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A Simple Approach to Achieving Ultrasmall III-Nitride Microlight-Emitting Diodes with Red Emission

Peng Feng,⁺ Ce Xu,⁺ Jie Bai,⁺ Chenqi Zhu, Ian Farrer, Guillem Martinez de Arriba, and Tao Wang*



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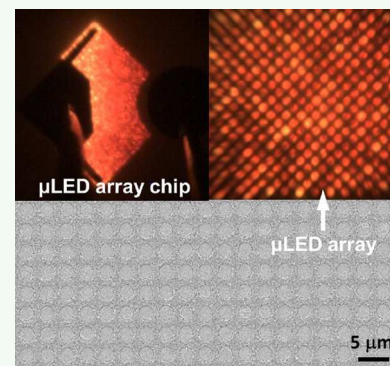
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ABSTRACT: The microdisplays for augmented reality and visual reality require ultrasmall microlight-emitting diodes (μ LEDs) with a dimension of $\leq 5 \mu\text{m}$. Furthermore, the microdisplays also need three kinds of such μ LEDs each emitting red, green, and blue emission. Currently, in addition to a great challenge for achieving ultrasmall μ LEDs mainly based on III-nitride semiconductors, another fundamental barrier is due to extreme difficulty in growing III-nitride-based red LEDs. So far, there has not been any effective approach to obtain high indium content InGaN as an active region required for a red LED while maintaining high optical performance. In this paper, we have demonstrated a selective epitaxy growth approach on a template featuring microhole arrays. This allows us to not only obtain the natural formation of ultrasmall μ LEDs but also achieve InGaN with enhanced indium content at an elevated growth temperature, at which it is impossible to obtain InGaN-based red LEDs on a standard planar surface. By means of this approach, we have demonstrated red μ LEDs (at an emission wavelength of 642 nm) with a dimension of $2 \mu\text{m}$, exhibiting a high luminance of $3.5 \times 10^7 \text{ cd/m}^2$ and a peak external quantum efficiency of 1.75% measured in a wafer form (i.e., without any packaging to enhance an extraction efficiency). In contrast, an LED grown under identical growth conditions but on a standard planar surface shows green emission at 538 nm. This highlights that our approach provides a simple solution that can address the two major challenges mentioned above.



KEYWORDS: InGaN, microLED, selective epitaxy growth, patterned template, MOVPE, EQE

1. INTRODUCTION

There is a growing interest for developing microdisplays with compact screens of $\leq 1/4$ " diagonal length, which have a wide range of applications in smart watches, smart phones, smart bands, and augmented reality and virtual reality (AR & VR) devices.^{1–5} Their individual pixel elements typically consist of a large number of microscale visible LEDs mainly based on III-nitride semiconductors, which are referred to as microLEDs (μ LEDs). For instance, the microdisplays for AR and VR require μ LEDs with an ultrasmall dimension of $\leq 5 \mu\text{m}$.^{6–8} Such devices are typically utilized in a scenario where spaces are small or the devices need to be close to the eyes. Therefore, the devices require high resolution, high contrast ratio, high luminance, and high external quantum efficiency (EQE).^{9,10} Of course, a microdisplay needs three kinds of individual μ LEDs as a single pixel each emitting red, green, and blue emission (i.e., RGB), respectively.

InGaN semiconductors have direct bandgaps across their whole content ranging from 0.7 eV for InN to 3.43 eV for GaN, covering part of the infrared region, the full visible spectrum, and part of the ultraviolet (UV) region. So far, InGaN-based μ LEDs with reasonably good performance in the blue and green spectral region have been reported. However, red LEDs still rely on AlGaInP materials. Although a large area

AlGaInP red LED with a high efficiency of $>50\%$ can be obtained,¹¹ the efficiency reduces dramatically when its dimension is reduced to the microscale, namely, μ LEDs. This is due to an enhancement in the surface recombination rate and the long diffusion lengths of carriers.^{12–15} Moreover, the efficiency of AlGaInP red LEDs is sensitive to their junction temperature,^{16,17} and thus, AlGaInP red LEDs generally suffer from a severe leakage current at a high temperature, generating a severe efficiency thermal drop. All these fundamental issues indicate that it is indispensable to develop III-nitride-based red LEDs to meet the requirements for the fabrication of a full color microdisplay.

InGaN with high indium content ($>20\%$) is necessary for obtaining long wavelength emission. Unfortunately, it is greatly challenging to achieve high indium content InGaN while maintaining high optical performance.^{18,19} A typical method to achieve high indium content in InGaN is to lower the growth

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62 temperature for InGaN. However, it is clear that this method is
63 not ideal because it causes a significant degradation in crystal
64 quality.

65 In general, vapor–solid thermodynamic equilibrium can be
66 modified by stress, making the solid-phase epitaxial composi-
67 tion reduce toward lattice-matched conditions. This is the
68 major reason why it is difficult to increase indium
69 incorporation into GaN.^{20–24} Therefore, the growth of
70 InGaN on a relaxed layer is beneficial for obtaining high
71 indium content in InGaN. However, bear in mind that the
72 formation of a relaxed layer is often associated with the
73 generation of extra defects if a heterostructure with a large
74 lattice mismatch is used to generate a relaxed layer. This leads
75 to degradation in optical performance. Furthermore, the stress
76 status of an underlying layer plays a critical role in determining
77 indium incorporation into GaN. Generally speaking, tensile
78 stress tends to enhance indium incorporation into GaN,
79 offering a unique advantage for growing red LEDs on silicon
80 substrates as GaN on Si suffering tensile stress.^{25,26} In contrast,
81 GaN grown on sapphire substrates exhibits compressive strain.

82 The growth of InGaN-based red LEDs has been reported by
83 means of inserting a thin AlN or an AlGaIn layer into each
84 InGaIn quantum well as an emitting region, leading to an
85 enhancement in strain that pushes the emission wavelength of
86 InGaIn quantum wells toward longer wavelength.^{27,28} So far,
87 this approach has become a popular method for the growth of
88 long wavelength emitters, in particular, red LEDs.^{25–30}
89 Furthermore, by combining the idea of inserting a thin AlN
90 or AlGaIn and further adjusting the in-plane strain of a GaN
91 template by tuning the GaN thickness, 633 nm-wavelength red
92 LEDs with an external quantum efficiency (EQE) of 1.6% has
93 been reported, where an extremely thick GaN (8–10 μm)
94 template has been employed.^{26,29} A more recent work has
95 demonstrated that a peak EQE as high as 4.5% has been
96 achieved on red InGaIn μLEDs .³¹ However, it is worth noting
97 that the approach of strain enhancing also leads to a reduction
98 in internal quantum efficiency.

99 We expect that an enhanced relaxation can be achieved by
100 using a selective epitaxy growth approach on a microhole-
101 patterned template, which we have developed recently, where
102 μLEDs can be naturally formed but without employing any
103 dry-etching techniques because selective epitaxy growth can
104 take place only within these microholes.^{7,8} In this work, we are
105 proposing to employ this approach to achieve ultrasmall red
106 μLED arrays with enhanced quantum efficiency but without
107 inserting any thin AlN or AlGaIn into InGaIn quantum wells as
108 an emitting region. It is expected that no lateral confinement
109 during the selective epitaxy growth process leads to strain
110 relaxation effectively and naturally. By this mechanism, 642 nm
111 red μLEDs with a dimension of 2 μm have been achieved by
112 our selective epitaxy growth conducted at an elevated
113 temperature, at which a red LED cannot be achieved on a
114 standard planar GaN surface. The resultant external quantum
115 efficiency is 1.75%. For comparison, only 538 nm green LEDs
116 on a standard planar GaN template can be obtained even
117 under identical growth conditions. Our X-ray diffraction
118 measurements have confirmed that a significant enhancement
119 in indium content in InGaIn has been achieved by our
120 approach.

2. RESULTS AND DISCUSSION

121 In this work, two different InGaIn-based LED samples have
122 been designed and then grown, aiming to study the influence

of selective epitaxial growth on the optical performance of III-
nitride LEDs grown on a prepatterned template featuring
microhole arrays. A μLED array sample is obtained by our
selective epitaxy growth on the prepatterned n-GaN template
as mentioned above and is denoted as LED A. The other one is
a normal LED sample grown under identical growth but on a
standard planar n-GaN template without any features and is
denoted as LED B.

Silicon-doped n-GaN epiwafers are first grown on *c*-plane
(0001) sapphire substrates using the standard two-step
approach by a metalorganic vapor phase epitaxy (MOVPE)
technique. Initially, a 25 nm GaN nucleation layer is prepared
at a low temperature after the substrate is subject to a
thermally annealing process at a high temperature of 1150 $^{\circ}\text{C}$,
followed by a 1 μm GaN buffer layer, and then another 500 nm
silicon-doped n-GaN layer both grown at a high temperature of
1120 $^{\circ}\text{C}$. For LED A, the n-GaN template is further patterned
into microhole arrays using SiO_2 masks on its top, which is
then used as a prepatterned template for our selective epitaxial
growth.

Figure 1a shows the schematics of our selective epitaxy
growth approach, allowing us to naturally achieve μLED arrays

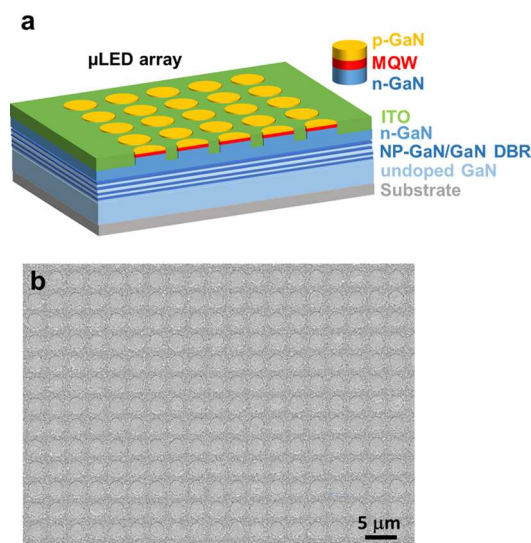


Figure 1. (a) Schematic of selective epitaxy growth and (b) plan-view SEM image for the μLED array epiwafer, showing a diameter of 2 μm and an interpitch of 1.5 μm .

without involving any dry-etching process, i.e., LED A. For the
detailed information on fabricating the prepatterned templates,
refer to the [Experimental Methods](#) section.

Afterward, a standard III-nitride LED structure is selectively
grown on the micropatterned template by MOVPE, namely, a
silicon-doped n-GaN layer is first prepared, followed by an
 $\text{In}_{0.05}\text{Ga}_{0.95}\text{N}/\text{GaN}$ superlattice (SLS) structure as a prelayer,
five periods of InGaIn/GaN multiple quantum wells (MQWs)
as an emitting region, then a 20 nm *p*-type $\text{Al}_{0.2}\text{Ga}_{0.8}\text{N}$ as an
electron blocking layer, and a final 150 nm *p*-type GaN layer.
The total thickness of the overgrown layers is 500 nm, which
matches the thickness of the SiO_2 masks. Due to the SiO_2
masks, the growth of the LED structure takes place within the
microholes only, naturally forming regularly arrayed μLEDs .

A Raith 150 scanning electron microscopy (SEM) system
has been used to characterize the surface morphology of our
regularly arrayed μLEDs . Figure 1b shows a typical plan-view

162 SEM image of our regularly arrayed μ LEDs wafer (i.e., LED
163 A), exhibiting a nice circular shape with an excellent high
164 uniformity in shape, diameter, and interpitch. All μ LEDs are 2
165 μm in diameter and only 1.5 μm in interpitch. Such a small
166 diameter and an interpitch are crucial for manufacturing a
167 high-resolution microdisplay in a compact manner. Further-
168 more, the μ LED pixels share a common n contact while all the
169 p contacts are left open. As a result, our regularly arrayed
170 μ LED epiwafers well match any existing manufacturing
171 technique of microdisplays, for instance, the pick-and-place
172 technology, which has been widely used,³² and the integrating
173 technique using driving transistors based on the silicon CMOS
174 IC to achieve individually addressable μ LED-based microdis-
175 plays.³³

176 A high-resolution X-ray diffractometer (HRXRD) (Bruker
177 D8) has been employed to determine the indium content of
178 the InGaN MQWs by performing ω - 2θ scan measurements
179 along the (002) direction, together with a fitting using the
180 Bruker JV-RADS simulation software. Figure 2a,b shows the

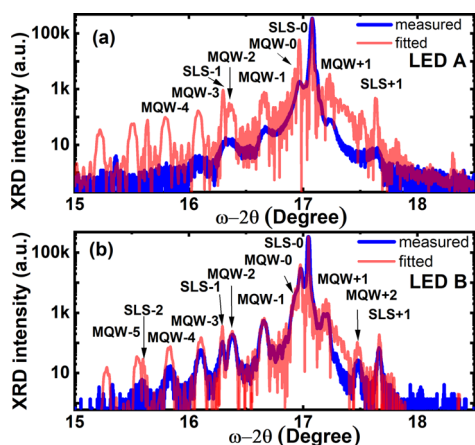


Figure 2. HRXRD ω - 2θ scan curves of the μ LED array sample on a patterned template, i.e., (a) LED A, and the LED sample grown on a standard planar template, i.e., (b) LED B under identical conditions. Fittings have also been provided to determine the indium content in InGaN MQWs.

181 HRXRD ω - 2θ scan curves of our regularly arrayed μ LED wafer
182 (i.e., LED A) and the standard LED wafer (i.e., LED B),
183 respectively. In both cases, the satellite peaks with up to 4 or 5
184 orders from the InGaN/GaN MQWs have been clearly
185 observed. The satellite peaks from the SLS structure as a
186 prelayer have also been observed. Based on a detailed fitting, it
187 can be determined that the indium content of the InGaN
188 MQWs of LED A is 31% and that the thicknesses of the
189 InGaN quantum well and the barrier are 2.2 and 13.8 nm,
190 respectively. In contrast, LED B exhibits 24% indium content
191 in the InGaN MQWs with a 2.6 nm quantum well and a 14.1
192 nm barrier. The XRD fittings are conducted based on fully
193 strained InGaN MQWs for both LEDs. It is well known that
194 strain relaxation will reduce the strain-induced quantum-
195 confined Stark effect (QCSE), leading to a blue-shift in the
196 emission. It means that if the InGaN MQWs are assumed to be
197 strain-relaxed, the indium content should be even higher. In
198 consideration of a higher chance of strain relaxation for the
199 μ LEDs, the fitted values of indium contents represent the least
200 difference between the two LEDs. This direct comparison
201 indicates that enhanced indium content in InGaN MQWs can

be obtained by using our selective epitaxy growth approach on 202
a prepatterned template featuring microhole arrays. 203

Finally, both the regularly arrayed μ LED wafer (i.e., LED A) 204
and the standard LED wafer (i.e., LED B) have been fabricated 205
into LED devices with an area of $330 \times 330 \mu\text{m}^2$. For the 206
detailed information about device fabrication, refer to the 207
Experimental Methods section. For LED A, each LED device 208
consists of a few thousands of 2 μm μ LEDs connected. In this 209
work, the μ LEDs in LED A share a common p contact and n 210
contact, which are driven simultaneously in all electro- 211
luminescence (EL) measurements. However, it is worth noting 212
that our arrayed μ LEDs are designed to make the p contacts of 213
each μ LED left open, providing an opportunity in the future to 214
allow indium bumps to be bonded to an active matrix driving 215
transistors. This means that our regularly arrayed μ LED 216
structure entirely matches any existing individually addressable 217
 μ LED microdisplays. 218

For a direct comparison, the LED B wafer has also been 219
processed under identical conditions in the same batch. All the 220
characteristics of our μ LED chips in the present study have 221
been carried out on bare chips, meaning that we did not use 222
coating or passivation or epoxy or reflector for improving 223
extraction efficiency. Current–voltage (I - V) characteristic and 224
EL measurements have been performed at room temperature 225
in continuous wave (CW) mode using a Keithley 2400 226
sourcemeeter on a probe station equipped with an optical 227
microscopy system. 228

The EL spectra have been measured on the two LED devices 229
under identical conditions aiming to make a direct comparison. 230
For instance, Figure 3a,b shows the EL spectra of the two LED 231
devices measured at a current density of 10 A/cm^2 , 232
respectively. Both spectra exhibit a single emission peak. The 233

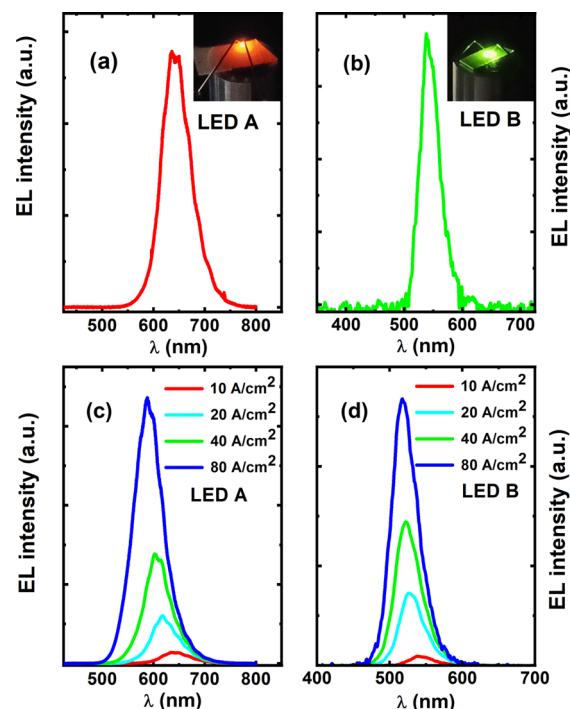


Figure 3. EL spectra measured at 10 A/cm^2 for the μ LED array device, i.e., (a) LED A and (b) LED B, where the insets show their respective emission images. EL spectra measured at increased current densities from 10 to 80 A/cm^2 for (c) LED A and (d) LED B, respectively.

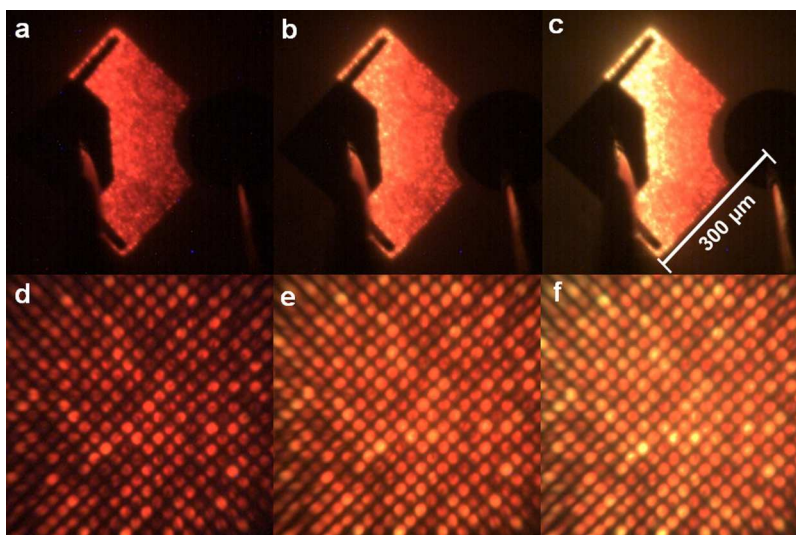


Figure 4. Emission images of the μ LED array device taken using an optical microscopy system as a function of injection current density (4, 8, and 12 A/cm^2) under (a–c) a low magnification and (d–f) a high magnification, respectively.

234 μ LED array device shows a strong emission at an emission
 235 wavelength of 642 nm in the red spectral region. The inset of
 236 Figure 3a exhibits an emission image of the μ LED array chip,
 237 demonstrating red light. In contrast, Figure 3b displays a
 238 strong green emission at 538 nm from the LED B device also
 239 measured at 10 A/cm^2 , and the inset displays its emission
 240 image. This means that the selective epitaxial growth on a
 241 prepatterned template featuring regularly arrayed microholes
 242 results in a red-shift of about 100 nm in emission wavelength in
 243 comparison with the LED grown on a standard planar GaN
 244 surface, although both are grown under identical growth
 245 conditions. As discussed earlier, the growth of InGaIn on a
 246 relaxed layer is beneficial for obtaining high indium content in
 247 InGaIn. Due to no lateral confinement during the overgrowth
 248 within the microholes, the overgrown n-GaN is very likely
 249 strain-relaxed, which leads to an enhancement in the indium
 250 content in the overlying InGaIn MQWs. Combined with the
 251 XRD results, it has been confirmed that our selective epitaxy
 252 growth approach can enhance indium incorporation into GaN
 253 significantly. Figure 3c,d shows the EL spectra of LED A and
 254 LED B, both measured as a function of injection current
 255 density ranging from 10 to 80 A/cm^2 , respectively.

256 In order to demonstrate emitting μ LED pixels, optical
 257 microscopy images have been taken using a micro-EL
 258 measurement system where emissions are collected through
 259 two objective lenses (one 10 \times magnification lens with NA =
 260 0.28 and another 50 \times magnification lens with NA = 0.43).
 261 Figure 4a–c displays the emission images of our μ LED array
 262 chip taken under 4, 8, and 12 A/cm^2 current density,
 263 respectively, while Figure 4d–f provides their corresponding
 264 emission images taken under a high magnification, showing
 265 strong red emissions from individual 2 μm μ LED pixels even
 266 under low current densities. It is worth mentioning that such
 267 low current densities used for the operation of our μ LEDs are
 268 lower than a typical current density (22 A/cm^2) for the
 269 operation of a conventional broad area LED.

270 Both light output powers and luminous flux have been
 271 measured on the bare-chip LEDs bonded on TOS-headers in
 272 CW mode using a LCS-100 integrating sphere equipped with a
 273 CCD APRAR spectrometer. Figure 5a–c shows the output
 274 power, luminance, and EQE of the μ LED array device (i.e.,

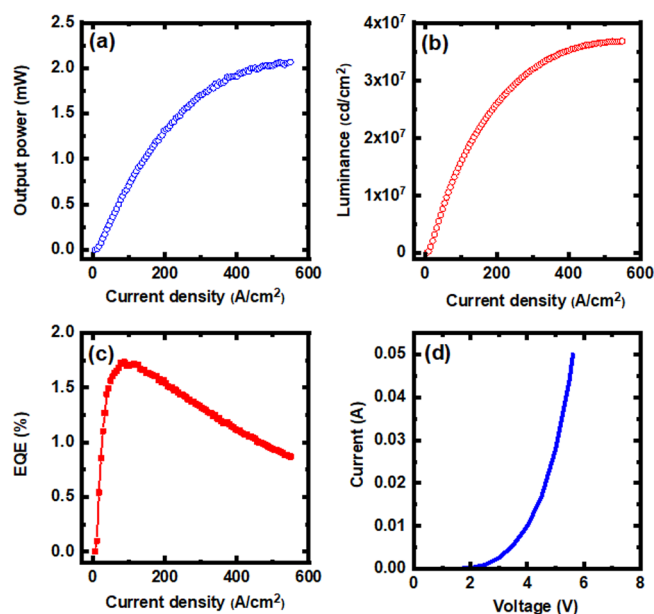


Figure 5. (a) Output power, (b) luminance, (c) EQE, and (d) current–voltage characteristics of the μ LED array device (i.e., LED A).

275 LED A) as a function of injection current density. This
 276 demonstrates that the output power and luminescence increase
 277 monolithically with increasing current density up to 450 A/cm^2
 278 and that a high luminance of 3.5×10^7 cd/m^2 has been
 279 achieved. The peak EQE is about 1.75%. It is worth
 280 highlighting that although there are not any heat-sink
 281 components used, our ultrasmall μ LEDs can still sustain a
 282 high current density of above 450 A/cm^2 , also confirming the
 283 high crystal quality of our μ LED array sample achieved by our
 284 selective epitaxy growth approach. Figure 5d displays the
 285 typical I – V characteristics of LED A measured as a function of
 286 bias, which is similar to that of the LED B device. This also
 287 shows the good electrical property of our μ LED array device.

3. CONCLUSIONS

In summary, we are proposing to employ a selective epitaxy growth approach on a microhole patterned template to significantly enhance strain relaxation, allowing us to not only obtain the natural formation of regularly arrayed μ LEDs but also achieve enhanced indium content in the InGaN/GaN MQWs used as an active region for the μ LEDs. By means of this approach, we have demonstrated red InGaN-based μ LED arrays with a dimension of $2\ \mu\text{m}$ and an interpitch of $1.5\ \mu\text{m}$. A high luminance of $3.5 \times 10^7\ \text{cd/m}^2$ and a peak EQE of 1.7% have been achieved for the red μ LED array chip in a wafer form without any packaging. In contrast, the standard LED grown under growth conditions but on a standard planar GaN template demonstrates green emission. This means that our approach paves the way for achieving long wavelength InGaN-based μ LEDs with ultrasmall dimensions at an elevated growth temperature, at which it is impossible to obtain InGaN-based red LEDs on a standard planar template.

4. EXPERIMENTAL METHODS

4.1. Fabrication of Prepatterned Templates. A 500 nm SiO_2 dielectric film is deposited on the n-GaN template by a plasma-enhanced chemical vapor deposition (PECVD) technique, followed by employing a standard photolithography and then a dry etching technique to selectively etch the SiO_2 dielectric layer down to the n-GaN surface by inductively coupled plasma (ICP), forming regularly arrayed microholes with a diameter of $2\ \mu\text{m}$ and an interpitch of $1.5\ \mu\text{m}$. This prepatterned template will then be used for further selective epitaxy growth. Finally, μ LEDs will be selectively grown only within SiO_2 microhole regions, naturally forming regularly arrayed μ LEDs.

4.2. Device Fabrication. Indium-tin-oxide (ITO) is deposited and then undergoes an annealing process in air at $600\ ^\circ\text{C}$ for 1 min, forming transparent *p*-type contact, while Ti/Al/Ni/Au alloys are prepared as *n*-type contact. Ti/Au alloys are used as *p*-type and *n*-type electrodes. All the characteristics of the LEDs in this paper are conducted on bare chips, namely, no coating, no passivation, no epoxy, or no reflector, which are often employed for obtaining enhanced extraction efficiency.

AUTHOR INFORMATION

Corresponding Author

Tao Wang – Department of Electronic and Electrical Engineering, The University of Sheffield, Sheffield S1 3JD, United Kingdom; orcid.org/0000-0001-5976-4994; Email: t.wang@sheffield.ac.uk

Authors

Peng Feng – Department of Electronic and Electrical Engineering, The University of Sheffield, Sheffield S1 3JD, United Kingdom
Ce Xu – Department of Electronic and Electrical Engineering, The University of Sheffield, Sheffield S1 3JD, United Kingdom
Jie Bai – Department of Electronic and Electrical Engineering, The University of Sheffield, Sheffield S1 3JD, United Kingdom
Chenqi Zhu – Department of Electronic and Electrical Engineering, The University of Sheffield, Sheffield S1 3JD, United Kingdom
Ian Farrer – Department of Electronic and Electrical Engineering, The University of Sheffield, Sheffield S1 3JD, United Kingdom
Guillem Martinez de Arriba – Department of Electronic and Electrical Engineering, The University of Sheffield, Sheffield S1 3JD, United Kingdom

Complete contact information is available at:

<https://pubs.acs.org/10.1021/acsaelm.2c00311>

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Author Contributions

[†]P.F., C.X., and J.B. contributed equally to this work.

Author Contributions

T.W. conceived the idea and organized the project. T.W. and J.B. prepared the manuscript. P.F., X.C., and C.Z. grew all the samples. P.F., X.C., and I.F. performed the material characterization. J.B. and G.M.D.A. fabricated the prepatterned templates and carried out the device fabrication. J.B. conducted the device characterization.

Notes

The authors declare no competing financial interest.

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