Innovative Development of Next Generation and Energy Efficient Solid State Light Sources for General Illumination

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Abstract

This two year program resulted in a novel broadband spectrally dynamic solid state illumination source (BSDLED) that uses a dual wavelength light emitting diode (LED) and combinations of phosphors to create a broadband emission that is real-time controllable. Four major focuses of this work were as follows: (1) creation of a two terminal dual wavelength LED with control of the relative intensities of the two emission peaks, (2) bandgap modeling of the two terminal dual LED to explain operation based on the doping profile, (3) novel use of phosphor combinations with dual LEDs to create a broadband spectral power distribution that can be varied to mimic a blackbody radiator over a certain range and (4) investigation of novel doping schemes to create tunnel junctions or equivalent buried current spreading layers in the III-nitrides. Advances were achieved in each of these four areas which could lead to more efficient solid state light sources with greater functionality over existing devices. The two-terminal BSDLED is an important innovation for the solid-state lighting industry as a variable spectrum source. A three-terminal dual emitter was also investigated and appears to be the most viable approach for future spectrally dynamic solid state lighting sources. However, at this time reabsorption of emission between the two active regions limits the usefulness of this device for illumination applications.
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Executive Summary

The focus of this two year program was to develop new technical innovations that address, high risk bridging research and development that would enable the more rapid short term progress of new solid-state lighting sources. This approach is critical for the early and more immediate adoption of solid-state lighting technology into general illumination. This need is well illustrated in the energy savings detailed in the Arthur D. Little, Inc. report [1] and other road mapping documents produced by the DOE [2]. Solid-state lighting technology has the possibility of breaking the ‘vicious cycle’ within the lighting industry with its emphasis on ‘low-first-cost’ that slows the development and adoption of improved lighting solutions as detailed in Vision 2020 [3]. Solid-state lighting has interest to the DOE because it is clean and energy efficient compared with many of the traditional general illumination sources. This results in both energy conservation and a corresponding reduction in fossil fuel use.

Currently, all of the high brightness blue LEDs that are used for bi-color YAG phosphor white light sources are grown on sapphire or, to a lesser extent, SiC substrates. The large lattice and coefficient of thermal expansion (CTE) mismatch between the sapphire (or SiC) and the GaN results in a high density of defects ($10^{8-10}$ cm$^{-2}$) throughout the device structure. This was not an issue for the first devices produced using GaInN layers in the active region to produce emission in the blue/green (450-525nm) because exciton localization due to compositional fluctuations meant that the defects were not optically active. However, the need to increase the power density in the device to increase the total lumen output for lighting applications and the need to shift the emission wavelength of the device towards the UV has meant that dislocations are now an issue as the mechanism of exciton localization in InGaN is now negated. While growth on native GaN substrates would reduce dislocation densities, these are not widely available or cost-effective. Thus the defect density of GaN grown on non-native substrates must be reduced. This was one of the original objectives of the program, and the year one milestone of achieving III-N growth with a defect density of $10^{7-8}$ cm$^{-2}$ on a non-native substrate was met on schedule, as shown in Table 1. However, after year one the program narrowed its focus to developing the dual emitter and appropriate phosphor combination needed achieve the spectrally dynamic white LED, on which this report is focused.

<table>
<thead>
<tr>
<th>Objective</th>
<th>Metric</th>
<th>Complete</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lattice-matched emitters on non-native III-Nitride substrates</td>
<td></td>
<td></td>
</tr>
<tr>
<td>III-Nitride defect density</td>
<td>$10^{7-8}$ cm$^{-2}$</td>
<td>Yes</td>
</tr>
<tr>
<td>Substrate removal active area</td>
<td>1 cm$^2$</td>
<td>Yes</td>
</tr>
<tr>
<td>Next generation solid state emitters</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phosphor efficiency</td>
<td>50-60%</td>
<td>Yes</td>
</tr>
<tr>
<td>Optimized emitter wavelength</td>
<td>400 nm</td>
<td>Yes</td>
</tr>
<tr>
<td>Equivalent lumen output</td>
<td>30</td>
<td>Yes</td>
</tr>
</tbody>
</table>
The result of this program is a functionally new broadband white LED using a dual wavelength emitter to preferentially pump phosphors in a mixture. The spectral power distribution (SPD) of the source can be dynamically tuned by varying the drive current of the device which can then be pulse width modulated to normalize the output intensity. A two terminal dual wavelength LED was used to pump a mixture of phosphors with varied excitation spectrum. One of the LED emission bands was used to preferentially excite part of the phosphor mixture while the second emission band was used to excite the remaining phosphors in the mixture. Varying the drive current of the LED allowed tuning of the relative intensity of the two LED emission bands. Careful selection of the phosphors and LED structure achieved a CCT range for the final device of 3200K – 5300K.

Several novel structures and materials were investigated and developed for use in the final device including: p-type doping schemes, transition metal doping of junctions, doping of quantum well barriers, phosphor combinations of varied excitation spectra. Two device designs were considered for the dual wavelength pump device. The first was a three terminal design incorporating a tunnel junction as a buried p-type contact. The extremely high p-type doping required for a suitable tunnel junction prohibited the use of the three terminal LED as a pump source. However, substantial work investigating novel p-type doping schemes was performed achieving hole concentrations above 1 \( \times 10^{18} \text{ cm}^{-3} \).

A novel hybrid doping scheme employed high and low temperature GaN:Mg layers to achieve high hole concentrations (~1 \( \times 10^{18} \text{ cm}^{-3} \)) while maintaining material quality (rms surface roughness < 10 Å). Tunnel junctions were achieved using hybrid p-type layer as the p++ layers; however, suitable performance (< 0.5 V @ 20 mA) for the tunnel junction reverse current was not achieved. Silicon and magnesium doped short period superlattice structures consisting of 5 Å periods of GaN/InGaN were also investigated as p++ layers. Hole concentrations ~ 3 \( \times 10^{18} \text{ cm}^{-3} \) were achieved; however, very little reverse current was observed when the SPS layers were incorporated into a tunnel junction structure.

The final dual wavelength LED employed was a two terminal LED consisting of 3 quantum wells (QW) designed to emit at ~460 nm followed 3 QWs designed to emit at 400 nm. The barrier between the third and fourth QWs was doped with silicon to limit hole injection across the barrier and enable control of the carrier recombination in each QW region. Controlling the carrier recombination allowed for control of the SPD of the LED. The variation range for the LED was less than that that could be achieved with a three terminal device, but was suitable for pumping the phosphors chosen.

The final phosphor combination incorporated 5 phosphors including a UV-to-white, green, YAG, orange and red. The final phosphor combination was developed to work with the two terminal dual wavelength LED and achieved a CCT range of 3200K-5300K.

Efficiency in these devices is not expected to be drastically different from other high brightness LEDs because the fundamental processes are not drastically changed. On the other hand some gain in efficiency could be seen by properly choosing phosphors and pump wavelengths to minimize stokes shift losses seen in other phosphor converted LEDs. Further optimization of the two terminal dual wavelength LED will achieve the most efficient operating point and minimize the amount of drive current variation needed to tune the SPD. Finally, optimization of these devices can enhance solid state light sources and potentially increase the efficiency of phosphor converted LEDs.
Introduction

A novel solid state illumination source has been developed. A two terminal dual LED has been created with the ability to control the relative intensities of the two emission peaks by varying drive current. The development of a three terminal dual LED as a pump source was prohibited by the need for a III-nitride tunnel junction, that proved unattainable in the scope of this work due to the need to degenerate dope the tunnel junction. Doping profiles have been used to extend the dynamic range of the two terminal dual LED over other reported devices. Operation of the two terminal dual LEDs is explained as a function of drive current. In addition, novel use of phosphor mixtures allows the creation of a broadband spectral power distribution that can be varied using a dual LED as an excitation source. Combinations of phosphors that have varied excitation spectra provide the ability to selectively excite different phosphors with the different LED emission peaks. First and second generations of the two terminal dual LED and the phosphor combination are discussed. The final source has the ability to mimic the light of a blackbody radiator over a range of 3200 K - 5300 K.

Theory

The development of the broadband spectrally dynamic LED required a number of other developments including a phosphor combination, dual wavelength LED and a tunnel junction. Several experiments for each of these structures are described. Novel p-type doping schemes were a major part of the work.

High hole concentrations in GaN are difficult to achieve due to the high thermal ionization energy (~180 meV) of Magnesium (Mg) in GaN. At room temperature less than 1% of Mg acceptors are typically electronically active. Therefore, in order to achieve acceptable hole concentrations, high Mg incorporation far above normal doping densities is required. This high doping concentration, however, decreases the crystalline quality of the film, which consequently increases its resistivity and makes it difficult to obtain good electrical contacts to the material. Resistive films cause current crowding in LEDs. A semitransparent contact is deposited onto the entire surface of the p-type region to compensate for current crowding; this is followed by a smaller but thicker bonding pad. The cost of using this semitransparent current spreading layer is a reduction in light extraction from the device by not entirely transmitting the incident light. Post-growth treatment of GaN:Mg is another important issue. Typically, material is annealed in a non-\text{H}_2\text{ environment to break the compensating Mg-H bond that occurs in metal organic chemical vapor deposition grown GaN:Mg. Reports show that optimization of the annealing process can lead to more effective activation of Mg acceptors. Some reports suggest that annealing in air (containing O}_2\text{) can further increase Mg activation in GaN[4].}

Current crowding poses a problem for buried p-type contacts in an LED. It is not possible to deposit a metal current spreading layer similar to a top p-type contact and a simple bonding pad will not provide enough current spreading for an efficient device. Hence, a tunnel junction is one solution to contacting buried p-layers. In this work, p+ GaN layers have been developed for use in a tunnel junction that may be used in two and three terminal devices for solid state lighting applications.
Tunnel Junction Theory

Electron tunneling results from a quantum mechanical phenomenon from the wave nature of electrons [5]. When an electron wave is incident with energy less than the energy barrier height, there is a probability that this electron can penetrate this energy barrier, and the penetration depth depends on the barrier height. Figure 1 shows a rectangular barrier with potential height of V, and thickness of d.

![Potential barrier seen by an electron.](image)

For an electron wave with energy $E_v$, the tunneling probability is given by Equation 1:

$$T(E_v) = \frac{1}{1 + \frac{V^2 \sinh^2(kd)}{4E_v(V - E_v)}}$$

$$k = \sqrt{\frac{2m(V - E_v)}{\hbar^2}}$$

where $m$ is the effective mass of an electron, $\hbar$ is the plank constant. The tunnel probability is high for very thin barriers. For arbitrary shape of potential barriers, a numerical solution can be obtained by approximating the potential barrier with multiple sub-potential layers, each sub-potential layer can be treated as a constant potential layer. Equation 1 is used to calculate the tunneling probability for each of these constant potential barriers.

In semiconductor materials, carriers can be generated by band to band tunneling when a strong electric field presents, for example, the electric field greater than 10 mV/cm. For a reversed biased, heavily doped, thin pn junction, if the electron penetration depth is larger than the depletion width of this pn junction electrons at valence bands at p side can penetrate into conduction bands at n side to become free carriers. This is the so called tunneling junction. The uses of a tunnel junction (TJ) in III-Nitride LEDs have been demonstrated recently [6]. The reverse biased TJ is to improve the lateral current spreading. With a TJ, n-GaN can be used to replace p-GaN as a top cap layer. Commonly, a so called semi-transparent p-Pad is usually applied on the top of p-GaN to overcome the low conductivity of p-GaN In conventional LEDs. The n-GaN has much higher conductivity, about 100 times, compared to p-GaN , and the reverse biased TJ supplies holes to p-GaN adjacent to the active region through lateral current. As a result, the application of n-GaN top contact layer can improve the current spreading of the LED, and increase light extraction by avoiding using semitransparent contact. Another benefit of using a TJ is the simplification of the LED fabrication process because n-GaN can be used as to implement both the top p-contact and lower n-contact as shown in Figure 2.
The TJ usually is fabricated using heavily doped p++/n++ either InGan/GaN or GaN/GaN bi-layer. One example use an InGan/GaN bi-layer with layer thicknesses of 15 nm/30nm [7]. The p-type doping Mg level is about $10^{20} \text{cm}^{-3}$, and n-type doping Si level is about $6 \times 10^{19} \text{cm}^{-3}$ for this tunneling junction. This TJ is incorporated into a resonant cavity LED with violet light emission. The resonant cavity of this LED uses two pairs of DBRs: the top one is a SiO$_2$/HfO$_2$ above the TJ, and the bottom one is an AlGaN/GaN DBR grown on a GaN buffer layer. A TJ in the second example is fabricated using a GaN/GaN layer with layer thicknesses of 10nm/10 nm. The p+ GaN layer is doped with $3 \times 10^{19} \text{cm}^{-3}$ of Mg; the averaged Si doping level in the n+ GaN layer is about $10^{20} \text{cm}^{-3}$. A n-GaN is grown on the top of this TJ to replace the conventional p-GaN as a top contact layer. One disadvantage of the TJ is its large forward voltage, about 1 volt higher than that of a conventional LED because the TJ add a series resistance to the LED. However, the overall optical power of a LED is claimed to double compared to the conventional LED with a top semitransparent p-Pad [6]. A necessary characteristic to tunneling is the negative differential resistance (NDR) region in the forward bias operation of a tunnel diode, because it shows the existence of band-to-band tunneling (Figure 3) [8].
In the NDR region a decrease in the density of states leads to a decrease in tunneling probability and a subsequent decrease in current density [9]. Kane derived an equation for the current density of a tunnel diode as shown in Equation (2) [9].

\[ j = \frac{e}{18h^3} m^* \frac{-\pi \sqrt{m} E^3}{E_s} e^{2\sqrt{2hE}} \left( \frac{E}{2} \right) \int \left[ f_1(E_1) - f_2(E_2) \right] \left[ 1 - e^{-\frac{2E}{E_s}} \right] dE \]

where \( E_s \) is the smaller of \( E_1, E_2 \). Kane’s work will be used as a basis for understanding and quantifying the observed tunneling mechanisms in the GaN device. Full analysis of the current-voltage characteristics will lead to a better understanding of the device, although tunnel junctions for this work will be operated in the reverse bias region. A better understanding of GaN based tunnel junctions will help to optimize them for incorporation as more efficient contacting layers in LEDs. Currently, the presence of a tunnel junction increases the series resistance of the device by 33% [10] over a typical LED with semitransparent current spreading layer. A goal of this research will be to reduce the series resistance of GaN tunnel junctions.

**Hybrid p-type Doping**

Hybrid doping is a novel technique developed for this project in which dopant layers are grown at alternating high and low temperatures in order to achieve both the good crystal quality of high temperature growth and the high doping potential of low temperature materials. This was the first doping scheme employed in the tunnel junction development. Three sets of runs were performed to optimize hole concentration and material quality. The first set of growths was at high temperature (~1030°C) in which the molar flow ratio of \( \text{Cp}_2\text{Mg}/\text{TMGa} \) was changed from 0.00556 to 0.0667 in order to incorporate more Mg into the films and maintain a high quality epitaxial layer. The second set of runs repeated the change in molar flow ratio of \( \text{Cp}_2\text{Mg}/\text{TMGa} \) from the high temperature runs, but with a lower deposition temperature (~980°C). The third set of growths, hybrid p-type doping, consisted of alternating layers of high and low temperature p-type GaN repeated 10 times. The high temperature layers were doped with a molar flow ratio of 0.0334 \( \text{Cp}_2\text{Mg}/\text{TMGa} \), while the low temperature layers were highly doped with a molar flow ratio of 0.0459 \( \text{Cp}_2\text{Mg}/\text{TMGa} \). It is important to note that in the first and second set of growths the molar flow ratio was set by changing the \( \text{Cp}_2\text{Mg} \) flow and maintaining a consistent TMGa flow. However, in hybrid p-type doping, the molar flow ratio was set by changing either the TMGa flow or the \( \text{Cp}_2\text{Mg} \) flow. It is believed that decreasing the amount of available gallium allows for higher rates of Mg incorporation. For all growths, \( \text{NH}_3 \) flow was held constant.

Several tunnel junctions were grown where the p++ layer was varied according to the three sets of growth runs described in this work (a general schematic of the devices is shown in Figure 4).
A GaN buffer layer was grown on top of a bare sapphire substrate at low temperature (500°C to 600°C) followed by high temperature (~1050°C) undoped GaN and then an n-type GaN with SiH₄ as an n-type dopant at a molar flow ratio of 3.51 to TMGa. The TMGa was switched with TEGa for the n+ layer (highly doped n-type GaN), which was grown at low temperature (~980°C), and SiH₄ was used as an n-type dopant at molar flow ratio of 48.3 to TEGa. To ensure a better interface between the n+ and p+ layers a thin layer of undoped GaN was introduced as a barrier to magnesium diffusion. The device was capped with high quality p-type GaN.

The activation process was also changed, in addition to varying the temperatures and molar flow ratios of Cp₂Mg/TMGa. A standard rapid thermal annealing (RTA) process was used at 800°C with N₂ ambient. Also, lower temperature (500°C) atmospheric anneals were performed after some reports indicated that oxygen could enhance the activation process [4]. The atmosphere anneals were done at times ranging from 5 minutes to 3 hours.

Secondary Ion Mass Spectroscopy (SIMS) was used to measure Mg incorporation in the layers. Hall measurements were used to determine hole concentration and atomic force microscopy was used to measure surface quality. Scratch diodes and lithographically fabricated diodes were used to test for tunneling in the tunnel junction devices.

III-Nitride Tunnel Junctions with Hybrid Doping

III-nitride tunnel junctions were grown by MOCVD. Various novel doping schemes to create highly doped p-type GaN were employed with Mg as the acceptor ion. Highly doped p-type material (p++) is necessary for creating efficient tunnel junctions that act as buried current spreading layers and provide high quality contacts to p-type GaN. Molar flow ratios and growth temperature were varied to achieve maximum doping concentration. SIMS and Hall Effect were used to measure Mg incorporation and carrier concentration, respectively. The maximum carrier concentration achieved employed a hybrid growth scheme that achieved high carrier concentration and high quality material by alternating layers of high and low growth temperature material. The hybrid growth scheme developed here is a novel growth process that enhances p-type GaN. Successful GaN tunnel junctions were grown using the hybrid p-type doping scheme developed in this work. I-V measurements were taken on these devices to evaluate the effectiveness of the devices.
Short Period Superlattice Doping

GaN/InGaN short period superlattice (SPS) structures doped with Si or Mg were investigated as possible doping schemes for a III-nitride tunnel junction. SPS structures have been suggested as and reported to have the potential for achieving higher carrier concentrations than bulk layers. The superlattices in this work had periods of 5 Å GaN and 5 Å InGaN. The test structures were each comprised of 20 periods. Hall and PL spectroscopy measurements were performed on the SPS test structures to evaluate their quality. Junctions were then grown using the Mg doped SPS and Si SPS structures and evaluated as tunnel junction devices.

SPS test structures were grown with varying levels of silicon and magnesium doping. The test structure consisted of 20 periods of alternating Si/Mg:InGaN and GaN layers of 5 Å thickness grown on top of 1.5 µm of undoped GaN on a sapphire substrate. Both the InGaN and GaN layers were grown at ~800°C, which is the normal growth temperature for InGaN. However, high quality GaN is normally grown at ~1000°C. In this structure the quality of the material is preserved as characterized by AFM, XRD, and PL.

Mn Doping

Mn doping was introduced into a tunnel junction structure similar to the one mentioned previously that used the hybrid p-type doping scheme to assess its effect on tunneling behavior. Mn dopants tend to have energy levels in the middle of the bandgap for GaN, which may lead to defect enhanced tunneling. Normally, as the tunnel junction is driven into reverse bias, tunneling may occur when the valence band of the p-side is at higher potential than the conduction band of the n-side. The tunneling probability is a function of the width of the tunnel barrier, however, if a state exists in that barrier it may act as a transition point for carriers, and improve the tunneling probability as shown in Figure 5.

![Figure 5: Tunnel junction in reverse bias illustrating defect assisted tunneling](image_url)

Several devices were grown and fabricated. The layer structures shown in Figure 6 are modified from the tunnel junctions employing the hybrid doping scheme mentioned before.
Phosphor Test Bed

First Generation Phosphor Test Bed

The basis for a BSDLED was demonstrated by pumping a combination of phosphors with multiple wavelengths of excitation, and then varying the relative intensities of those pumps. Such a device is a novel concept and will play an innovative role in solid state general illumination.

A wide variety of phosphors that are excited by a range of pump wavelengths exists. In fact, a single phosphor may respond differently to different pump wavelengths. Thus, a combination of multiple phosphors with differing and exclusive excitation spectra can be simultaneously pumped by multiple wavelengths to produce white light. The relative intensities of the pump peaks can then be varied to change the output of the total device and therefore allow dynamic tuning of the correlated color temperature (CCT). Such a device combines the controllability seen in RGB-LEDs and the broadband characteristics of PC-LEDs. It is believed that such innovations will prove instrumental in the success of solid state general illumination sources that require high quality dynamic white light.

Second Generation Phosphor Test Bed

A wider range of phosphors were combined to create a mixture that would have CIE coordinates closer to the Planckian Locus. In addition to the phosphors used in the first generation test bed, YAG:Ce and an orange emitting phosphor were used to enhance the spectral power distribution of the source. Another stipulation to the second generation phosphor test bed was that it work with the second generation dual LEDs described below. These devices do not have the dynamic range that is achievable with two discrete devices, which can have $\beta$ ranging from 0 to $\infty$.

Dual LEDs

First Generation Two Terminal LED Structures

LED structures with multiple quantum well regions designed to emit at different wavelengths have been grown. It has been observed that the emission mechanism for each of these MQW regions is the same as that seen in a single MQW region LED.
Dual MQW region LEDs, Figure 7, for efficient pumping of multiple phosphors have been grown by MOCVD for use in broadband, white solid state light sources.

![Figure 7: Layer schematic of 1st generation Dual LED](image)

Blue (460 nm) and Violet/UV (~400-420 nm) emitting MQW regions were incorporated into a single device and show recombination mechanisms similar to single MQW region devices. The introduction of a spacing region successfully separated the electroluminescent emissions, and two distinct emission peaks were observed. A large factor in the efficient radiative recombination in these devices is the localization of carriers by indium compositional fluctuations in the InGaN quantum wells. Photoluminescence measurements were carried out to determine the physical mechanism behind light emission in these devices. Optical recombination in low-dimensional InGaN quantum structures strongly depends on the localization of carriers in quantum dot-like structures [11,12,13,14]. Inhomogeneities in the indium concentration on the nm-scale provide potential fluctuations in the band gap, and the carriers are trapped in islands that may provide 3D-quantisation up to elevated temperatures [13]. An increased overlap of hole and electron wave functions is induced by the localization of carriers in the nanoscale islands. The average localization energies were determined by an Arrhenius plot of the luminescence intensity following the approach suggested by Adelmann et al [15]. The two activation processes are necessary to describe the data in both low and high temperature regimes.

All the LEDs in this work consisted of InGaN/GaN quantum wells in the active region. The emission wavelength was controlled by changing the growth temperature of the wells, which affected the growth rate and the indium incorporation. The barrier was grown in two stages; the first stage was grown without hydrogen and the second stage was grown with hydrogen and at a higher temperature and rate than the first stage. Multiple growths were completed to optimize the growth rate of the well and the two stage barrier.

A series of systematic growth runs were completed at various temperatures to better understand the effects of temperature on the indium incorporation in the quantum wells. Active regions were developed specifically for 460 and 400-420 nm emission. X-ray diffraction and structure simulation were used to study changes in the indium concentration with respect to the growth temperature and the thickness of the wells against the emitted and PL wavelength.
Next, a series of devices containing a 460 and a 400-420 nm set of multi-quantum well regions, a dual MQW region structure, separated by an undoped spacing region was grown as shown schematically in Figure 7. The dual MQW region emitter incorporated the optimized MQW growth conditions for the respective emission wavelengths. The device structure development focused on the 3 layers shown below:

- First set of MQW (460 nm)
- Spacing layer between the two MQWs
- Second set of MQW (400-420 nm)

The 460 nm MQW was placed at the bottom so that it does not reabsorb the emission from the 400-420 nm MQW. A spacing layer was introduced between the two MQWs to mitigate interaction. The two respective MQWs were split into three growth stages: the quantum well and a two stage barrier. As discussed above, temperature was used to vary the indium incorporation into the well to control the desired emission wavelengths. During device optimization it was determined that varying well and barrier thickness and different growth temperatures were the best approach for the two sets of MQWs.

Thin and thick undoped GaN, n-type GaN and low aluminum concentration GaN layers were studied for the spacing layer at both low (below 800˚C) and high (above 900˚C) temperature growths. In addition, different precursors for gallium (TMGa and TEGa) were evaluated to optimize the interface layer.

**Second Generation Two Terminal Dual Wavelength LEDs**

The first generation of two terminal dual wavelength had an emission spectrum with two distinct peaks which could be used to efficiently couple into multiple phosphors of different excitation spectra. However, dynamic control of the relative intensities of the two peaks was limited in these devices. A second generation of devices were designed to increase the ability to control the relative intensity of the two peaks and simplify the structure. A series of experiments were performed to determine the behavior of these devices. A basic schematic of the device layer structure is shown in Figure 8.

![Figure 8: Layer schematic for 2nd generation Dual LEDs](image)

The samples were grown in the highly modified commercial MOCVD reactor used for the rest of the work. The modifications included changes to the switch block, which enables the use of novel precursors, which was used for Mg doping. The active region consisted of six quantum wells with three designed to emit at 400 nm and three designed to emit at 460 nm. The first three grown were designed to emit at 400 nm and
will be referred to as QWs 1-3, while the last three grown were designed to emit at 460 nm and are referred to as QWs 4-6. Below the active region is an n-type GaN region with carrier concentration of $10^{18} \text{cm}^{-3}$ electrons. Above the active region a p-type GaN region with hole concentration $10^{17} \text{cm}^{-3}$. The n and p-type regions used in these device are typical of common LED structures. In this study the most important metric was the relative intensity of the two emission peaks. $\beta$ is defined as the ratio of the long wavelength intensity to the short wavelength intensity. In the case of these samples the long wavelength quantum wells are positioned nearer the n-type GaN layer and the short wavelength quantum wells are positioned near the p-type gallium nitride. In effect, $\beta$ is a measure of the amount of electron hole pairs recombining in the long wavelength quantum wells as compared to the short wavelength quantum wells, which is dependent on the current injection into both of those regions. $\beta$ will be used throughout this section to denote this ratio in the text and figures.

**Results and Discussion**

**Hybrid p-type Doping**

The variation of $\text{Cp}_2\text{Mg}/\text{TMGa}$ ratio at high temperatures had little effect on hole concentration as measured by Hall effect or Mg incorporation as measured by SIMS. At higher deposition temperatures with Mg incorporation of $5 \times 10^{19} \text{cm}^{-3}$, hole concentrations of $1 \times 10^{17} \text{cm}^{-3}$ were achieved for RTA and atmospheric annealed samples, and surface morphologies remained good, up to a $\text{Cp}_2\text{Mg}/\text{TMGa}$ ratio of 0.0459 as shown in Table 2.

<table>
<thead>
<tr>
<th>Mg/Ga Precursor Flow Ratio</th>
<th>0.014</th>
<th>0.050</th>
<th>0.100</th>
<th>0.025 (HT)/0.045 (LT)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RMS Roughness (Å)</td>
<td>3.3</td>
<td>9.7</td>
<td>25.5</td>
<td>7.2</td>
</tr>
<tr>
<td>Hall Effect (cm$^{-3}$)</td>
<td>$1 \times 10^{17}$</td>
<td>$3 \times 10^{17}$</td>
<td>$2 \times 10^{18}$</td>
<td>$2 \times 10^{18}$</td>
</tr>
<tr>
<td>AFM Scan (4µm x 4µm)</td>
<td>![Scan Image]</td>
<td>![Scan Image]</td>
<td>![Scan Image]</td>
<td>![Scan Image]</td>
</tr>
</tbody>
</table>

However, at lower deposition temperatures with Mg incorporation of $1 \times 10^{20} \text{cm}^{-3}$, hole concentrations $>1 \times 10^{18} \text{cm}^{-3}$ were achieved with atmosphere annealing and increased $\text{Cp}_2\text{Mg}/\text{TMGa}$ flow ratio, but these samples exhibited poor surface morphology. Finally, the hybrid p-type doping, described above, achieved good surface morphology and hole concentrations $>10^{18} \text{cm}^{-3}$ using the standard RTA activation process, while no atmospheric annealing was performed on these samples as mentioned in the preceding section. In the hybrid doping, the balance between the Mg incorporation and the surface quality depended on the TMGa molar flow and the layer thickness of the respective temperature growths. Different $\text{Cp}_2\text{Mg}$ molar flows were evaluated for the high and low
temperature growth layers while keeping the TMGa molar concentration constant, but regardless of temperature they lead to poor surface quality and thus poor contacts. Lowering the Cp₂Mg molar flow greatly improved the surface quality, but it also reduced the carrier concentration and a working tunnel junction could not be achieved. A substantial improvement in the carrier concentration, from $1 \times 10^{17}$ cm$^{-3}$ to $1 \times 10^{18}$ cm$^{-3}$, without diminishing surface quality was seen when the Cp₂Mg molar concentration was kept constant for both the low temperature and high temperature layers, while the TMGa molar concentration was reduced for the low temperature layers.

Tunnel junctions employing p+ layers from each of the three growth runs exhibited different properties. Junctions with a high temperature p+ layer exhibited no tunneling, only typical diode behavior; this was true for both annealing processes. On the other hand, low temperature p+ layers showed tunneling when the sample was atmosphere annealed, but not after the standard RTA, Figure 9.

![Tunnel Junction with Low Temperature p+ Layer](image)

**Figure 9:** Comparison of an atmosphere anneal to standard RTA, and its affect on p+ layer, and tunneling

RTA under a N$_2$ ambient showed no effect on surface morphology as viewed under an optical microscope. Conversely, the atmosphere annealing caused a deterioration of the surface and was not used with the hybrid doped devices. Diodes (tunneling and normal) processed with the atmosphere anneal showed higher resistance.

**Discussion of Hybrid Doping**

The first method for p+ doping did not allow for enough Mg incorporation into the GaN. Ga incorporation is more efficient at higher temperatures, blocking the incorporation of Mg. However, the quality of GaN is preserved, as observed by surface inspection. At lower growth temperatures it is possible to incorporate more Mg and activate more acceptors, but the quality of the crystal surface tends to deteriorate. If the lower temperature p+ layers are used as a buried contact, the overgrown layers suffer from poor crystallinity, and a complex device becomes impossible.
Hybrid doping of the p+ region combines the good crystallinity of higher temperature GaN:Mg with the higher doping potential of low temperature materials. Higher Mg incorporation rates are achievable, by decreasing the TMGa flow relative to the Cp2Mg flow, due to the lack of available Ga that would normally prevent Mg incorporation. In addition layers with high Cp2Mg/TMGa molar ratios were kept thin to minimize disturbance to the crystal. Tunneling was achieved as shown by the I-V curves in Figure 10. The inset shows an I-V curve displaying N-type negative differential resistance, which is an identifying characteristic of tunnel junctions.

![Tunnel Junctions Using Hybrid p+ Doping](image)

Figure 10: Current-Voltage data for several Tunnel junctions using the hybrid growth scheme
Inset plot shows N-type differential resistance

Such a device is a good candidate for a buried contacting and current spreading layer in other devices, eliminating the need for the p-type material to be the top layer. Tunnel junctions have also been demonstrated as fully transparent current spreading layers on the top of devices [6].

Atmospheric annealing in normal atmospheric conditions was also evaluated as a way of increasing the activation of Mg in GaN. While some improved carrier concentrations were observed with Hall measurements, a tradeoff came in material quality deterioration. Although there may be some optimal condition for activation without unacceptable material degradation, such a method is unsuitable for a buried p+ layer, because it is essentially capped from the oxygen that is believed to cause higher activation efficacy.

**Short Period Superlattice Doping**

Both n- and p-type SPS doping structures were fabricated. The n-SPS structures were doped with Si, while the p-SPS structures were doped with Mg. Both of these structures were grown on sapphire substrates with a 2 μm GaN buffer layer followed by 20 periods of alternating 5 μm thick doped InGaN and 5 μm thick GaN. Figure 11 illustrates the structure for the Mg doped p-SPS structure.
Figure 11: Schematic of p-SPS structure

**Si Doped SPS Structures**

AFM measurements provide a RMS surface roughness of 10 Å, while Figure 11 is PL data for the SPS structure showing a GaN peak at 3.4 eV. Hall effect measurements showed electron carrier concentrations in excess of $1 \times 10^{20} \text{ cm}^{-3}$.

**Mg Doped SPS Structures**

Magnesium doping was introduced instead of silicon to investigate the properties of a pSPS structure. The growth was performed at 800°C for both the GaN and InGaN. The doping concentration was maximized using feedback from PL spectroscopy and Hall effect measurements. PL spectra of three samples are shown in Figure 12, Figure 13 and Figure 14.
Figure 12: PL spectra of baseline pSPS doping

Figure 13: PL spectra of twice baseline pSPS doping
Figure 14: PL spectra of half baseline pSPS doping

The flow of Mg into the chamber during growth is the only difference in these structures. The growth was performed at 800°C for both the GaN and InGaN. Sample A is shown as a baseline doping concentration. Sample B has twice the Mg flow and Sample C has half the Mg flow. Each of the PL spectra shows strong Fabry-Perot oscillations indicating high quality interfaces in the superlattice. Each spectra also shows a peak near 2.8 eV arising from the superlattice InGaN composition. Samples A and B have another peak in their spectra that is due to defects in the material from over doping of Mg. The relative intensity of the defect band to the SPS peak indicates the level of defects in the material. For instance, in sample B which has twice as much Mg flow as sample A, the defect band nearly washes out the SPS peak. However, in sample C there is no defect band, indicating that there is no overdoping from Mg. This explanation is corroborated by the results of Hall effect measurements shown in Table 3.

Table 3: Hall measurements for p-GaN samples

<table>
<thead>
<tr>
<th></th>
<th>Sample A</th>
<th>Sample B</th>
<th>Sample C</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hole Concentration</strong></td>
<td>$1.99 \times 10^{18}$</td>
<td>$8.86 \times 10^{17}$</td>
<td>$3.65 \times 10^{18}$</td>
</tr>
<tr>
<td><strong>Mg Flow</strong></td>
<td>Baseline</td>
<td>Twice</td>
<td>Half</td>
</tr>
</tbody>
</table>

Carrier concentrations for pSPS samples. The hole concentration for sample C is higher than that of samples A and B which indicates that the samples are suffering from compensation effects from over doping of Mg.

**SPS junctions**

The SPS test structure described in the previous sections were then used to form a pn junction. Two variations of the structure are shown in Figure 15.
The n SPS layer was the same as above and the pSPS layer was the same Sample C. Current voltage measurements were performed on the devices to assess them as tunnel junctions, however, very little reverse current could be achieved. It is unclear whether further optimization of these structures would lead to a tunnel junction.

**Mn Doped Junction**

Mn has been introduced into the intrinsic region at the junction between the n++ layer and p++ layer. Table 4 shows the expected Mn concentrations in each of the samples. All other parameters were held constant except the flow of Mn entering the reactor.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mn Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low Mn Doping</td>
<td>0.5%</td>
</tr>
<tr>
<td>Medium Mn Doping</td>
<td>1%</td>
</tr>
<tr>
<td>High Mn Doping</td>
<td>2%</td>
</tr>
</tbody>
</table>

Mn doping for tunnel junctions  The current voltage results are shown below in Figure 16.
It can be seen that each of the samples containing Mn in the junction has reduced reverse current as compared to the junction without Mn. One explanation for this is that the presence of Mn impurities in the junction increases carrier scattering causing more carriers to give off their energy in the form of photons. A more plausible explanation maybe that the Mn impurities diffused into the p++ layer partially compensating the hole concentration resulting in a fermi level further away from the valence band. A decrease in the p-type doping would increase the tunneling barrier seen by carriers as and decrease the amount of tunneling current.

**Phosphor Test Bed**

**First Generation Phosphor Test Bed**

The analyses of several phosphors for use in a dynamic phosphor converted illumination source are reported here. Such a device can be constructed using readily available phosphors and pumped with standard GaN LED emission wavelengths. Pump wavelengths of 400 nm and 460 nm were chosen based on a number of criteria including availability of phosphors and desired output spectrum. The phosphors analyzed here include: UV to white (A), SrGa$_2$S$_4$ (B), SrCa:Eu (C). Each phosphor was excited by the 400 nm and 460 nm light independently and simultaneously and the output power spectra observed. Phosphor A was strongly excited by 400 and not 460, while the converse was true for Phosphor C as shown in Table 5.
Table 5. Summary of phosphor excitation data

<table>
<thead>
<tr>
<th>Excitation</th>
<th>Phosphors</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SrGa$_2$S$_4$ (Green)</td>
</tr>
<tr>
<td>460 nm</td>
<td>Strong</td>
</tr>
<tr>
<td>400 nm</td>
<td>Strong</td>
</tr>
</tbody>
</table>

Phosphors A, B and C were then combined based on the results in Table 5 to achieve a source with CCT similar to an incandescent bulb. Next, the dual wavelength source was used to excite the phosphors as the relative intensities of the two wavelengths were varied from singular 460 nm or 400 nm pumping to equal pumping from both wavelengths. Figure 17 a and b shows the change in phosphorescence as the intensity of one wavelength is held constant and the other varied for both cases.

The shape of the emission curve is shown to change by varying the relative intensity of the two pump wavelengths. Figure 17 shows an increase in the red phosphorescence (~655 nm peak) as the 460 nm pump relative intensity is increased (red curve). In addition, Figure 17 shows that an increase in 400 nm relative intensity is accompanied by an increase in the yellow/green (~560 nm peak) phosphorescence (green curve). Finally, the black curve represents equal pumping intensity by both pumps.

Figure 18 shows the varied spectra plotted on a 1931 Commission Internationale De L'eclairage (CIE) diagram showing the variation of chromaticity achievable with this combination of phosphors.
Pumping relatively harder with the 400 nm light excites more of the yellow/green phosphor emission moving the chromaticity coordinates closer to the yellow edge of the spectrum. Increasing the relative intensity of the 460 nm pump excites more of the red phosphorescence as well as adding blue light from the pump to the spectrum, thus, shifting the chromaticity coordinates toward the violet edge (bottom left) of the CIE diagram. A large range of points can be accessed with this combination of phosphors and pump choices. In addition, correlated color temperatures (CCT) ranging from 3588 K to 4793 K are achievable. Another important note is the high color rendering index (CRI) of 85 and 82 as listed in Figure 18. These CRIs rival many fluorescent lamps and are attributed to the broad emission from the combined phosphor emission.

These results show great potential for BSDLEDs. Such a source is possible by using a three terminal device as described previously and phosphor combinations similar to those analyzed here. Innovations such as this will help solid state illumination sources gain a competitive advantage over conventional illumination sources. This will be necessary for solid state lighting to realize success in the general illumination market.

**Second Generation Phosphor Test Bed**

YAG responds very well to pumping with 460-470 nm light, but not to ~400 nm light. On the other hand, the orange phosphor used responded equally well to pumping from 400-480 nm light. The range of chromaticity achievable with the second generation phosphor combination is shown in Figure 19.
This was achieved using the two terminal dual LED described later. When $\beta$ of the source is 0.08, the spectrum has a CCT of 3228K, and is very similar to an incandescent light bulb. As $\beta$ is increased by increasing current density, the spectrum shifts due to added long wavelength content from the LED as well as additional long wavelength components from the red and orange phosphors. The high CCT end also shows a slight upward curve that is caused by saturation of the long wavelength phosphors and more LED light passing through. Shift in the CIE coordinates for the standalone pump source is shown on the bottom left corner of Figure 19. The fact that the direction of shift for the final device and standalone pump differs illustrates that the phosphor emission significantly contributes to the final SPD.

**Dual LEDs**

**First Generation Two Terminal LED Structures**

*Fabrication*

Following the growth of the dual LED material, several standard semiconductor processing techniques were employed to fabricate individual devices. These included rapid thermal annealing (RTA), photolithography, reactive ion etching (RIE), and electron beam lithography. This section will step through the fabrication process used in this work for LEDs and other diode devices. Some discussion of the reasons and issues for each step will be included.

*Solvent Cleaning Routine:* A LED wafer is removed from the MOCVD growth chamber, and is either fabricated whole or cut into pieces for device fabrication.
The sample is then cleaned using a solvent cleaning routine. The sample is sprayed with Trichloroethane (TCE) first to remove any organic matter that may have deposited on the sample. Acetone is then used to remove the TCE and any residual contaminants. Methanol is then sprayed on the sample to remove the Acetone and other residual contaminants. The sample is then thoroughly washed with deionized water to remove any methanol and other contaminants. Finally, the sample is thoroughly dried with a nitrogen gun.

**Anneal - pGaN Activation**: The sample is annealed in the RTA system at 800°C for 4 minutes under N\textsubscript{2} ambient. This steps serves to drive hydrogen out of the pGaN layer on top of the device thereby activating the Mg dopants as acceptors in the lattice.

**Metal Deposition - Semitransparent Contact**: A semitransparent metal layer is deposited next. 5 nm of Ni followed by 5 nm of Au are deposited directly on the annealed sample. This metal layer serves as a current spreading layer that allows light to pass. In the next step, it will be made to help form an ohmic contact to the pGaN surface.

**Anneal - Current Spreading Contact**: Annealing the previous Ni/Au layer in an oxygen ambient causes oxygen to diffuse into the metal layer and form NiO at the metal semiconductor interface which aids in making an ohmic contact to the pGaN surface. The sample is annealed in the RTA at 500°C for 2 min in standard atmosphere which contains oxygen.

**Pattern transfer – Mesa**: Photolithography is used next to transfer a mesa pattern to the sample that will define the semitransparent Ni/Au pattern as well. Photoresist is spun on the sample at 2500 rpm for 30 seconds, followed by a second photoresist layer at the same parameters. The photoresist layer is then soft-baked at 100°C for 1.5 minutes, then exposed with the mesa pattern mask shown in Figure 27 for a total of 60 mJ/cm\textsuperscript{2} of photonic energy (for example 10 seconds of light with intensity of 6 mW/cm\textsuperscript{2}). A crosslinking-bake is then performed at 115°C for 2.5 minutes. A flood exposure is then performed without a mask for a total of >180 mJ/cm\textsuperscript{2}. Finally, the photoresist is developed in a solution of 5:1 H2O:AZ 400K for 1 minute.

**Metal Etch – Semitransparent**: The sample is then submerged in potassium iodide which removes any metal not covered with photoresist. Potassium iodide is typically used as a gold etchant, but is used here to etch both the Ni and Au.

**Hard-bake**: A hard-bake in a laboratory oven at 120°C for 30 minutes is required before the sample can be dry etched. The photoresist layer must be cured in the oven to prevent its etching in the dry etch process. This will insure that the material underneath the photoresist remains intact.
Dry Etch – Mesa: Reactive ion etching of the sample is performed to define the mesa structure of the device and expose the n-type GaN layer for contacting. A typical etch depth for an LED device in this work is 1.2 \( \mu \text{m} \).

Photoresist Stripping: Reactive ion etching causes some damage to the photoresist increasing the difficulty of totally removing the photoresist layer. However, an ultrasonic bath of acetone for 15 minutes removes all residue of the photoresist. Methanol is then used to remove and acetone residue, and de-ionized water is used to remove any methanol. Drying with a nitrogen gun is the final setup before the next patterning step is performed.

Pattern Transfer - pGaN Bonding Pad: The pattern transfer for the pGaN bonding pad is very similar to the mesa pattern transfer, except only on layer of photoresist is deposited, and the soft-baking and crosslinking-baking times are reduced to 1 minute and 2 minutes respectively. The mask and sample patterns must be aligned so that the pGaN bonding pad is at the proper place relative to the mesa for each segment of the pattern. Metal Deposition - pGaN Bonding Pad Electron beam evaporation of 50 nm of Ni followed by 250 nm of Au creates the metal layer for the pGaN bonding pad. The thickness of the Au layer determines how well a wire bonder will be able to bond a lead wire. If the Au is too thin it will peel off with the wire bonding process.

Metal Liftoff - pGaN Bonding Pad: Metal deposition covers the entire sample including the areas covered by photoresist. An ultrasonic acetone bath strips away the underlying photoresist and any metal that is covering it leaving metal only on the areas that were not covered by photoresist. This process is called metal lift-off because the metal is lifted away with the underlying photoresist leaving only the desired pattern.

Pattern Transfer - nGaN Bonding Pad: The pattern transfer for the nGaN bonding pad is exactly the same as the pGaN bonding pad, except that the nGaN bonding pad mask is used instead of the pGaN mask.

Metal Deposition - nGaN Bonding Pad: Electron beam evaporation is used to deposit 10 nm Ti, 50 nm Al, 10 nm Ti and 250 nm Au in that order. The gold capping layer must be thick enough to allow wire bonding.

Metal Liftoff - nGaN Bonding Pad: The metal lift-off for the nGaN bonding pad is exactly the same as for the pGaN bonding pad.

Figure 20 shows all the photomasks used for patterning the device overlayed. Figure 21 shows a completed device under electroluminescence. As can be seen from Figure 21, the devices are 350 \( \mu \text{m} \) squares, which give a device area of 1.225 \( \times 10^{-3} \text{ cm}^2 \).
Characterization

A strong quantum well emission dominated the PL of all samples. This emission was comparable in intensity to single III-N LEDs produced in the same facilities. Temperature dependent PL data for a blue emitter is shown in Figure 22. Inset shows peak position and intensity as a function of the temperature facilitating the determination of activation energies of 4 meV and 44 meV.
Similar results were obtained for the UV emitter (not shown here). A significant S-shape behavior of peak energy, which indicates the presence of QD-like potential fluctuations, is revealed by temperature dependent PL measurements for low excitation energies (Figure 22). Evaluation of MQW emission determines the activation energies to be $E_{\text{act}}(1) = 4 \text{ meV}$ and $E_{\text{act}}(2) = 44 \text{ meV}$ following the approach suggested by Adelmann et al.[15], Equation 3.

$$I = C \left[ 1 + A e^{E_1/T} + B e^{-E_2/T} \right]^{-1}$$

(3)

where, $k$ is the Boltzmann factor, and $A$ and $B$ are scaling factors. The scaling factor of the latter process is more than 200 times larger than the first. Interface roughness and/or one monolayer fluctuations of the QW thickness typically provide localization centers with localization energies below 5 meV. Therefore, the smaller energy can be attributed to imperfections of the interfaces in the MQW. A more thermally stable localization is indicated by the second $E_{\text{act}} = 44 \text{ meV}$. It is assigned to the localization of carriers in nanoscale islands caused by fluctuations in the indium concentration. Site selective PL spectroscopy was also performed on the dual MQW device, shown in Figure 23.
For each peak a mobility edge can be determined: 2.94 for the blue emission and 3.08 for the UV emission. This indicates the existence of isolated localization sites on the low-energy side of the mobility edges, confirming the radiative recombination in the device is generated in zero dimensional centers. Hence, the dual MQW devices show the same emission mechanism as detected for each single MQW devices.

Bright electroluminescence was also observed from the devices. The EL data for dual MQW region device is shown in Figure 24.
Two distinct peaks are observed in the dual MQW region device, showing separate luminescence from each of the MQWs. Similar results are observed for devices with 400 nm and 460 nm peaks.

The enabling light emission mechanism in GaN-based LEDs with a single MQW region, namely carrier localization [16], has been successfully employed in a device with dual MQW regions tailored to emit in the blue and near UV regions of the spectrum. This was important to preserve the same high brightness characteristics from standard GaN LEDs to the new device. Temperature and excitation energy dependent PL measurements confirmed the existence of carrier localization first in the single MQW region devices and then in the dual MQW region devices.

Bright emission derived from the carrier localization is also seen in EL (Figure 24). These devices show that dual wavelength emission is possible from the same device. Figure 25 shows the I-V curve for the device.

![I-V curve for a dual LED](image)

**Second Generation Two Terminal Dual Wavelength LEDs**

First a sample with no doping in any of the barriers was grown, fabricated and tested. The electroluminescent spectra at varied drive current is shown in Figure 26.
Figure 26 shows that at low currents the long wavelength emission is greater than the short wavelength emission. As drive current is increased the relative intensity of the short wavelength emission is increased to a point where the effect saturates. The maximum $\beta$ for this device is 1.47 and the minimum is 0.89. Next the barrier between QWs 3 and 4 was doped with Si to study the affect it has on the minimum and maximum $\beta$. Three samples with different doping levels in the barrier between QWs 3 and 4 were grown, fabricated and tested. The first sample, had a carrier concentration of $6\times 10^{18} \text{ cm}^{-3}$ electrons. EL spectra in Figure 27 show that at low currents the short wavelength dominates with a $\beta=0.1$. 

![Normalized EL spectra of Dual LED with undoped barriers](image-url)
It is not until very high current densities (220 A/cm$^2$) that $\beta$ begins to change. The maximum $\beta$ reached before the device is destroyed by high current is ~1.1, as shown in Figure 28, where $\beta$ is plotted against current density.
Another sample was grown with twice the Si doping in the barrier (1.2x10^{19} \text{ cm}^{-3}), however, no change in $\beta$ was observed for current densities that the device could sustain. A fourth sample with half the Si doping (3x10^{18} \text{ cm}^{-3}) was also grown. The effect of drive current on $\beta$ was much greater and started to show at current densities 100 A/cm^2, as seen in Figure 29.

![Figure 29: $\beta$ vs J for 3x10^{18} \text{ cm}^{-3} doped barrier](image)

The maximum $\beta$ attainable for this device was ~2.5, for which the EL spectrum is shown in Figure . The results for these devices are tabulated in Table 6.

<table>
<thead>
<tr>
<th>Doping (cm^{-3})</th>
<th>Min $\beta$</th>
<th>Max $\beta$</th>
<th>$\beta=1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>0.89</td>
<td>1.47</td>
<td>$4\times10^{-4}$</td>
</tr>
<tr>
<td>$3 \times 10^{18}$</td>
<td>0.08</td>
<td>2.5</td>
<td>$220 \text{ A/m}^2$</td>
</tr>
<tr>
<td>$6 \times 10^{18}$</td>
<td>0.08</td>
<td>1.1</td>
<td>$305 \text{ A/m}^2$</td>
</tr>
<tr>
<td>$1.2 \times 10^{19}$</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>
Discussion of Dual LED

Doping of the barrier between QWs 3 and 4 with Si achieves significant control over the $\beta$ by varying drive current. A model to explain this behavior considers two barriers to current injection in the different QW regions. A band diagram for the QW region of a device with doping in the barrier between QWs 3 and 4 is shown in Figure 30.
In Figure 31 the n-type region lies to the left of the QWs and the p-type region is to the right. The relatively high In concentration in the QWs 1-3 leads to electron trapping and blocking, making it more difficult for electrons to travel to QWs 4-6. This effect is best observed in a dual LED with no doping where the QWs 4-6 are preferentially excited at low currents. In the case of no doping holes are more free to move to QWs 1-3 than electrons are to move to QWs 4-6 at low currents. As current is increased the electrons fill higher energy states and are able to travel to QWs 4-6. At this point the mobility of the carrier influences $\beta$. QWs 4-6 are more preferentially pumped because holes have lower mobilities and do not travel as far as the electrons, thus recombination is more likely in QWs 4-6. On the other hand, the doping of the barrier between QWs 3 and 4 creates a barrier to hole transport to the QWs 1-3. In the case of the doped barrier the hole blocking washes out the effect of electron blocking at low currents. The holes are trapped in QWs 4-6 until current is high enough to promote some holes over the doped barrier and into QWs 1-3. At this point the electron trapping effect influences $\beta$ recombination occurs in QWs 1-3.

**Conclusion**

Several methods for forming a III-nitride tunnel junction were explored, however, a suitable device for a buried current spreading layer was not achieved. Hybrid doping showed the best results as a tunnel junction however the voltage need to achieve reverse currents suitable for LED operation were too high. SPS doping showed strong potential for obtaining high concentrations of holes in p-type material, however, structures incorporating these layers did not create suitable tunnel junctions. Lastly, Mn doping of an intrinsic region in the junction to create defect assisted tunneling also proved to be not suitable for creating a tunnel junction. Though ultimately unsuccessful in the development of a suitable tunnel junction, this doping study is the first systematic effort.
to examine stand-alone tunnel junction structures on GaN-based materials. It is also the first to investigate SPS structures as buried current spreading layers.

The results shown here describe successful use of phosphor combinations with a dual LED to create a dynamic spectrum light source. The first generation phosphor test bed was pumped with a source in which $\beta$ ranged from 0 to $\infty$. However, the second generation phosphor test bed had a constraint that it must work with a $\beta$ range of 0.08 to 2.5. A suitable set of phosphors was used to create a source of which the CCT can be varied from 3228 K to 5339 K by increasing the drive current used in the LED.

A spectrally dynamic broadband source was developed with ability to tune CCT from 3200K to 5300K. The novel use of multiple phosphors which can be selectively excited by two wavelengths of light enabled this development. A two terminal dual LED has been created with the ability to control the relative intensities of the two emission peaks by varying drive current. The development of a three terminal dual LED as a pump source was prohibited by the need for a III-nitride tunnel junction that proved unattainable in the scope of this work. Doping profiles have been used to extend the dynamic range of the two terminal dual LED over other reported devices. Operation of the two terminal dual LEDs is explained as a function of drive current. Combinations of phosphors that have varied excitation spectra provide the ability to selectively excite different phosphors with the different LED emission peaks. First and second generations of the two terminal dual LED and the phosphor combination are discussed. Such innovations may prove important in the success of solid state lighting as a general illumination solution.

The current injection mechanisms for the two terminal dual wavelength LED are not fundamentally different from those of conventional high brightness LEDs. Thus, it is expected that the efficiency of optimized two terminal dual wavelength LED would be near that of other high brightness LEDs. Common enhancement techniques such as chip shaping and flip-chip bonding can be applied to the dual wavelength device as easily as any conventional high brightness LED. Furthermore, if suitable phosphors and pump wavelengths are selected, efficiency can be enhanced by minimizing the stokes shift loss seen in other phosphor converted white LEDs.

The broadband spectrally dynamic source can be viewed as an enhancement of more conventional LEDs and should exhibit many of the same properties as other high brightness LEDs. This will in part depend on the optimization of drive current control over SPD. With further investigation it is believed that this could be achieved. However, the two terminal dual emitting device developed in this project is not as versatile as the three terminal tunnel junction device investigated in this project, which would allow independent control of both of the component LEDs. Such devices with blue and green emitters have been reported, but these were not designed for use with phosphors [17, 18]. The tunnel junctions in these devices produced an increase in the necessary forward voltage for LED operation of the bottom device, as well as an increase in equivalent series resistance. Further improvement of III-N tunnel junction performance is critical for efficient three terminal devices, which have the potential to be viable solid state lighting sources.
References


[2] The promise of solid-state lighting for general illumination: LEDs and OLEDs’, Technology roadmaps co-sponsored by DOE (BTS) and OIDA, published by OIDA (oida.org)


Technical Papers Resulting from this Work


