EFFICIENT LIGHT EXTRACTION METHOD AND DEVICE

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Abstract
A tight emitting device comprises at least one p-type layer and at least one n-type layer and a microlens array surface. A method for improving light efficiency of a light emitting device, comprises depositing polystyrene microspheres by rapid convection deposition on surface of light emitting device; depositing a monolayer of close-packed SiO₂ microspheres onto the polystyrene microspheres; and seal treating to convert the polystyrene microspheres into a planar micro-layer film to provide a surface comprising substantially two-dimensional (2D) hexagonal close-packed SiO₂ colloidal microsphere crystals partially imposed into a polystyrene monolayer film.

14 Claims, 5 Drawing Sheets
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<th>Publication Number</th>
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<tr>
<td>6,958,497 B2</td>
<td>10/2005</td>
<td>Emerson et al.</td>
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<tr>
<td>2003/0020085 A1</td>
<td>1/2003</td>
<td>Bour et al.</td>
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<tr>
<td>2006/0017061 A1</td>
<td>1/2006</td>
<td>Sakamoto et al.</td>
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**OTHER PUBLICATIONS**


Deposition of close-packed monolayer PS microspheres via rapid convection deposition (112)

Deposition of close-packed monolayer SiO₂ microspheres (114)

Polystyrene microspheres converted to planar layer via heat treatment ~ 140°C (116)
In$_{0.19}$Ga$_{0.81}$N QW LEDs
$\lambda_{\text{peak}}$=480-nm

Fig. 4
PL with HeCd Laser at 325 nm T=300 K

Fig. 5
EFFICIENT LIGHT EXTRACTION METHOD AND DEVICE

CROSS REFERENCE TO RELATED APPLICATION

This application claims the benefit of U.S. Provisional Application No. 60/871,823, filed 24 Dec. 2006, which is incorporated herein by reference. This invention was made with government support under Contracts Nos. W9111NF-07-2-0064 and 07014121 respectively awarded by the Department of Defense—Army Research Lab and by the National Science Foundation. The government has certain rights in the invention.

BACKGROUND OF THE INVENTION

The invention relates to a light emitting device, particularly to an improved light extraction efficiency method and light emitting diode (LED). A light-emitting diode is a semiconductor diode that emits incoherent narrow-spectrum light when electrically biased in the forward direction of a p-n junction. This effect is a form of electroluminescence.

An LED typically comprises a small area source, often with extra optics added to the chip that shapes its radiation pattern. Color of emitted light depends on semiconductor material composition and can be infrared, visible, or near-ultraviolet. The LED can comprise a chip of semiconducting material impregnated or doped with impurities to create the p-n junction. Charge-carriers—electrons and holes—flow into the junction from electrodes with different voltages. When an electron meets a hole, it falls into a lower energy level and releases energy in the form of a photon (light) causing current flow from the p-side, or anode, to the n-side, or cathode. The wavelength of the light emitted, and hence its color, depends on the band gap energy of the materials forming the p-n junction.

Refractive index of an LED package material should match the index of the semiconductor, otherwise produced light will be partially reflected back into the semiconductor, where it may be absorbed and turned into efficiency lowering heat. Efficiency lowering reflection also occurs at the surface of the package if the LED is coupled to a medium with a different refractive index such as a glass fiber or air. The refractive index of most LED semiconductors is quite high, so in almost all cases the LED is coupled into a much lower-index medium. The large index difference makes the reflection quite substantial (per the Fresnel coefficients), and both the Fresnel reflection and critical angle limitations are usually the dominant causes of LED inefficiency. Often more than half of emitted light is reflected back at the LED-package and package-air interfaces.

Several approaches have been implemented to improve light extraction efficiency of LEDs, such as: surface roughening, photonic crystals and nano-pyramids. The disadvantages of surface roughening are related to difficulty in controlling the process as well as in achieving good roughness repeatability. The photonic crystal and nanoplyramid approaches require costly e-beam lithography, which are not applicable for large scale production of nitride LEDs.

A need continues for enhanced, inexpensive and repeatable LED light extraction efficiency.

BRIEF DESCRIPTION OF THE INVENTION

The invention provides enhanced and repeatable light emitting device extraction efficiency at a reasonable cost. The invention is described in an embodiment as a light emitting device comprising at least one p-type layer and at least one n-type layer and a microsphere array surface.

Also, the invention is a method for improving light efficiency of a light emitting device, comprising: depositing polystyrene microspheres by rapid convection deposition onto a quantum well; depositing a monolayer of close-packed SiO₂ microspheres onto the polystyrene microspheres; and heat treating to convert the polystyrene microspheres into a planar multilayer film to provide a surface comprising substantially two-dimensional hexagonal close-packed SiO₂ colloidal microsphere crystals partially imaged onto a polystyrene monolayer film.

Another embodiment is an optoelectronic device comprising: a multilayer semiconductor structure comprising a GaN layer and an active region, the active region comprising at least one quantum well layer of InGaN and GaN, wherein the electron quantum well layers and hole quantum well layer form a first quantum well stage, and wherein the active region comprises a plurality of quantum well stages adjacent to each other having the same structure as the first quantum well stage; and a microsphere array surface on the multilayer semiconductor.

In another embodiment, the invention is a method for making an optoelectronic device, comprising: providing a multilayer semiconductor structure comprising a GaN layer and an active region, the active region comprising at least one quantum well layer of InGaN and GaN, wherein the electron quantum well layers and hole quantum well layer form a first quantum well stage, and wherein the active region comprises a plurality of quantum well stages adjacent to each other having the same structure as the first quantum well stage; and depositing a microsphere array surface onto the multilayer semiconductor.

Still another embodiment is a method for generating optical emission from an optoelectronic device, comprising: providing a GaN layer and an active region, the active region comprising at least one quantum well layer of InGaN and GaN, wherein the electron quantum well layers and hole quantum well layer form a first quantum well stage, and wherein the active region comprises a plurality of quantum well stages adjacent to each other having the same structure as the first quantum well stage; depositing a microsphere array surface onto the GaN layer and an active region; and exciting the active region to produce optical emission.

BRIEF DESCRIPTION OF THE DRAWING

FIG. 1 is a schematic representation of an LED structure; FIG. 2 is a process flow schematic; FIG. 3A is a confocal laser scanning microscopy image and FIG. 3B is a scanning electron microscopy image; FIG. 4 is a graph of photoluminescence (PL); and FIG. 5 is a graph showing a power and current comparison.

DETAILED DESCRIPTION OF THE INVENTION

LED light extraction efficiency is the portion of emitted electromagnetic radiation that is transmitted usable for human vision. It is a ratio of emitted luminous flux to radiant flux. The present invention relates to an LED with improved light extraction efficiency.

An LED can comprise a chip of semiconducting material impregnated or doped with impurities to create a p-n junction. Current flows from the p-side or anode, to the n-side or cathode, but not in the reverse direction. Charge carriers—electrons and holes—flow into the junction from electrodes
with different voltages. When an electron meets a hole, it falls into a lower energy level and releases energy in the form of a photon.

A quantum well is an LED potential well. The term “quantum well” or “QW” used herein refers to a thin-layer structure comprising alternate layers consisting of a first semiconductor layer with a thickness smaller than the de Broglie wavelength of about 200 Å to 300 Å with respect to electrons or holes and at least a second semiconductor layer with a band gap greater than that of the first semiconductor layer. A “substrate” is an underlying template or substratum can such as a sapphire template, GaN substrate, a Si substrate, SiC substrate or ZnO substrate.

A QW structure can be formed by sandwiching a semiconductor thin layer of a narrow band gap between semiconductor layers of a large band gap. If a single semiconductor thin layer constitutes a quantum well for both electrons and holes, the quantum well is called a type I quantum well. In this case, the semiconductor layer of a narrow band gap is called a well layer, and the semiconductor layers of a large band gap are called barrier layers. A type 1 multi-quantum well structure can be formed by alternately laminating semiconductor layers of narrow and broad band gaps. A type 11 quantum well structure has a first semiconductor layer forming a quantum well for electrons, a second semiconductor layer forming a quantum well for holes formed on the first semiconductor layer and third semiconductor layers sandwiching the first and second semiconductor layers as barrier layers to the electrons and holes. A type II multi-quantum well structure can be formed by alternately laminating first semiconductor layers, second semiconductor layers and third semiconductor layers.

An embodiment, the invention relates to an optoelectronic device that includes a GaN layer and a quantum well active region with a microsphere array surface. The active region can comprise at least an InGaN hole quantum well layer and electron quantum well layers adjacent to the hole quantum well layer. At least one of the electron quantum well layers is GaN. The electron quantum well layer and hole quantum well layer form a first quantum well stage. The active region can include a plurality of the quantum well stages adjacent to each other having the same structure as the first quantum well stage. The structure can include a transitional layer of GaN between each quantum well stage.

A QW structure can be grown by III-V semiconductor MOCVD/MBE epitaxy and molecular beam epitaxy (MBE). However, for manufacturing considerations such as high-throughput, the use of metal organic chemical vapor deposition (MOCVD) growth may be preferred.

Quantum efficiency ($\eta_{QW}$) of a QW depends on the injection efficiency ($\eta_{inj}$), radiative efficiency ($\eta_{rad}$), and light extraction efficiency ($\eta_{extraction}$). Low carrier mobility, carrier lifetime, and recombination losses in p-type doping and polarization-induced electric fields impact the injection efficiency and radiative efficiency particularly of an III-Nitride LED. Large refractive index contrast at a GaN/air interface results in low light extraction efficiency.

Surface roughening has been implemented to improve the light extraction efficiency of an InGaN QW LED. The disadvantage of surface roughening is related to control and repeatability difficulties. Photonic crystal and microprism approaches have been proposed to improve light extraction efficiency. However, these approaches require e-beam lithography to obtain closely controlled dimensions, but, e-beam lithography is not applicable for low-cost and large scale production.

The invention provides an applied surface texture that improves light extraction efficiency of an LED. In an embodiment, the invention relates to an InGaN QW-based LED structure that utilizes SiO$_2$ microspheres to provide significantly improved light extraction efficiency. The SiO$_2$ microspheres provide a low-cost and straight-forward improvement in light extraction efficiency that eliminates the need for costly e-beam lithography.

The SiO$_2$ microsphere surface texturing can be formed by depositing PS microspheres onto a light emitting surface of the LED, depositing a monolayer of the SiO$_2$ microspheres, and then heat treating to convert the PS to a planar microlayer. In one procedure, the PS microspheres are deposited by rapid convection deposition. The PS microspheres can be deposited in a suspension media such as water. The PS suspension can be a 5% to 25% volume fraction PS in a suitable suspending media such as water. Preferably the PS volume fraction is 8% to 12% or about 10%. Suitable PS microsphere arrays are available from NanoLab, Inc., 55 Chapel Street, Newton, Mass. 02458. The microspheres can be deposited in a temperature range of 10 to about 50°C, preferably 20 to about 30°C.

Then a monolayer of close-packed SiO$_2$ microspheres is deposited as a suspension onto the PS microspheres. The SiO$_2$ suspension can be a 5% to 25% volume fraction SiO$_2$ in a suitable suspending media such as water. Preferably the SiO$_2$ volume fraction is 10% to 15% or about 13%. Suitable size spheres can be in the range from 0.05 microns up to 4 microns in diameter.

Suitable SiO$_2$ microspheres can be prepared, for example, by hydrolysis of tetraethoxysilane or, by a sol gel process or by spray drying a colloidal silica gel. The microspheres can be deposited in a temperature range of 10 to about 50°C, preferably 20 to about 30°C. Suitable size spheres can be in the range from 0.05 microns up to 4 microns in diameter, preferably 0.1 micron to 2 microns. Suitable SiO$_2$ microspheres are available from Bangs Laboratories, Inc., 9025 Technology Drive, Fishers, Ind. 46038-2886 and are available in quantities with a standard size deviation of less than 1%.

Then the SiO$_2$/PS is heat treated to convert the polystyrene microspheres into a planar microlayer film to provide a surface comprising substantially two-dimensional hexagonal close-packed SiO$_2$ colloidal microsphere crystals partially imbedded into a polystyrene monolayer film. The heat treatment can be conducted at a temperature in the range of 120°C to 200°C, preferably 130°C to 150°C. The resulting film thickness can be in the range of 0.01 micron up to about 3 micron.

The invention may be embodied in various types of optoelectronic devices including amplifiers, light emitting diodes and edge emitting and surface emitting lasers that incorporate optical feedback to provide laser action. The invention may find application in solid state lighting, solid state displays, lasers, light emitting diodes (LEDs), biomedical therapy and diagnostic devices, medical lasers, eye surgery devices and DVD lasers.

The invention provides an InGaN quantum well LED structure utilizing a SiO$_2$/polystyrene (PS) microsphere array that enhances light extraction efficiency from a top surface of an LED. The dimensions of the SiO$_2$ microspheres can be from 0.1 μm to 2 μm, preferably from 0.25 μm to 1 μm. Microsphere size can be controlled accurately to provide roughening repeatability. The SiO$_2$ spheres exhibit good adhesion to III-V materials. The spheres are and transparent to light emission from InGaN LEDs. The deposition of substantially two dimensional close-packed SiO$_2$ colloidal crystals is simple and quick compared to electron-beam lithography or to complex wafer fabrication techniques. The SiO$_2$
microscopes can be coated to the LED top emission area to avoid detrimental influence on the I-V characteristics of the LED.

These and other features of the invention will become apparent from the drawings and following detailed discussion, which by way of example without limitation describe preferred embodiments of the invention.

EXAMPLES

In these EXAMPLES, a layer of polystyrene (PS) and a monolayer of two-dimensional hexagonal close-packed SiO2 colloidal crystal were deposited on a top surface of an InGaN QW-based device. FIG. 1 illustrates the LED device 10 including a 3-μm GaN template 14 (grown at 1080°C) on c-plane sapphire substrate 12 using a low-pressure vertical-type MOCVD system, employing a low temperature buffer layer of 30-nm GaN (grown at 535°C). Active region 16 of the device structure comprised 4-period 2.5-nm InGaN QWs 18 and 12-nm GaN barriers 20. The device 10 included p-GaN layer 22, n-contact 24 and p-contact 26 as shown.

The LED device 10 included a microlens array surface 28 comprising SiO2 microspheres 32 embedded in a polystyrene monolayer film 34 prepared in accordance with the process illustrated in FIG. 2. The SiO2 microspheres 32 with diameter of 1.0 μm were semi-buried in the PS film 34, thereby forming a close-packed lens-like array 28. The refractive index of GaN in the visible spectrum is 2.5, while the refractive indices of a PS film 34 and SiO2 microspheres 32 are 1.58 and 1.46, respectively. The array 28 on the top surface of the LED 10 allowed photons emanating from the QW to scatter out from the LED 14 structure with larger ‘effective’ photon escape cone, thus leading to increase in the device external quantum efficiency, as hereinafter described in detail.

FIG. 2 schematically represents a process 110 of depositing a PS film 34 and monolayer of two-dimensional hexagonal close-packed SiO2 colloidal crystals 32. In FIG. 2, polystyrene spheres 30 were deposited by rapid convection deposition 112. Then a monolayer of close-packed SiO2 microspheres 32 was deposited 114 onto the PS microspheres 30. Finally the PS microspheres 30 were converted 116 to a planar microcrystal film 34 via heat treatment at 140°C. The final surface comprised substantially two-dimensional hexagonal close-packed SiO2 microsphere crystals 32 partially impeded into a planar PS microcrystal film 34 as shown in 3A and 3B.

In the FIG. 2 process, volume fractions of a 1.0 μm-diameter PS suspension and a SiO2 microsphere suspension were 10% and 13%, respectively.

A droplet volume of a 10 μL PS colloidal suspension was injected onto the InGaN QW LED sample. The volume used was just sufficient to cover the sample surface, about 10 microliters. A leveler (a deposition glass plate in this EXAMPLE) was swiped across the surface at an angle of 25°±10° controlled by a linear motor at a speed of 45 μm/s. After depositing one ML of PS microspheres 30, the process was repeated to deposit one ML of SiO2 microspheres 32 onto the PS microspheres. The coated samples were then heated using a hotplate at 140°C to melt the PS microspheres 30, thereby capturing the SiO2 microspheres 32 in a planar PS film 34 without significant rearrangements to the SiO2 packing structure.

Photoluminescence (PL) and LED test samples were grown using a vertical-type metalorganic chemical vapor deposition (MOCVD) reactor. The PL samples were grown on a 3-μm u-GaN template on c-plane sapphire at a temperature of 1080°C, employing a low temperature 30-nm u-GaN buffer layer. The PL structure was then grown on top of the u-GaN template. Active regions consisted of four-period 2.5-nm InGaN QW and 12-nm GaN barriers, grown at a temperature of 710°C. The In-content of the InGaN QW of the PL sample was found as 12%, as calibrated via X-ray diffraction.

In-content of the investigated GaN QW 10 ranged from 12% up to 20%, calibrated via X-ray diffraction. The n-GaN layer 14 was Si-doped at a level of ~ 2x10^{19} cm^{-3}. The PL samples did not employ p-GaN 22. The p-GaN 22 was grown utilizing 50-nm Mg-doped GaN at growth temperature of 970°C, followed by N2 activation annealing at a temperature of 780°C for a duration of 5 min.

The ex-situ rapid convective deposition of the microsphere layers from colloidal suspensions was subsequently conducted on top of each of the InGaN QW PL and LED samples. Strong capillary forces at a meniscus between a substrate and a colloidal suspension induced crystallization of spheres into a 2D array. The use of colloidal self-assembly exploits the tendency of monosized sub-micrometer spheres to spontaneously arrange into a close-packed 2D crystal.

The SiO2 microspheres 32 were semi-buried in the PS film 34, forming a hexagonal close-packed SiO2/PS microlens array 28. FIG. 3A and FIG. 3B are confocal laser scanning microscopy and scanning electron microscopy images of the SiO2/PS microlens array 28 on top of the GaN layer of the LED structure. The scanning microscopy image shows substantially close packed SiO2 microspheres 32. As illustrated in FIG. 3B, the SiO2 colloidal crystal forms a 2D hexagonal closed-packed microlens array 28. The spherical SiO2 microspheres 32 had diameters of 0.8-1 μm and were close-packed and semi-buried in the PS. Substantially close packed means that the microspheres were arranged as a lattice of spheres that substantially took up the greatest possible fraction of a 2-dimensional plane. The close-packed SiO2 microspheres 32 imparted a controlled surface roughness that allowed light rays emanating from the QW to diffuse out from the LED 10 thus leading to increased external quantum efficiency.

PL comparison studies were conducted on In0.15Ga0.85N QW samples emitting with peak wavelength (λpeak) at 419.3 nm. The PL measurements were conducted on samples (grown at the same time) with and without the microlens array 28, utilizing an He-Cd excitation laser (λ=325 nm) from the backside of the samples at room temperature. PL luminescence was collected from the top surface of the samples.

FIG. 4 shows the PL spectra of the In0.15Ga0.85N QW PL samples with and without the microlens array. PL luminescence peak intensity of the samples covered with microlens array 28 exhibited 233.6% improvement over that of samples without microlens array. Integrated PL luminescence for samples with the microarray 28 also showed 269.7% improvement over that of an uncoated sample. The multi-peaked emission of the PL spectra for the uncoated sample is an artifact of the Fabry-Perot cavity effect in the cavity formed by sapphire/GaN/air, resulting in interference effect. The microlens array 28 on the surface of the sample scattered the emitted photons from the active media, resulting in suppression of the Fabry-Perot cavity effect.

The SiO2/PS microlens array was deposited on an LED sample, employing 4-period of 2.2 nm±0.15 nm thick (calibrated via transmission electron microscopy) In0.15Ga0.85N QWs with GaN barriers, emitting with λpeak=480 nm. In-content of the InGaN QW of the LED sample was found as 19%±1%, as calibrated via X-ray diffraction. The LED structure was grown on a 3.0-μm n-GaN template on c-plane sapphire substrate. The n-GaN was Si-doped with n-doping of 2x10^{19} cm^{-3}. The p-GaN was grown utilizing 80-nm
thick Mg-doped GaN (p-doping=3×10^{17} \text{ cm}^{-3}) at growth temperature of 970°C, followed by N_{2} annealing at a temperature of 780°C for a duration of 5 minutes.

Continuous wave (CW) power measurements were performed at room temperature for LED devices with and without the SiO_{2}/PS microspheres array 28. On-wafer power measurement was done in a light proof dark chamber, using a large-area planar-diffused silicon photodiode, and a benchtop optical power meter. FIG. 5 shows the output power as a function of the driving current for up to 100 mA for both LEDs with an area of 1 mm². The CW power-current measurements exhibit 219% improvement in the output power of the LED device with SiO_{2}/PS microspheres array 28 at a current level of 100 mA, as compared to that without microspheres array.

The current invention provides a significant increase in LED output power. While there is no intention to be bound by the following explanation, it is believed that the increase in the output power for devices with microspheres is attributable to an increase in an effective photon escape cone between emitter and air. Escape cone is a virtual cone defined by a critical angle (θc) of impact of a photon at an interface. The critical angle defines a limit to the escape of incident photons from an emitter. Photons that are incident to the interface at angle θc or less, escape from the emitter while photons that are incident at a greater than θc angle are reflected back into the emitter where they may be lost by absorption. See Pocius et al., 6987613, incorporated herein by reference in its entirety. The inventive SiO_{2}/PS film serves as an intermediate refractive index material that provides multiple angles of incidence. The multiple angles of incidence of the curved hemisphere SiO_{2} surfaces provide increased opportunities (as compared to a flat surface) for photon escape, thereby expanding the effective escape cone between emitter and air.

In addition to increase in effective photon escape cone, the use of SiO_{2} (n=1.46)/PS (n=1.58) as the intermediate refractive index materials for the microspheres array also leads to reduced Fresnel reflection in the GaN/PS/SiO_{2}/air interface by as high as 4.7% for normal incidence, as compared to that for GaN/air interface. The Fresnel reflection for GaN/PS/ SiO_{2}/air interface was calculated using transfer matrix propagation matrix method for normal incidence.

Relative external quantum efficiency as a function of injection current of the LED was obtained by differentiating the relative output power with the injection current. As shown in FIG. 5, there was an overall improvement in the relative external quantum efficiency of LEDs with SiO_{2}/PS microspheres array 28 as compared to that of the LEDs without the microspheres array. At low current level (1-5 mA), the improvement of LED efficiency with microspheres was about 4.3 times. The improvement in the relative external quantum efficiency in of 3.32 times was observed for LEDs with microspheres at current level of 100 mA. This reduction in the improvement in LEDs with microspheres at high current level can be attributed to the escape effect. The proof-of-concept experiments show promising results, and further understanding and optimization on the thermal distribution of the LEDs with microspheres array are still required.

The EXAMPLES demonstrate enhancement of light extraction efficiency of InGaLN QW LEDs by using SiO_{2}/PS microspheres. The utilization of SiO_{2} microspheres led to enhancement of ~3.3 and ~4.3 times the peak luminescence intensity and the integrated luminescence of the InGaLN QW LEDs. Improvement of output power by ~23% for the LEDs with coated microspheres was also observed. This low-cost and controllable microsphere process provides a method for enhancing the light extraction LED efficiency.

Utilization of SiO_{2}/PS microspheres deposited via rapid convective deposition on InGaN QW PL sample led to improvement of 233.6% and 269.7% for its peak luminescence intensity (\lambda_{peak}≈493 nm) and integrated luminescence, respectively. Improvement of output power by 219% (at current level of 100 mA) for the electrically-injected InGaN QW LEDs emitting with \lambda_{peak}≈490 nm with microsphere array was also obtained, presumably due to the increase in the effective photon escape cone and reduced Fresnel reflection.

The process of forming the array provides adequate adhesion to III-Nitride materials at room temperature. The array materials are relatively transparent to visible light emission from the InGaN QW. Also, the deposition of 2D close-packed SiO_{2}/PS colloidal crystal is practical and straightforward, as compared to e-beam lithography or complex wafer fabrication techniques. As the SiO_{2} microspheres are deposited as a final step on the top emission area of the LEDs, this approach avoids any degradation on the electrical characteristics of the LEDs.

While preferred embodiments of the invention have been described, the present invention is capable of variation and modification and therefore should not be limited to the precise details of the Examples. The invention includes changes and alterations that fall within the purview of the following claims.

What is claimed is:

1. A method for improving light efficiency of a light emitting device, comprising:
   - depositing polystyrene microspheres by rapid convection deposition on the top surface of light emitting device;
   - depositing a monolayer of close-packed SiO_{2} microspheres onto the polystyrene microspheres; and
   - heat treating to convert the polystyrene microspheres into a planar microsphere film to provide a surface comprising substantially two-dimensional hexagonal close-packed SiO_{2} colloidal microsphere crystals partially imaged into a polystyrene monolayer film.

2. The method of claim 1, comprising providing a semiconductor substrate; and forming on the substrate a succession of layers to provide a QW comprising at least one InGaN layer and at least one InGaN layer adjacent a GaN barrier layer; and depositing the polystyrene microspheres onto a structure comprising the formed QW.

3. The method of claim 1, comprising forming a SiO_{2} microsphere suspension.

4. The method of claim 1, comprising:
   - forming a 1.0 μm-diameter polystyrene suspension; and
   - depositing a droplet volume of the polystyrene suspension between the InGaN QW LED sample; and
   - sweeping a deposition plate at an angle of 25°±1° to the deposited droplet volume of polystyrene to form the film.

5. The method of claim 1, comprising forming on the substrate a succession of layers to form an inner contact layer and a multiple quantum well stack comprising the active region, and an outer contact layer.

6. The method of claim 1, wherein the heat treating is at a temperature in the range 120° to 200°C.

7. The method of claim 1, wherein the heat treating is at a temperature in the range 130° to 150°C.

8. The method of claim 1, wherein the close-packed SiO_{2} microspheres are deposited as a 5% to 25% volume fraction suspension.

9. The method of claim 1, wherein the close-packed SiO_{2} microspheres are deposited as a 10% to 15% volume fraction suspension.
10. The method of claim 1, wherein polystyrene microspheres are deposited as a 5% up to 25% volume fraction suspension.

11. The method of claim 1, wherein polystyrene microspheres are deposited as a 8% to 12% volume fraction suspension.

12. The product of the process of claim 1.

13. A method for making an optoelectronic device, comprising:

providing a multilayer semiconductor structure comprising a GaN layer and an active region, the active region comprising at least one quantum well layer of InGaN and GaN, wherein the electron quantum well layers and hole quantum well layers form a first quantum well stage, and wherein the active region comprises a plurality of quantum well stages adjacent to each other having the same structure as the first quantum well stage; and
depositing a polystyrene microsphere array surface onto the multilayer semiconductor structure; depositing close-packed SiO₂ microspheres onto the polystyrene microsphere array surface; and

heat treating to convert the polystyrene microsphere array surface into a microlayer film to provide a surface comprising close-packed SiO₂ colloidal microsphere crystals partially imposed into a polystyrene monolayer film.

14. A method for generating optical emission from an optoelectronic device, comprising:

providing a GaN layer and an active region, the active region comprising at least one quantum well layer of InGaN and GaN, wherein the electron quantum well layers and hole quantum well layer form a first quantum well stage, and wherein the active region comprises a plurality of quantum well stages adjacent to each other having the same structure as the first quantum well stage;
depositing a polystyrene microsphere array surface onto the active region;
depositing close-packed SiO₂ microspheres onto the polystyrene microsphere array surface;
heat treating to convert the polystyrene microsphere array surface into a microlayer film to provide a surface comprising close-packed SiO₂ colloidal microsphere crystals partially imposed into a polystyrene monolayer film; and
exciting the active region to produce optical emission.