High Internal Quantum Efficiency Ultraviolet Emission from Phase-Transition Cubic GaN Integrated on Nanopatterned Si(100)

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Supporting Information

ABSTRACT: Ultraviolet emission characteristics of cubic (c-) GaN enabled through hexagonal-to-cubic phase transition are reported. Substrate patterning and material growth are shown to affect phase purity and emission characteristics of c-GaN as studied by electron backscatter diffraction, and photo- and cathodoluminescence, respectively. Raman study shows a tensile strain in the c-GaN. Time-resolved photoluminescence reveals c-GaN band edge emission decay time of 11 ps. The ultraviolet emissions from both phases of GaN are linearly polarized in the same direction, which is along the ⟨112̅0⟩ and ⟨110⟩ directions of hexagonal GaN and c-GaN, respectively. Temperature-dependent (5.7 to 280 K) cathodoluminescence studies reveal an internal quantum efficiency of ∼29% at room temperature along with intrinsic and extrinsic defect energy levels of ∼124 and ∼344 meV, respectively, of the phase-transition c-GaN. Using the IQE value and carrier decay lifetime, a radiative lifetime of 38 ps is extracted. Overall, photonic properties of phase-transition c-GaN and their dependence on substrate patterning and material growth are reported.

KEYWORDS: cubic phase, gallium nitride, cathodoluminescence, electron backscatter diffraction, photoluminescence, ultraviolet emitter

Gallium nitride (GaN) and its alloys are the semiconductors of choice for driving the solid-state lighting revolution through the white light emitting diodes (LEDs). Such LEDs are hexagonal (wurtzite) phase semiconductors and conventionally grown on the polar (0001) plane of the thermodynamically stable hexagonal (h-) phase of GaN. However, the lack of inversion symmetry in h-crystal structure leads to strong spontaneous and piezoelectric polarization fields (∼MV/cm) along the ⟨0001⟩ growth direction. Such large electric fields have many detrimental effects in photonics (reducing the radiative recombination rates in LEDs due to the quantum confined Stark effect) and electronics (dictating a normally on operation in high electron mobility transistors, HEMTs).

As such, unconventional crystal facets such as the nonpolar ⟨1100⟩ m-plane of h-GaN with zero polarization fields are being explored. However, these substrates suffer from low availability, low scalability, and high defectivity. This necessitates the search for novel GaN photonic materials with zero polarization nature and yet on an inexpensive, scalable platform.

GaN also forms in the cubic (zincblende) phase. The cubic phase of GaN (c-GaN) is inherently polarization-free in the ⟨100⟩ growth direction due to its centrosymmetric atomic arrangement. Additionally, c-GaN has benefits such as high carrier mobility, high p-type conductivity, high electron drift velocity, small Auger losses, high optical gain, high radiative efficiency, and cleavage planes.

Most notably, the band gap of c-GaN (E_g = 3.22 eV) is 0.2 eV smaller than that of h-GaN (E_g = 3.42 eV). The conventional high-indium-content h-InGaN green LED is 4 times less efficient than the blue and red LEDs; this creates the
To date, the Article a,c,e, and their representative cross-sectional Here we employ this approach to enable 3C-SiC 10.1021/acsphotonics.7b01231 Thus, our approach As shown in Figure 1, the U-groove consists of two vertical dielectric sidewalls conffned by h-GaN (10T1) planes (red). In (e) and (f), the U-groove is overdeposited (h > h_c), resulting in a defective, mixed phase material (purple).

Figure 1. Phase-transition GaN drawings (a, c, e) and SEMs (b, d, f) under various conditions. In (a) and (b), the U-groove is optimized (h = h_c), enabling complete c-GaN (blue) surface coverage. In (c) and (d), the U-groove is underdeposited (h < h_c), leading to a dual-phase surface of c-GaN (blue) flanked by h-GaN (10T1) planes (red). In (e) and (f), the U-groove is overdeposited (h > h_c), resulting in a defective, mixed phase material (purple).

The most important issue hampering c-GaN is its metastability when grown directly on conventional cubic substrates (MgO,27 GaAs,18 3C-SiC17), limiting its practicality. GaN grown via such an approach results in highly defective (>10^10 cm^-2) and mixed h-/c-phase materials, as GaN thermodynamically prefers the h-GaN crystal formation.19

An alternative approach, hexagonal-to-cubic phase transition, has recently been introduced to synthesize single-crystalline stable c-GaN.21−23 Here we employ this approach to enable high phase purity and high crystal quality c-GaN via metalorganic chemical vapor deposition (MOCVD) on a CMOS-compatible on-axis nanopatterned Si(100) substrate.24 By meeting critical substrate patterning and growth conditions, we circumvent the need for any postgrowth processing techniques (e.g., ion beam milling) to enable complete phase-transition c-GaN surface coverage.25 Thus, our approach enables us to study and report on the structural and optical properties of the as-grown phase-transition c-GaN and demonstrate the effect of growth and patterning on the crystal quality and luminescence characteristics. In addition, our employment of Si(100) as the substrate reduces cost, increases scalability, and opens up heterointegration opportunities.

The hexagonal-to-cubic phase transition of GaN is facilitated on a U-shape nanogrooved Si(100) substrate.24 As shown in Figure 1, the U-groove consists of two vertical dielectric sidewalls confning two opposing Si(111) planes and a Si(100) plane in the center. We recently showed that U-groove parameters, which are dielectric opening width (p), silicon etch depth (t_s), and dielectric sidewall angle (α), affect the phase-transition c-GaN surface coverage atop the U-grooves. Complete c-GaN surface coverage occurs when the deposition thickness (h) of GaN deposited above the Si(100) substrate surface equals the critical thickness (h_c):25

\[
h_c = \frac{1.06p - 0.75t_s}{1 - \tan \alpha/0.71}
\]

RESULTS

Three different c-GaN surface coverage outcomes (complete, underdeposition, overdeposition) are schematically presented in Figure 1a,c,e, and their representative cross-sectional scanning electron microscopy (SEM) images are shown in Figure 1b,d,f. In the U-grooves, h-GaN (colored red) nucleates on the (111) planes of the silicon substrate, making an angle of 54.7° to the Si(100) plane. The two opposing h-GaN wings then merge at twice the angle (54.7° × 2 = 109.4°) at a seam point where phase transition from h-GaN (colored red) to c-GaN (colored blue) initiates. Thus, the c-GaN (100) plane is parallel to that of the Si(100) substrate. A void forms below that seam point. Optimized growth and patterning conditions (h = h_c) are presented in Figure 1a,b.

The two nonideal deposition scenarios are under- and overdeposition. In the former case (Figure 1c,d) the phase-transition c-GaN does not grow over the h-GaN wings. Hence, the (10T1) planes of the h-GaN wings, making an angle of ~7° to the Si substrate, are observed. Achieving complete c-GaN coverage requires further GaN deposition, much needed to increase h to the level of h_c. In the latter case (Figure 1e,f), overgrowth above the c-GaN rhombus becomes mixed phase, which is due to h-GaN growth from the sidewalls. In order to quantify between under-, ideal-, and overdeposition cases, we define the h/h_c ratio as the fill factor (ff) where an ff less than 100% suggests underdeposition, whereas an ff more than 100% suggests overdeposition.
Electron backscatter diffraction (EBSD) studies are conducted to investigate the phase transition of GaN. The Kikuchi bands formed by a 15 kV electron beam is used to distinguish the crystal phases, orientation, and grain (phase) boundaries of the GaN in U-grooves with a spatial resolution of \( \sim 50 \) nm. To best study the phase transition, EBSD was conducted on the undergrown sample (Figure 1c,d) with 20 nm \( \times \) 20 nm pixels.

The 70° tilted view SEM image of the undergrown U-grooves is shown in Figure 2a. The c-GaN “plateau” can be seen sandwiched between two h-GaN “wings”, which are neighbored by the “SiO\textsubscript{2}” dielectric sidewalls. The EBSD phase detection of the region marked by the red rectangle in Figure 2a is shown in Figure 2b, with red, blue, and black pixels representing hexagonal, cubic, and indistinguishable crystal phases, respectively. The inverse pole figures (IPF) of the [100], [010], and [001] directions of the same region are plotted in Figure 2e,f,g. The color indices for IPFs of c-GaN and h-GaN are provided in Figure 2c,d. It is seen that the plateau region of the U-groove is composed of solely c-GaN with no visible grain boundary and crystal misalignment. Similar to the Si(100) substrate, the c-GaN plateau has its (100) plane pointing in the normal direction and the (110) planes along and perpendicular to the grooves. These results indicate the phase transition of GaN in the U-grooves is uniform and controllable and leads to a single-crystalline c-GaN surface. These results demonstrate an opportunity for studying the optical properties of phase-transition c-GaN.

A transmission electron microscopy (TEM) study is conducted to characterize the crystal quality of the GaN materials in a U-groove, as shown in Figure 3: (a) bright field and (b) dark field images. A highly defective and thin (\( \sim 15 \) nm) layer of AlN is seen covering the Si substrate and the amorphous oxide barriers. Near the h-GaN nucleation sites on the Si(111) planes, both TEM images show crystals containing dislocations and stacking faults. The h-GaN crystals that are more than 50 nm away from the nucleation sites show increased quality, and dislocations are rarely observed. A \( \sim 20 \) nm long seam oriented perpendicular to the substrate is observed in the middle of the groove between the two opposing h-GaN wings directly above the void. It terminates into two distinctly straight interfaces extending toward the oxide sidewalls to form a V-shaped boundary. Enclosed in the V-boundary is the c-GaN cubic crystal, which is further confirmed by TEM electron diffraction taken from the triangular area shown in Figure 1a. Stacking faults parallel to...
the interface, i.e., lying on the c-GaN (111) and h-GaN (0001) planes, are observed to accommodate the transition between the two crystal structures, but are limited to within a few layers of atoms thick (not shown). At the c-GaN/oxide interface near the surface, defects (stacking faults) are observed to orient away from the oxide (Figure 3b). Defective and mixed-phase material are observed above these stacking faults and the aforementioned critical height, \( h_c \). This demonstrates the importance of silicon substrate patterning and GaN growth optimization, as inadvertent relaxation of c-GaN to mixed-phase material can occur at the c-GaN/barrier.

Raman spectroscopy is used to study the crystallinity and strain of GaN materials grown in various-sized U-grooves. Using a Horiba Raman confocal imaging microscope with a depolarized 405 nm laser and 1800 lines/mm grating, Raman spectra of U-grooves with \( ff' \)’s ranging from 0% to 205% are collected and plotted in Figure 4 to demonstrate the effect of optimization on the crystal quality.

![Figure 4](image)

**Figure 4.** Confocal Raman spectroscopy of U-grooves with \( ff' \)’s from 0% (purely h-GaN) to 210% (overgrown-mixed phase GaN). In (a), four peaks are labeled: \( A_2(LO) \) and \( E_{22}(TO) \) of h-GaN, Si, and \( A_1(LO) \) of the AlN nucleation layer. In (b), as phase transition occurs, c-GaN’s TO-phonons can be observed. At \( ff' = 205\% \), the extremely overdeposited case, c-GaN’s TO-phonon disappears, and the fwhm of h-GaN’s signal increases, indicating a reduction in crystallinity.

Four peaks in Figure 4a are identified as Si substrate (520 cm\(^{-1}\)), \( E_{22}(TO) \) and \( A_2(LO) \) of h-GaN (567 cm\(^{-1}\), 733 cm\(^{-1}\)), and \( E_{21} \) or \( A_1(LO) \) of AlN (618 cm\(^{-1}\)). Zoomed-in Raman spectra of the TO-phonons are shown in Figure 4b, in which the 558 cm\(^{-1}\) peak, attributed to the \( E_{22}(TO) \) shift of h-GaN, is observed. Another peak at 551 cm\(^{-1}\), attributed to TO-phonons of c-GaN, emerges as \( ff' \) increases from 0% to 95%, signifying the increasing presence of c-GaN.\(^{26}\) The peak at 733 cm\(^{-1}\) intensities (Figure 4a), suggesting an involvement of the LO-phonons of c-GaN in addition to the \( E_1(LO) \) phonons of h-GaN. Both peaks of c-GaN show a red-shift in their Stokes shift compared to the literature (555 and 742 cm\(^{-1}\) for TO- and LO-phonons, respectively), indicating c-GaN is under a tensile strain,\(^{26}\) which is in agreement with other studies on phase-transition c-GaN on Si(100).\(^{22,23}\) This is due to the lattice mismatch between Si(111) and h-GaN being passed to the c-GaN. At the extreme case of overdeposition (\( ff' = 205\% \)), all GaN-related Raman signals decrease in intensity and increase in their full-width at half maximum (fwhm), and the TO-phonon Raman mode of c-GaN becomes undetectable. We attribute this to the mixed-phase material on the surface in overgrowth samples.\(^{27}\)

We perform time-integrated and time-resolved photoluminescence (PL) studies to investigate the luminescence lifetimes in h- and c-GaN. The luminescence lifetime can reveal the carrier loss mechanisms and quantitatively compare their rate and how suitable the material is for photonic applications. The measured PL lifetime (\( \tau_{\text{PL}} \)) is related to the radiative lifetime (\( \tau_{\text{rad}} \)) and nonradiative lifetimes (\( \tau_{\text{nr}} \)) through

\[
\tau_{\text{PL}}^{-1} = \tau_{\text{rad}}^{-1} + \tau_{\text{nr}}^{-1}
\]

It is known that the radiative lifetime in semiconductors is determined by the scattering from the excitonic branch to the photonic branch of the exciton–polariton dispersion curves. Hence, the radiative lifetime for a 200 nm thick GaN film can be expressed as:\(^{28}\)

\[
\tau_{\text{rad}} = \frac{1}{2} \left( \frac{5\pi}{3} \right)^{3/5} \left( \frac{M}{\nu_0} \right)^{2/5} \left( \frac{\hbar c^2}{\alpha_p^5} \right)^{1/5} \left( \frac{C^*\alpha^5}{C^*\alpha^5} \right)^{1/5}
\]

where \( M/\nu_0 \) is the mass per unit volume of the unit cell (\( \sim 3 \) kg m\(^{-3}\)), \( \alpha_p \) is the polarizability (\( 4 \times 10^{-5} \)), \( L \) is the thickness of the excited layer (\( \sim 200 \) nm), \( \hbar \omega \) is the photon energy of the excitonic transition (3.42 eV), and \( C \) is the deformation potential (4 eV). This equation yields a theoretical radiative lifetime of \( \sim 470 \) ps for h-GaN. The corresponding radiative lifetime of c-GaN is estimated to be \( \sim 47 \) ps, an order of magnitude shorter than that of h-GaN, as suggested by the literature.\(^{13,29}\) This is due to the greater electron–hole wave function overlap, which results in a greater oscillator strength, in the absence of polarization fields in c-GaN.\(^{13}\)

Experimentally, a time-resolved PL study is carried out with a frequency-tripled Ti:Al\(_2\)O\(_3\) laser (\( \lambda = 266 \) nm) and a streak camera system with a temporal resolution of \( \sim 1 \) ps to investigate the carrier dynamics of phase-transition c-GaN. Figure 5 shows the PL spectrum with two peaks at 3.42 and 3.22 eV, which correspond to the free exciton transitions of h-GaN and c-GaN, respectively.\(^{26}\) A time-resolved PL study of the selected spectral ranges, shown in Figure 5 as insets, shows exponential decays with lifetimes of 20 ps for 3.42 eV (h-GaN) and 11 ps for 3.22 eV (c-GaN).

Studies have shown that lateral epitaxial overgrowth h-GaN on sapphire exhibits a PL lifetime in the range of 80–210 ps at room temperature\(^{31,32}\) and 530 ps for GaN grown by hydride vapor phase epitaxy (HVPE).\(^{33}\) For c-GaN, studies reported an extremely short PL lifetime of \( \sim 15 \) ps.\(^{34}\) For h-GaN, our data show an extremely fast decay compared to the high-quality material in the literature, suggesting an additional carrier loss mechanism in the U-grooves. However, the phase-transition c-GaN shows a comparable PL decay lifetime to the 1 \( \mu \)m thick c-
Ga\textsubscript{N} films on 3C-SiC, indicating a similar carrier recombination rate.\textsuperscript{24} Using eq 2 and a theoretical \(r_{\text{rad}}\) of 470 ps (47 ps) for h-GaN (c-GaN), the nonradiative decay dominates the transition with a \(r_{\text{rad}}\) of 21 ps (14 ps) for the h-GaN (c-GaN) in the U-grooves. We believe that the short radiative lifetimes of the two phases of GaN in the U-grooves are dominated by two mechanisms: material interfaces and strong oscillator strength. The stacking faults along the hexagonal-cubic phase-transition interfaces may act as nonradiative recombination centers.\textsuperscript{35} The h-GaN–silicon and c-GaN–air interfaces may also provide another carrier loss mechanisms. On the other hand, a fast PL lifetime could also be the result of a strong radiative recombination resulting from the higher degree of confinement provided by the quasi 1D nature of the U-grooves. It has been reported that c-GaN has a significantly stronger oscillator strength with its smaller band gap and the lack of polarization fields.\textsuperscript{36} We are conducting further studies to better understand the origins of ultrafast lifetimes in these materials.

Polarization-dependent PL can reveal the nature of the radiative recombination and the escape path of the photons from the material. The phase-transition c-GaN in the U-grooves exhibits polarization in the light emitted through either a grating-like effect\textsuperscript{36} and/or strain-induced valence band splitting.\textsuperscript{37} In this work, the U-grooves are orientated parallel to the passing axis of the polarizer at 0°, and the normalized result is plotted in Figure 6. The band edge emissions from h-GaN and c-GaN are shown by red squares and blue circles, respectively. The emissions from both phases of GaN are linearly polarized in the same direction, which is along the (1120) and (110) directions of h-GaN and c-GaN/Si, respectively. The magnitude of the polarization of the emitted photons is characterized by the degree of polarization \(\rho\), which is

\[
\rho = \frac{I_\parallel - I_\perp}{I_\parallel + I_\perp}
\]

where \(I_\parallel\) and \(I_\perp\) are the intensity when the passing axis of the polarizer is parallel and perpendicular to the U-grooves, respectively. The values are found to be 0.12 for h-GaN and 0.20 for c-GaN. The polarization of h-GaN emission (3.42 eV) in a grating-like setting has been reported to have a strong linear polarization parallel to the direction of the grating,\textsuperscript{38} which is in good agreement with the results shown in Figure 6. The polarization observed in this work is believed to be the result of a combination of the grating effect and tensile strain, which is determined through Raman studies.

Cathodoluminescence (CL) is used to study phase purity, defect distribution, impurity levels, and internal quantum efficiencies of various GaN materials in U-grooves. Particularly, CL can map the luminescence with a spatial resolution of <1 \(\mu\)m, providing useful insights on the spatial distribution of the radiative recombination centers. Our study was conducted via a JEOL JSM 7000F field emission scanning electron microscope outfitted with a Gatan MonoCL3 for panchromatic/monochromatic imaging and spectroscopy with a wavelength resolution of 2 nm fwhm and a helium-flow cryo-stage capable of temperature control from 5.65 K to >300 K. The electron beam is set to emit a beam current of 0.1–3.2 nA at 2 kV, which yields a maximum electron penetration depth of \(\sim47\) nm.\textsuperscript{39} This allows for a good combination of maximizing signal intensity while minimizing the amount of excitation of the h-GaN wings buried underneath the c-GaN isosceles triangle (Figure 1a).

As shown by Raman spectroscopy and EBSD, the \(ff = 95\%\) is the most optimized U-groove sample for c-GaN and provides the largest surface coverage of phase-transition c-GaN with the highest crystal quality. As such, a CL study conducted on this U-groove configuration reveals the optical properties of the highest quality c-GaN enabled via phase transition. Figure 7 shows the CL spectrum of the \(ff = 95\%\) U-grooves conducted at 5.7 K with variable electron beam current in raster scan mode, which gives a good representation of the luminescence properties of the top 47 nm layer over a 20 \(\times\) 30 \(\mu\)m\textsuperscript{2} area.

Three luminescence centers are observed and identified as bound exciton (BX) and donor–acceptor pairs (DAP\textsubscript{1} and DAP\textsubscript{2}) at 3.28, 3.17, and 2.95 eV, respectively. LO-phonons of the DAPs are also observed at energy intervals of \(\sim90\) meV from the zero-phonon line, with DAP\textsubscript{2} showing a stronger phonon coupling due to the deeper acceptors involved, as expected. No luminescence at 3.48 eV, which is the excitonic

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**Figure 5.** Room-temperature PL spectra of the U-grooves. Near band edge emissions of h-GaN at 3.42 eV (red) and c-GaN at 3.22 eV (blue) are observed. The time-resolved PL luminescence decay of the marked spectral ranges are shown as insets.

**Figure 6.** Polarization-dependent photoluminescence of the near band edge emission of h-GaN (red squares) and c-GaN (blue circles). The passing axis of the polarizer angle is parallel to the direction of the U-grooves. The minimum occurs when the polaroid is 90° to the direction of grooves for both materials, with c-GaN showing a higher degree of polarization than h-GaN.
recombination of h-GaN, is observed. A broad emission centered at 3.67 eV with an fwhm of 220 meV is reported. Injection current dependent CL is used to determine the nature of this radiative recombination center. A power-law study reveals a linear relationship between its intensity ($I_{CL}$) and injection current ($J_b$) with a slope of 0.9 ± 0.2, which indicates an unsaturated, band-to-band-like transition across the injection current range of 0.1 to 3.2 nA ($I_{CL}^{3.67eV} \propto J_b^{0.9}$). Since the emission energy is above both cubic and hexagonal phase GaN, it is likely from neither crystal phases. We attribute this to the presence of the exposed AlN nucleation layer on the SiO$_2$ sidewalls, considering the persistent AlN phonon shifts observed in Raman spectroscopy (Figure 3) across different $ff$’s.

Cathodoluminescence imaging was conducted to investigate the spatial distribution of the luminescence centers at 5.7 K. High-magnification SEM images of grooves with various $ff$’s from 30% to 205% are shown in Figure 8a, and a large-scale SEM image is shown in Figure 8b. The monochromatic CL images of the grooves shown in Figure 8b at $\lambda = 355$ nm (3.49 eV, h-GaN BX), 390 nm (3.18 eV, c-GaN DAP$_1$), and 420 nm (2.95 eV, c-GaN DAP$_2$) are shown in Figure 8c,d,e, respectively.

The luminescence of h-GaN BX (Figure 8c) in the undergrown U-grooves ($ff = 30\%$ and 40%) shows very uniform and intense luminescence, which indicates the presence of h-GaN on the surface, while the intermediate U-grooves ($ff = 65\%, 70\%, 90\%$) show a decreasing amount of luminescence at this energy, indicating the decreasing amount of h-GaN that is excited within the top ∼47 nm layer. The most optimized U-grooves, $ff = 95\%$, show no observable h-GaN emission, indicating a very good coverage of the c-GaN triangle on the surface. Beyond the optimized condition for phase transition at $ff = 100\%$, traces of h-GaN BX luminescence can be observed scattered randomly across the image, indicating a reappearance of h-GaN in an irregular distribution. The luminescence of DAP$_1$ (Figure 8d) of c-GaN at 3.18 eV can be seen scattered randomly across the images in small and

Figure 8. Plain view SEM and CL images of GaN on various U-grooves. (a) High-magnification SEM images with a scale bar of 500 nm. (b) Low-magnification SEM image with scale bars of 5 μm, and its corresponding monochromatic CL images at (c) the band gap of h-GaN at 355 nm, (d) DAP$_1$ of c-GaN at 390 nm, and (e) DAP$_2$ of c-GaN at 425 nm.
bright clusters. The intensity and concentration of these clusters seem to have a weak dependence on \( f_f \). At the extreme end of overgrowth \( (f_f = 20\%) \), the concentration of DAP\( _1 \) increases dramatically, but the cluster-like distribution persists. This suggests an increase in point defect formation (e.g., lattice site vacancies, \( V_{\text{vac}} V_{\text{Ga}} \)) or slip planes (e.g., basal stacking faults) rather than an extrinsic impurity (C, O, etc.) since the growth condition has not changed between the U-grooves. The formation of stacking faults is directly observed in Figure 3b near the c-GaN/oxide interface. The luminescence of c-DAP\( _2 \) (Figure 8c), while showing a similar cluster-like distribution observed in c-DAP\( _1 \), follows the trend of the cubic optimization. It is at its strongest when \( f_f \) is the closest to 100\% and becomes weaker as \( f_f \) deviates from 100\%. This suggests this luminescence center is unique to crystalline c-GaN and is not observed when a large degree of phase mixing or poor crystalline quality is present.

Temperature-dependent CL is conducted on c-GaN to calculate the energies of the donors and acceptors involved in DAP\( _1 \) and DAP\( _2 \). Figure 9 shows the Arrhenius plot of the integrated CL intensity of the three luminescence centers of c-GaN. The integrated intensity obeys

\[
I(T) = \frac{I_0}{1 + C \exp(-E_{\text{act}}/k_B T)}
\]

where \( I_0 \) and \( C \) are fitting constants, \( k_B \) is Boltzmann’s constant, and \( T \) is the temperature. Fitting of the experimental data reveals a donor activation energy \( (E_{\text{act}}) \) of 20, 16, and 16 meV for BX (exciton binding energy), DAP\( _1 \) (shallow donor), and DAP\( _2 \) (shallow donor), respectively. The proposed band diagram for each radiative recombination is shown as the inset.

Figure 9. Arrhenius plot of c-GaN CL emissions at 5.7 K. Fitting shows an activation energy of 20 ± 9, 16 ± 7, and 16 ± 4 meV for BX (exciton binding energy), DAP\( _1 \) (shallow donor), and DAP\( _2 \) (shallow donor), respectively. The proposed band diagram for each radiative recombination is shown as the inset.

Temperature dependence is also shown. The IQE for c-GaN increases very rapidly as the temperature decreases. IQE and CL imaging suggest the importance of U-grooves (the U-grooves shown in the bottom row of Figure 3) near the c-GaN/oxide interface. The luminescence of c-DAP\( _1 \) is at its strongest when \( f_f \) is the closest to 100\% (the U-grooves shown in the bottom row of Figure 3), while showing a similar cluster-like distribution observed in c-DAP\( _2 \), follows the trend of the cubic optimization. It is at its strongest when \( f_f \) is the closest to 100\% and becomes weaker as \( f_f \) deviates from 100\%. This suggests this luminescence center is unique to crystalline c-GaN and is not observed when a large degree of phase mixing or poor crystalline quality is present.

The IQE value of 29\% indicates phase transition and nanopatterning are capable of synthesizing crystals with excellent quality without the need for thick and intricate buffer layers. This is especially clear when compared to commercially available 3-μm-thick h-GaN on sapphire, 325-μm-thick HVPE-grown freestanding GaN, and GaN on Si(111), which exhibit IQE values of 12\%, 8\%, and 2\%, respectively (see Supporting Information Figures S1, S2, S3).

The inset in Figure 10 shows the IQE of c-GaN for various \( f_f \) from 0\% (one h-GaN wing without any phase transition) to 210\% (the U-grooves shown in the bottom row of Figure 7). The IQE for c-GaN increases very rapidly as \( f_f \) approaches 100\% and drops down to very small values once \( f_f \) exceeds unity. IQE and CL imaging suggest the importance of U-pattern optimization and regrowth height. If GaN is under-deposited, c-GaN is unable to cover the entire surface, resulting in poor quantum efficiency. If GaN is overdeposited, defect formation dominates the material growth, resulting in poor quantum efficiency. There exists a relatively small window in which IQE is maximized for c-GaN crystals given a specific U-
groove. This is when complete c-GaN coverage is achieved through eq 1.

Another approach to estimate $\tau_{\text{rad}}$ of c-GaN is to utilize the IQE calculated from eq 6. Using IQE = 29% and the measured $\tau_{\text{rad}} = 11$ ps of c-GaN, the $\tau_{\text{rad}}$ can be extracted using eq 7. This yields a $\tau_{\text{rad}}$ of 38 ps, which is ~12.4 times shorter than the value for h-GaN, and a $\tau_{\text{rad}}$ of 22 ps. The finding in the TRPL for c-GaN is in good agreement with the other studies in the literature. This method, which does not necessarily assume the relationship between the radiative lifetime of c- and h-GaN, independently verifies the one-order faster radiative recombination lifetime in c-GaN than that of h-GaN observation.

$$\eta_{\text{IQE}} = \frac{\tau_{\text{rad}}}{\tau_{\text{nu}}}$$

Fundamental material properties can be extracted using the observed radiative lifetime (38 ps) and eq 3. Assuming similar mass density and deformation potential between h-GaN and c-GaN, the polarizability, $\alpha$, can be calculated for c-GaN using the 3.22 eV for $\alpha$ and 200 nm for $L$. The polarizability of c-GaN is calculated to be $8.3 \times 10^{-11}$, which is approximately 7 orders of magnitude smaller than that of h-GaN.

### CONCLUSION

We demonstrate that phase transition of h-GaN to c-GaN is a reliable c-GaN growth technology that can be precisely controlled by substrate patterning and material growth. A high-crystallinity and -uniformity phase-transition c-GaN surface can be reproduced when the critical condition, $h_c = (1.06p - 0.75\alpha)/(1 - \tan \alpha/0.71)$, is met. Phase-transition c-GaN is observed to be under tensile strain, and its light emission has 0.2 degree of polarization along the pattern direction. Low-temperature CL spectroscopy identified two defect luminescence centers at 144 and 344 eV above the valence band, and imaging of the distribution of these acceptors suggests these are intrinsic defect and extrinsic impurity levels, respectively. The band-to-band emission of phase-transition c-GaN internal quantum efficiency is calculated independently via temperature-dependent CL and time-resolved photoluminescence to be 29%, making this technology suitable for next-generation photonic devices.

### ASSOCIATED CONTENT

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsphto-0000-0001-9696-8830.

Temperature-dependent photoluminescence spectra of conventional 3-μm-thick h-GaN on sapphire; temperature-dependent photoluminescence spectra of HVPE-grown 325-μm-thick freestanding h-GaN; temperature-dependent photoluminescence spectra of h-GaN on Si(111) (PDF)

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### REFERENCES


**Notes**

The authors declare no competing financial interest.


Supporting Information

High internal quantum efficiency ultraviolet emission from phase-transition cubic GaN integrated on nanopatterned Si(100)

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(2 pages, 3 figures)

To compare with the literature, we conducted IQE measurements of conventional samples (purchased commercially) of GaN on Sapphire, free-standing GaN, and GaN/Si wafers. Figures S1-3 show the photoluminescence spectra of 3 µm-thick GaN/sapphire, 325 µm-thick HVPE-grown freestanding GaN, and 500 nm-thick GaN/Si(111) at 1.4 and 300 K, respectively. The IQE values of GaN on Sapphire, free-standing GaN, and GaN/Si wafers. Band-edge emissions are measured to be ~12%, ~8%, and ~2% whereas our novel work with cubic GaN led to an IQE of ~29% - showing the promise of this technology.

Figure 1 PL spectra of 3µm-thick GaN/Sapphire at 1.4 and 300 K, showing an IQE of ~12%.
Figure 2 PL Spectra of 325 µm-thick HVPE-grown GaN, showing an IQE of ~8%.

Figure 3 PL spectra of 500 nm-thick GaN/Si, showing an IQE of ~2%.