result in a comparatively high internal quantum efficiency. The latter is obviously attractive for the realization of efficient light emitting devices in the ultraviolet range. However, we emphasize that for taking advantage of the effects reported here in any application, one would have to achieve a high level of control over the formation of the compositional gradients. To reach this aim, further studies are required to fully understand the mechanisms leading to the self-assembly of (Al,Ga)N nanowires in PAMBE. Moreover, this compositional self-assembly is likely to occur also for other mixed cation nanowires synthesized by diffusion-controlled growth techniques such as MBE. Hence, our findings may be relevant not only for deep UV emitters based on (Al,Ga)N nanowires, but for example also for MBE-grown (In,Ga)N nanowires for emission in the visible spectrum and solar energy harvesting.

**Methods**

**Nanowire synthesis**

The GaN/(Al,Ga)N nanowire ensembles are synthesized utilizing a DCA Instruments P600 molecular beam epitaxy system equipped with two radio frequency plasma sources for active N and solid-source effusion cells for Al and Ga. First, GaN nanowires are grown on Si(111) substrates relying on their spontaneous formation on this substrate under suitable experimental conditions. A two-step growth approach is employed in order to minimize the incubation time. After the formation of a GaN nanowire ensemble with a nominal length of 750 nm, the Ga flux is reduced and an Al flux is added for the synthesis of a nominally 800 nm long (Al,Ga)N segment with a nominal Al content of 0.3 using the same V/III ratio as for the GaN nanowires underneath. For the formation of a GaN nanowire ensemble with a nominal length of 750 nm, the Ga flux is reduced and an Al flux is added for the synthesis of a nominally 800 nm long (Al,Ga)N segment with a nominal Al content of 0.3 using the same V/III ratio as for the GaN nanowires underneath. For the formation of the GaN quantum disk, the Al shutter is simply closed for a duration corresponding to the intended disk thickness. Finally, the disk is capped by a nominally 35 nm long (Al,Ga)N segment. Three samples are grown in this way, differing only in the thickness of the GaN quantum disk with nominal values of 1.8, 3.3, and 6.3 nm. A sample without quantum disk, but otherwise identical (Al,Ga)N nanowires on a GaN base
serves as comparison. The nanowires thus obtained have an average length of 1.55 µm (close to the expected value) and an average equivalent disk diameter at their top of 125 nm. The mean density of nanowires in the ensemble is $3 \times 10^9$ cm$^{-2}$. Top- and sideview secondary electron micrographs of the sample with a 6.3-nm-thick disk are shown in the Supporting Information.

**Morphology and microstructure**

The morphological properties of the as-grown GaN/(Al,Ga)N nanowires are studied by scanning electron microscopy carried out in a Hitachi S-4800 field emission microscope using an acceleration voltage of 5 kV (see Supporting Information). Single nanowires in cross-sectional specimens are investigated by transmission electron microscopy. The cross-sections are prepared by mechanical grinding and dimpling followed by Ar-ion beam milling down to electron transparency. Scanning transmission electron microscopy is performed by using a JEOL 2100F field emission instrument equipped with a bright-field and a dark-field detector and operated at 200 kV. High-angle annular dark-field micrographs of the nanowires reflect the chemical contrast between the GaN quantum disk and the surrounding (Al,Ga)N segments.

**Composition**

The composition of single nanowires is investigated by energy dispersive x-ray spectroscopy and atom probe tomography. The former is performed using an EDAX Apollo XV silicon drift detector attached to a Zeiss Ultra55 field emission scanning electron microscope operated at 5 kV. The latter is carried out using a Cameca LEAP 4000X-HR system. Single nanowires from the ensemble with the 6.3 nm thick quantum disks are isolated in an FEI Nova Nanolab 600i. Welds are created by electron-induced metal deposition of Pt or Co. The selected nanowire is mounted on a tip, and the evaporation of atoms is triggered by a laser generating picosecond pulses at a wavelength of 355 nm. Since the (Al,Ga)N nanowire absorbs only weakly at this wavelength, a comparatively high pulse energy of 100 pJ is used. The field of view probed by the technique is about 40 nm.

**Spectroscopy**

For the investigation of the optical properties of the nanowire ensembles, the as-grown samples are mounted onto the cold finger of a liquid He cryostat allowing continuous control of the sample temperature between 10 and 300 K. Cathodoluminescence spectroscopy is carried out at room temperature with a Gatan MonoCL4 system fitted to a Zeiss Ultra55 field-emission scanning electron microscope operated at 5 kV with a probe current of 0.7 nA. The signal is spectrally dispersed by a monochromator and detected using a photomultiplier tube. Continuous-wave photoluminescence spectroscopy is performed by exciting the ensembles with the 244 nm (5.081 eV) line of a Coherent Innova 90C FreD Ar$^+$ ion laser focused to a spot of about 0.5 µm diameter yielding an excitation density on the order of 1 W/cm$^2$. The signal is spectrally dispersed by a monochromator and detected with a cooled charge-coupled device array. Time-resolved PL measurements are performed by exciting the ensemble with the second harmonic (325 nm) of 200-fs-pulses from an APE optical parametric oscillator synchronously pumped by a Coherent Mira 900 Ti:sapphire laser, which itself is pumped by a Coherent Verdi V10 frequency-doubled Nd:YVO$_4$ laser. The pulses with an energy of 4 pJ are focused onto the sample with a 75-mm plano-convex lens to a spot with a diameter of 30 µm. The transient PL signal is dispersed by a monochromator and detected by a Hamamatsu C5680 streak camera with a temporal resolution set to about 50 ps.

**Simulations**

The band profiles and electron densities are obtained by one-dimensional Schrödinger-Poisson simulations performed
with 1D Poisson. For the polarization of Al$_x$Ga$_{1-x}$N, we assume a value of $-2 \times 10^{-6}$ (1 + 2.1$x$ + 0.475$x^2$) C/cm$^2$ (the negative sign is due to the fact that the nanowires grow in the [0001] direction). This value is about half of that predicted theoretically, but in agreement with the majority of experimental results.

For the conduction band offset between GaN and (Al,Ga)N, we take the value reported by Tchernycheva et al. The spontaneous recombination rate is calculated from Fermi’s golden rule assuming $k$ conservation as $R_{sp} = R_0 \int_0^{\infty} \rho_r(E) f_c(E)[1 - f_v(E)] dE$, where $\rho_r(E)$ is the reduced density of states, $f_c(E)$ and $f_v(E)$ are the Fermi-Dirac distributions for the conduction and the valence bands, respectively, and $R_0$ is a prefactor. Details of this calculation are presented in the Supporting Information.

**Supporting Information Available**

The following files are available free of charge. Top- and sideview secondary electron micrographs of the sample with a 6.3-nm-thick disk, temperature dependence of the integrated photoluminescence intensity for excitation wavelengths of 325 and 244 nm, calculation of the spontaneous recombination rate.

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