GaN Nanowires Grown by Molecular Beam Epitaxy

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(Invited Paper)

Abstract—The unique properties of GaN nanowires grown by molecular beam epitaxy are reviewed. These properties include the absence of residual strain, exclusion of most extended defects, long photoluminescence lifetime, low surface recombination velocity, and high mechanical quality factor. The high purity of the nanowires grown by this method allows for controllable ntype doping. P-type doping presents more challenges but has been demonstrated in active light-emitting diode devices. The present understanding of nucleation and growth of these materials is also reviewed.

Index Terms—Crystal growth, nanotechnology, photoluminescence, semiconductor materials.

I. INTRODUCTION

S EMICONDUCTOR nanostructures have opened many new paths for device technology and fundamental studies. This is particularly true for (Al, In, Ga)N nanowires because of the difficulty in growing large single crystals in this alloy family to serve as substrates for epitaxial growth. Nanowire growth is, therefore, one of the few methods available to produce lowstrain, defect-free material.

This review describes the unique properties of GaN nanowires and their related alloys when grown by catalyst-free molecular beam epitaxy (MBE). Many of their properties stem directly from the growth method. Unlike catalyst-based methods commonly used to grow nanowires, MBE growth of GaN nanowires is driven by differences in surface energies, sticking coefficients, and diffusion coefficients on different crystal planes. The growth proceeds at high temperature and with low growth rate, allowing the crystals to fully relax and achieve a structure that annihilates crystalline defects. The high purity of the starting materials and ultrahigh vacuum environment prevents incorporation of chemical impurities.

The earliest reports of GaN nanowire growth by this method appeared from Sophia University, Tokyo [1], [2] and the Universidad Politécnica, Madrid [3], [4]. Since then, morphology and optical property descriptions have been reported by numerous other groups [5]–[9]. Electrical characterization and device re-

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sults are more infrequent, but they include single nanowire field effect transistors (FETs) [10] and light-emitting diodes (LEDs) made of ensembles of nanowires [11]. More recent study will be discussed in the main sections of this paper, along with the growth mechanism developments over the past several years.

GaN nanowires can also be grown by other mechanisms, though the quality of this material varies widely and in many cases has not been fully explored with sensitive methods such as low-temperature photoluminescence (PL) or carrier lifetime measurements. Although this study in general falls outside the scope of this review, excellent device results have been demonstrated with catalyst-grown GaN nanowires by use of organometallics vapor-phase epitaxy (OMVPE) for LEDs [12] and FETs [13]. LEDs have also been fabricated from ensembles of GaN nanowires grown by catalyst-free OMVPE with patterned substrates [14] and by hydride vapor-phase epitaxy (HVPE) [15]. Other crystal growth methods that have demonstrated GaN nanowire morphology include metalorganic MBE [16], OMVPE with anodic alumina templates [17], direct reaction [18]-[21], catalyst OMVPE [22], [23], and HVPE [24]–[29]. Because these materials have been defined more by their morphology than by their electrical or optical properties, what we refer to in this study as GaN nanowires have also been variously called GaN nanorods, nanocolumns, and nanowhiskers.

The broader field of nanowire growth includes many other materials and growth methods. The reader is referred to [30] for a review of inorganic nanowire growth, [31] for a highlevel review covering many materials mostly grown by catalyst methods, and [32] for a semiconductor nanowire review covering mostly catalyst and direct growth. An overview of catalystbased OMVPE nanowire growth is provided in [33].

II. GROWTH

A. Growth Parameters and Morphology

GaN nanowires grown by catalyst-free MBE form exclusively in the wurtzite (or hexagonal) crystal structure [34] with the growth axis parallel to the [0 0 0 1] crystal direction, also called the *c*-axis. The sidewalls of the nanowires conform to the $\{1 0 0\}$ family of *m*-planes. When the nanowires are well separated, their cross sections will be near perfect hexagons, as shown in the field-emission scanning electron microscopy (FESEM) images in Fig. 1(b). The nanowires generally have Ga-face polarity, i.e., the Ga–N bonds parallel to the growth axis are oriented such that Ga is nearest to the root of the bond and N is nearest to the tip [35], [36], although one study found mixed polarity [37].The Ga-face polarity in GaN films has generally been identified as the smoother surface, but there has been some controversy



Fig. 1 Examples of GaN nanowire morphology for well-separated nanowires. (a) and (c) Top and side views of the same growth run (66 h growth). (b) and (d) From a different run with higher Ga flux (72 h growth).

in measurements on bulk crystals [38]. Si(1 1 1) is the most commonly used substrate, and the nanowires will predominantly align azimuthally to the underlying Si such that the GaN $\langle 1 1 \overline{2} 0 \rangle$ directions are parallel with the Si $\langle 1 \overline{1} 0 \rangle$ directions [8], [39]. Thin AlN buffer layers are commonly used to prevent reactions between Ga and Si and to increase the uniformity of the nanowire nucleation process [35], [40]. GaN nanowires have also been grown on sapphire [1], [41] and Si(1 0 0) substrates. Growth on Si(1 0 0) leads to less regular morphology [40], [42] but has been shown to have PL similar to material grown on Si(1 1 1) [43], [44].

GaN nanowires typically form under MBE growth conditions of high active N flux relative to Ga flux and high substrate growth temperature, 740-830 °C being typical of the ranges reported. Group III elements (Ga, Al, In) are supplied by direct evaporation from molten metal in standard Knudsen cells. The N source is a gas injector with active excitation of the gas, typically provided by maintaining a plasma with radio-frequency (RF) excitation. This plasma-assisted or RF-assisted N₂ stream consists of a mixture of atomic N and excited molecular states that react readily with Ga at the growth surface. The beam can also contain ionized species, although many sources include biased plates designed to deflect any charged species from reaching the substrate. The relative abundances of these species vary with RF power, gas flow rate, plasma pressure, and details of the plasma tube and confinement fields. Optical emission from the plasma is frequently used as a semiquantitative measure of the relative abundance of these species [45], but aperture design and bias plates also influence the composition of the N flux that actually reaches the growing material. The variability in N_2 source operation coupled with the difficulty in monitoring the species produced is a major factor in the variability of results from different growth systems and long-time-period drift within a single system.

As described succinctly in [46], the morphology transitions from compact films to nanowires to the absence of growth from



Fig. 2. Phase diagram for GaN nanowire formation at fixed N₂ plasma conditions. Reprinted with permission from [46]. Copyright 2009, American Institute of Physics.

complete reevaporation of Ga. The transition temperatures depend on the Ga flux and N flux, as shown in the phase diagram given in Fig. 2. Within the nanowire (nanocolumn) growth regime, the density and diameter of nanowires can be influenced by changes in the Ga flux, substrate temperature and N₂ plasma conditions. Nanowire density is strongly affected by buffer layers and initial growth conditions. The amount of material that grows between the nanowires, which we call the matrix layer, also varies, from a relatively thick, faceted layer, such as that illustrated in Fig. 1, to a dense layer with no apparent matrix but irregular nanowire shapes [47] to very sparse nanowires with no matrix [39]. The matrix layer is often quasi-columnar, making the distinction between it and the nanowires difficult to identify in some cases. Additional examples of MBE GaN nanowire morphologies can be found in [48] and [49].

The Ga flux and substrate temperature also present some measurement challenges. Although most MBE systems are equipped with a beam flux monitor, the sensitivity of these gauges varies over time due to accumulation of deposits on active components. Changes in sensitivity by a factor of 2 have been reported [42]. More accurate results can be obtained by measuring the Ga flux under identical cell conditions but with the substrate held at lower temperatures where the Ga sticking coefficient can be assumed to be 1. The Ga flux can be calculated from in situ optical reflectance measurements [4], ex situ layer thickness measurements [50], or GaAs reflection high-energy electron diffraction (RHEED) intensity oscillations (if the MBE system has GaAs growth capability) [40]. The measurement of substrate temperature in MBE systems has a long history. The most accurate measurements at these high temperatures are typically achieved with optical pyrometry. The sample emissivity must be known for accurate calibration [51], and changes in emissivity during growth can lead to a systematic drift in the calibration. Using a calibrated pyrometer that collects light from the backside of the



Fig. 3. Schematic of growth propagation mechanism for GaN nanowires grown by MBE. The vertical growth is driven largely by the much lower sticking coefficient for Ga on the sidewall *m*-planes relative to the top surface *c*-planes.

substrate, where no deposition occurs, we have been able to estimate that substrate temperature can be inaccurate by as much as $15 \,^{\circ}C$ when uncorrected for changes in surface emissivity.

B. Nucleation and Growth Mechanisms

Because of the prevalence of catalyst-based nanowire growth, early workers in the field of GaN nanowire growth by MBE hypothesized that the growth proceeded through the formation of nanoscale Ga droplets acting as self-catalysis particles [49], [52]. A large body of evidence has accumulated in favor of other explanations, however, particularly because Ga droplets were never observed on nanowire tips. The high growth temperatures used for nanowires relative to MBE thin-film growth, the loss of Ga to re-evaporation, slow growth rate, sensitivity to surfactant effects, and consistent crystallography indicated that surface diffusion and variations in sticking coefficients were the primary drivers for growth [53]. This study led to the demonstration of a differential sticking coefficient model [40], [42], [54] of growth propagation, as illustrated in Fig. 3.

Further support for this model comes from recent theoretical predictions of surface energies and diffusion coefficients [55] and tests of growth under unusual conditions, including rapid cooling while continuing growth [42] and intentional Ga droplet formation [48]. As presented so far, this model has not been quantitatively applied to predict growth rates. A complete model would account for the fact that the three-dimensional geometry itself alters the predicted growth rates because the average Ga flux on the top surface is five to six times higher than that on the sidewalls [37], even in the absence of surface diffusion or sticking coefficient variations. There is evidence in [47] and [56] for a significant contribution to nanowire growth from Ga atoms that arrive either at the substrate surface or at the nanowire sidewalls and diffuse to the top surface. These mechanisms predict that thinner nanowires should grow longer than thicker ones. Experimental observations vary on whether this length variation indeed occurs, indicating that there is sufficient variation in growth conditions to argue for aspects of all these models. The length variation is most evident in growth where AlN buffers are omitted, growth periods are short, and there is a significant fraction of nanowires with diameters less than 50 nm.

Nucleation of nanowire growth is typically random, producing a wide range of densities and diameters within a single growth run. This variation increases for comparisons made among different growth systems. Nanowires have been found to nucleate on bare Si with a minimum nucleation diameter of 7 nm [57] to 15 nm [47] and no matrix layer growth. In [35], it was found that nanowires nucleated on AlN buffer layers arose from the top of GaN islands with a density that increased with increasing substrate temperature in the range of 740-800 °C while the nanowire growth rate declined. In another study [48] on nanowire growth on AlN buffer layers, increasing substrate temperature led to larger nanowire nuclei and thereby reduced the nanowire density. Several authors point out that despite the high N₂ flux used for GaN nanowire growth, the N₂ species can limit the vertical growth rates [4], [35], [58]. The N₂ plasma conditions can also affect nanowire nucleation independent of the nanowire growth rate [40]. This study also found that nanowires tended to nucleate inside faceted pits, rather than on the top of islands. A more recent study [59] of the initial GaN nanowire growth on 5-nm AlN layers found several stages of island formation and strain relaxation. High-resolution TEM of nanowire roots frequently shows thin layers of amorphous material, possibly SiN_x , between the substrate and the crystalline region of the nanowire [36], [48], [57]. The variety of possible seed crystal shapes may in part explain why so much variation is observed over relatively small ranges in growth parameter space.

C. Selective Epitaxy

The difficulty in controlling nanowire nucleation has led many workers to explore more complex nucleation schemes. In [11], a method of intentional generation of GaN seed crystals at low growth temperature prior to nanowire growth is described. Patterned epitaxy, in which substrates are masked with patterned layers that limit growth to specific areas, is well known in OMVPE growth and has been used for GaN nanowire growth in OMVPE [60], [61]. Because elemental Ga is more reactive with most mask surfaces, it has taken longer to develop selective epitaxy processes for MBE. Initial attempts with SiO₂ masks [4] and Al pedestal regions [62] produced differences in nucleation density but not true selectivity. Better results have been obtained with Ti masks that are intentionally converted to TiN_x in situ, and this method has proved critical in obtaining consistent diameters in nanowires for LEDs (see Fig. 4). Without the diameter control afforded by the selective epitaxy strategy, the In incorporation varies from nanowire to nanowire and the emission wavelength shifts. These experiments also suggest that optical emission from the InGaN disks is most intense when the InGaN quantum well grows on tilted facets rather than as a c-plane disk perpendicular to the growth axis. The faceted morphology avoids spontaneous polarization fields that separate electrons and holes spatially within a *c*-plane quantum well.



Fig. 4. Bird's eye view SEM and emission images excited by HeCd laser from InGaN/GaN nanowires with diameters as shown. The In incorporation increases with increasing nanowire diameter. Reprinted with permission from [2]. Copyright 2010, American Institute of Physics.



Fig. 5 Selective epitaxial of GaN nanowires through openings in a SiN_x mask covering an AlN buffer layer. The cross sections are equilateral hexagons with diameters that track the size of the mask opening.

The selectivity with TiN_x masks relies on close spacing of the openings to prevent random nucleation on the mask. Nearperfect selectivity, even in large mask regions, has recently been demonstrated with SiN_x masks on AlN buffer layers, as shown in Fig. 5. Details of this result are given in [63].

D. Crystalline Perfection

GaN nanowires grown by MBE are typically free of strain because strain mismatch between the substrate and the growing GaN can readily be relieved by dislocation formation perpendicular to the $(0\ 0\ 1)$ planes. The small diameter of the wire and ease of dislocation glide within the *c*-plane allow annihilation of the dislocations at the nanowire surface, leaving the crystal to grow defect-free after a few nanometers of axial growth [64]. Recent studies [59] using RHEED have identified several morphological steps in the nucleation of the nanowires and confirm the confinement of strain relaxation to dislocation formation



Fig. 6. Bright-field TEM images of GaN nanowires. (a) Bottom portions of two nanowires near their roots (indicated by block arrows); note occasional basal plane stacking faults (horizontal arrows). (b) Defect-free portion of a nanowire near its tip (indicated by block arrow); note band-like contrast associated with nanowire bending. (c) Middle portion of two coalesced nanowires; note a continuous array of dislocations at the NWs interface (indicated by vertical arrows). Scale bar is 100 nm. All images were recorded near the [1 1 0] GaN zone axis orientation. [86]

and annihilation very close to the nanowire-substrate boundary. The perfection of the crystalline structure may be lost if the nanowires coalesce during growth [65], [66]. Fig. 6(a) and (b) shows TEM images of nearly defect-free nanowires and a chain of nanowire defects formed as a consequence of coalescence of two nanowires. Screw dislocations have been observed in GaN nanowires grown on sapphire [67]. Although this observation is not common, elastic theory predicts that screw dislocations are in fact stable if formed near the center of a thin rod [68]. It is problematic to rely on TEM alone for defect identification because of the small number of samples that can be conveniently tested, and the need to examine different diffraction conditions and sample tilts to distinguish defects from bending contrast. Optical measurements, described in the next section, also provide evidence for the high crystalline quality of these materials.

By comparison, the structural properties of *a*-axis catalystgrown GaN nanowires are very different [69]. These nanowires are oriented such that dislocations tend to propagate along the nanowire length. Basal plane (*c*-plane) stacking faults are also common, leading to regions of the crystal with a structure that more closely resembles that of cubic GaN. The high speed of catalyst-based growth and the lower purity environments in which such growth often takes place can lead to high defect incorporation both from crystallographic imperfections and chemical impurities [70], [71]. These effects can be mitigated to some degree by employing the lowest possible growth rates and using high-purity environments such as MBE and OMVPE. A recent comparison [72] of catalyst versus catalyst-free MBE nanowires found that even under the same growth conditions, catalytic growth produced material with a high density of basal plane stacking faults, as determined by TEM. The PL intensity was also lower for catalyst-grown material, but the spectrum was free of yellow luminescence. OMVPE materials consistently contain a higher impurity background, presumed to be carbon, especially apparent in material grown at lower substrate temperatures. It is not clear, however, if carbon incorporation is necessarily detrimental to device operation provided that the concentration is below the point where significant subgap luminescence is observed [73].

E. Heterojunctions and Alloys

Nanowires of purely binary InN [74]-[76], AlN [77], and alloys of these compounds with GaN have been grown by MBE and other methods. As illustrated in the previous section, the addition of InGaN quantum disks in GaN nanowires [11], [78] is particularly useful for optoelectronic devices to enhance luminescence and tune the wavelength of the emission. Compelling results have also been obtained for GaN disks in AlGaN wires [79]. In [4], a comprehensive overview of these types of heterostructures is given, and we mention here just some highlights of this study. The interfaces between the AlGaN and GaN are atomically abrupt, and the quantization of the energy states in the quantum disks has been established [80]. Bragg reflector stacks of alternating AlN and GaN layers were also grown, demonstrating the ability of the nanowire structure to withstand strain without cracking [79]. One important outcome of this investigation was the determination that variations in strain between the core of the nanowire and its outer surface could lead to changes in the band structure that suppress radiative recombination of electrons and holes by spatially separating the two carrier types [81].

In Fig. 7, we illustrate another type of heterostructure, a coresleeve p-n junction. This structure was fabricated by coating n-type MBE-grown nanowires with p-type layers grown with HVPE. The image contrast depends on electron beam energy and likely arises from differences in work function between the p-type and n-type regions. The exact correlation between the physical junction and the boundary imaged by FESEM has not been established.

III. OPTICAL PROPERTIES

A. Photoluminescence (PL)

Investigation of the optical properties of MBE-grown GaN nanowires provided some of the earliest evidence for their exceptional quality. Room-temperature PL intensity from these nanowires usually compares favorably with that from reference



Fig. 7. FESEM images of core-sleeve *p-n* junctions formed by overgrowth of p-type HVPE GaN on MBE-grown GaN nanowires [87].



Fig. 8. Low-temperature photoluminescence of a single GaN nanowire showing that the spectrum consists of only exciton transitions and phonon replica peaks of free exciton transitions. The donor-bound exciton emission is shifted from the unstrained position, marked with the dashed line at 3.472 eV, by strain applied to the nanowire by differential thermal contraction of the nanowire relative to its substrate [88].

GaN films [53], [82], though the variety of quality among GaN materials prevents definitive comparisons. The low-temperature PL spectrum generally consists solely of donor-bound excitons, free excitons, and the associated phonon-mediated replicas of these transitions (see Fig. 8). In nanowires with radii below 40 nm, a two-electron satellite peak has also been reported [44] and its enhanced intensity attributed to interactions between the exciton pairs and nearby surfaces. For isolated nanowires, the donor-bound exciton D^0X_A peak is found at the strain-free position of 3.472 eV [83]–[85]. These data, in addition to X-ray diffraction on nanowire ensembles [8], [53], demonstrated that the growth mode of the nanowires promoted complete relaxation of lattice mismatch early in the nucleation stage without propagation of dislocations along the nanowire axis.

The absence of strain broadening allowed exceptionally highresolution spectroscopy relative to that obtainable from GaN specimens grown by other methods, so polarization anisotropy of the exciton peaks could be monitored from 3.2 to 300 K [83].



Fig. 9. Linear fit of PL decay time vs. nanowire diameter extracts the surface recombination velocity for this set of nanowires (from the same growth run).

Low-temperature PL peaks as narrow as 300 μ eV have been reported [89]. Yellow luminescence is generally absent from well-formed nanowires but is occasionally reported for larger, coalesced structures [65], material grown at lower temperatures [49], and specimens that contain matrix layer material [66]. Subgap luminescence in GaN (including yellow luminescence) is indicative of chemical impurities and/or structural defects [90].

The inherently strain-free state of the growth mode of the nanowires in turn permits the monitoring of strain states by monitoring shifts in the PL spectrum. The data in Fig. 8 show how the D^0X_A peak shifts to lower energy because of tensile strain imposed on the nanowire by its adhesion (via van der Waals bonding) to a quartz substrate during cooling to ~5 K. Compressive strain from atomic layer deposition of alumina produced positive shifts [88].

Time-resolved PL data indicate that the dominant recombination processes vary considerably with temperature. The lowtemperature regime is dominated by radiative recombination of bound excitons, and the thermalization of these exciton pairs away from the shallow donor that binds them causes the PL lifetime to decrease initially as the temperature increases [88]. The lifetime begins to increase in the range near 150 K as free exciton recombination becomes dominant, provided that nonradiative recombination processes are saturated. Near room temperature, surface recombination becomes the limit to the PL lifetime, so that thicker nanowires have longer lifetimes (see Fig. 9). Although this limit certainly invites attempts at surface passivation with AlGaN layers and other schemes applied in GaN epitaxial layers, we note that the actual value of the surface recombination velocity, 9×10^3 cm s⁻¹, is quite low compared with that of other compound semiconductor surfaces, e.g., GaAs [91].

This variation in recombination mechanisms over different temperature regimes underscores the need to make careful measurements of lifetime to estimate internal quantum efficiency (IQE). Ratios of room-temperature PL intensity to lowtemperature PL intensity do not compare the same phenomena and, therefore, do not reveal the room-temperature IQE. It cannot be assumed that the IQE is 100% at low temperature, nor is the "bulk" lifetime of the nanowire material likely to control recombination at room temperature. These caveats apply to epitaxial InGaN/GaN films as well [92]. By varying excitation conditions and sample temperature T until a characteristic $T^{3/2}$ dependence is observed in the PL lifetime, we have estimated that the IQE of one of our moderately doped GaN nanowires is 30% at room temperature [93]. This value is for excitation with 266-nm light in 8-ns pulses of 190 FJ/cm², with a 15 kHz repetition rate. The lifetime is limited by surface recombination, but the exact nature of the surface states is undetermined.

B. Cathodoluminescence

Several groups have examined GaN nanowires grown by MBE with cathodoluminescence (CL). This method offers more localized excitation and ready correlation between luminescence and structure [49] [85], [94]. The rapid diffusion of carriers from the absorption site, however, limits the lateral resolution of the method to within a factor of 2 or 3 of what can be achieved with PL. CL has been used to show that subgap luminescence (yellow or blue) is correlated with coalesced regions or other anomalous growth structures [65], [66]. Dispersed GaN nanowires readily acquire static charge in vacuum, causing a reversible decrease in CL intensity [66]. Electron-beam-induced deposits or damage must be considered in interpretation of the data. One interesting application of CL has been to map minority carrier diffusion length in OMVPE-grown GaN nanowires [95] by collecting luminescence with imaging systems or near-field scanning optical microscopy.

IV. ELECTRICAL PROPERTIES

A. Carrier Concentration and Mobility

Because ultrahigh-vacuum MBE technology and high growth temperature are employed in their fabrication, GaN nanowires have very low concentrations of chemical impurities. Their added exclusion of crystalline defects leads to very low background carrier concentrations, on the order of 10^{15} cm⁻³ or less. A wide range of n-type doping concentrations can readily be achieved with Si as an intentional dopant. The nanowire morphology, however, presents significant challenges in unequivocal determination of carrier type and measurement of free carrier concentration N, and drift mobility μ . P-type doping with Mg has been reported by several groups who have fabricated ensemble LEDs, but the absence of ohmic behavior for single p-type nanowires continues to impede true electrical characterization of their properties. This section will therefore be restricted to discussion of n-type nanowires.

Numerous articles have appeared illustrating the use of nanowire FET devices as test structures to infer these quantities [28], [96], [97]. Typical sample geometry consists of a nanowire dispersed onto an oxidized Si substrate with source and drain contacts fabricated at the two nanowire ends. The Si substrate itself is biased to form a "back gate," and the capacitance between the nanowire and the substrate is modeled as an ideal conducting wire embedded in a dielectric material far above a conducting plane. There are a number of systematic errors introduced by this approach [98], [99]. The full three-dimensional

Φ[eV]	$N [\mathrm{cm}^{-3}]$	$\mu [\mathrm{cm}^2 \cdot \mathrm{V}^{-1} \cdot \mathrm{s}^{-1}]$
Batch 1: intentional Si doping		
0.5 - 0.6	6—10 x 10^{17}	600—300
0.4	5—8 x 10^{17}	800—500
Batch 2: undoped with high background		
0.5 - 0.6	9—18 x 10^{16}	300—100
0.2 - 0.3	4—8 x 10^{16}	700—200

morphology cannot be ignored in efforts to reliably fit FET simulations to actual device behavior. These systematic errors can also depend on the nanowire diameter, and thus can confer diameter dependence to the mobility determination where none actually exists. Moreover, little is known regarding charge-trapping effects in gate oxide layers, or nanowire-metal interfaces in Schottky-gated nanowire MESFET devices. Fitted FET parameters (transconductance, threshold, and pinchoff) can thus vary widely. A few examples of nanowire FET gate hysteresis and memory effects have been published [10], [97]. Until better models are developed and processing-induced surface and interface artifacts can be eliminated or accounted for, nanowire transport properties inferred from FET behavior should be viewed with caution.

In efforts to infer transport properties from simpler twoterminal nanowire test structures, we have analyzed steady-state dark conductivity and photoconductivity (PC) in collections of these devices fabricated from common growth runs where the nanowires were chosen to have a distribution of diameters [100], [101]. We have found that modeling resistance as a function of nanowire length, radius, and contact area for sets of ten or more n-type (Si-doped) GaN nanowires from the same growth run gives carrier concentration and mobility results with moderate uncertainty. Key assumptions in this analysis are that the dopant concentration and surface potential are uniform across the sample set, the contact resistivity is known, and the surface potential can be assumed to fall within a range of a few electron volts.

These conductivity data allow us to restrict the space of allowable N and μ given a particular range of upward bending Φ of the conduction band at the nanowire surface. The latter controls the thickness of the surface depletion layer. Numerical results for two GaN nanowire growth runs are given in Table I. Disagreement exists in the literature concerning the proper range of Φ for air-exposed GaN. Hall mobility and XPS studies on (airexposed) MBE GaN films favor the higher μ and lower Φ ranges in Table I. For example, μ in the range 500–700 cm²·V⁻¹·s⁻¹ for GaN with $N \sim$ mid-10¹⁶ cm³ is consistent with the results in [102]. $\Phi \sim 0.5$ –0.3 eV is consistent with reports of XPS of clean and air-exposed GaN (0 0 0 1) surfaces [103]. However, Kelvin probe studies of air-exposed *c*-plane [104], [105] and *a*-plane [106] GaN surfaces consistently give $\Phi \sim 1$ eV and higher, so the matter is still open to debate.

We have discussed "fast" (\sim ns) surface recombination phenomena within the context of time-resolved PL. We also observe



Fig. 10. Fit of phenomenological PPC decay model to data is shown. The UV excitation is blocked at the point indicated, and the PPC decays in the dark. The fit recovers a free carrier concentration of $N = 4 \times 10^{17}$ cm⁻³ and capture cross section $\sigma = 8 \times 10^{-21}$ cm² for electrons, as explained in the text.

"slow" changes (over periods of seconds to hours) in surface band bending as revealed in the decay of persistent photoconductivity (PPC) transients. We presented a model describing this effect [101] that was based on a prior phenomenological theory of surface photovoltage on GaN films [104]. In these experiments, the current through a nanowire was monitored while UV light is turned on and off. The photoexcited holes were swept to the surface of the nanowire, reducing the surface band bending and depletion layer thickness. This change increased the diameter of the neutral region within the nanowire and hence the nanowire conductivity. When illumination ceased, equilibrium was restored via free electrons in the bulk of the nanowire surmounting the surface barrier and recombining with trapped holes. These recombination dynamics may be derived from an approximate Shockley-Read-Hall (SRH) analysis that assumes Fermi level pinning near midgap at the surface [107].

An example of fitting this SRH theory to PPC decay for a Batch 1 nanowire is illustrated in Fig. 10. The data are normalized from separate experiments and illustrate that the maximum relative photocurrent rises to 90% of the flatband value under continuous excitation of \sim 3.6 mW/cm² at 360 nm. The fitted PPC simulation is delayed until Φ has recovered sufficiently such that the cylindrical depletion approximation implicit to the model is valid [101]. The fitted solution returns $N = 4 \times$ 10^{17} cm^{-3} and a capture cross section $\sigma = 8 \times 10^{-21} \text{ cm}^2$ for electrons by the surface centers. The nanowire diameter between hexagonal vertices is 660 nm. The fitted magnitude of capture cross section is on the low end of the range generally reported for semiconductors (e.g., 10^{-22} to 10^{-12} cm²) [108]. On the other hand, it is consistent with the long PPC decay times observed and with "slow" surface recombination associated with surface adsorbates and oxide layers [105], [108]. Applying the same fitting procedure to Batch 2 samples discussed above returned N in the mid- 10^{16} cm⁻³ range with σ spanning 10^{-23} to 10^{-19} cm². These free carrier concentrations are consistent with results from the resistivity study summarized in Table I.

The spectral dependence of the PC is also an indicator of the quality of the MBE-grown nanowires because the subgap



Fig. 11. Weak subgap PC (in ambient room air, and in vacuum) for photoexcitation over the wavelength span indicated. The nanowire sample is chosen such that it is depleted in the dark in ambient air.

photoresponse is weak as compared to that for above-band-gap excitation. However, interpretation of these data should also consider ambient conditions. Fig. 11 illustrates the PC spectral dependence of a nanowire that is depleted in the dark while in room air. For excitation > 380 nm, the PC is at the dark (noise) level, which is roughly three orders of magnitude less than that under band-edge illumination. Under vacuum (mid- 10^{-3} Pa), the dark current increases and subgap PC becomes measurable. The excitation intensity over the spectral range indicated is within a factor of 2 of 0.2 mW·cm⁻². In contrast to the data in Fig. 11, PC studies on catalyst-grown GaN nanowires have identified specific subgap features [109].

B. GaN Nanowire FETs and High-Frequency Measurements

Although FET properties are not readily converted into measurements of carrier concentration and mobility, the performance of GaN nanowire FETs is promising for applications that require high sensitivity and on–off ratio. GaN FETs made with Schottky barrier gates exhibit subthreshold voltage swings approaching 60 mV/decade and on–off ratios $> 5 \times 10^8$ [10]. Fig. 12(a) shows an image of one such device, and Fig. 12(b) illustrates the pinchoff condition for increasing drain–source voltages. The low threshold voltage of -3.7 V indicates that this omega-gate architecture takes advantage of the large surface-tovolume ratio available in nanowires.

MBE-grown GaN nanowires fabricated with one ohmic and one Schottky contact have also been adapted for high-frequency measurements [110]. The broadband microwave reflection coefficients from about 1 to 10 GHz have been measured for one such structure, and the results fitted to estimate N. More sophisticated modeling and better understanding of interfacial effects are required to improve accuracy, but the demonstration shows that high-frequency capacitance measurements of N may be feasible.



Fig. 12. (a) Device structure and (b) drain–source current $(I_{\rm DS})$ versus gate– source voltage $(V_{\rm GS})$ for a single nanowire MESFET. The bottommost trace corresponds to a drain–source voltage $(V_{\rm DS})$ of 0.1 V, with $\Delta V_{\rm DS} = 0.1$ V between successive traces proceeding upward. Instrument artifacts produced the kink at $I_{\rm DS} = 5 \,\mu$ A. The inset is a log–linear plot of $I_{\rm DS}$ (solid) and a fit to the above-threshold linear region (dashed) versus $V_{\rm GS}$ for $V_{\rm DS} = 1$ V [10].

V. MECHANICAL PROPERTIES

One of the more interesting properties of MBE-grown GaN nanowires is that they display very high mechanical quality factors, Q, for an object with volume under 1 μ m³. The first evidence for high Q came from SEM observations of spontaneous resonances such as those illustrated in Fig. 13. These resonances can be enhanced with piezoelectric feedback to achieve Q > 1×10^6 in vacuum at room temperature [111]. These results compare quite favorably with catalyst-grown GaN nanowires, where Q < 3000 is typical [112]. The Q is found to be increasing as temperature increases, and is highly sensitive to absorption of small molecules or carbon deposits, indicating possible application as a mass sensor. Similar studies of nanowires coated with atomic layer deposition have shown that the frequency of oscillation can be tuned up or down depending on the stiffness of the coating. Individual nanowires generally recovered their original Q when the coatings were removed [113]. Measurements of GaN nanowire elastic properties with MEMS fixtures [114] and from modeling of the resonance [111] indicate that Young's modulus is equal to the modulus of bulk GaN within the rather large uncertainty. The high Q appears to be related instead to the smooth sidewalls and low chemical reactivity of GaN, so that these devices are free of damping from oxides that tend to reduce Q in silicon-based resonators.

Piezoresistive readout of GaN nanowire bridge structures has also been demonstrated in devices such as the one shown in Fig. 14. Because of the damping imposed by the metal contacts



Fig. 13 Two different mechanical resonances of the same GaN nanowire observed with SEM (insets) showing very high-Q under spontaneous excitation. Marker bars in insets are 100 nm; uncertainty in Q values is \pm 2000. Note that (a) one oscillation is along the wide diameter of the nanowire and (b) one is perpendicular to the sidewalls [115].



Fig. 14 Bridge structure with GaN nanowire suspended over gate metal to allow capacitive driving of nanowire oscillations and electrical readout of the oscillations [116].

on both ends, $Q \approx 1000-2500$ is typical for these structures. These devices can operate over a wide range of pressures, though damping by gas molecules also reduces Q.

VI. CONCLUSION

GaN nanowires grown by MBE currently represent some of the purest specimens of GaN and related alloys. This purity is reflected in their structural, optical, electrical, and mechanical properties. The smooth sidewalls and symmetric growth mode, especially under selective epitaxy processes, offer unique architecture for devices made from this technologically important material. Future study is expected to produce greater variety in heterostructures, parallel fabrication of large ensembles for device applications, and greater ability to measure key optical and electrical properties.

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