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White light-emitting diodes of GaN-based Sr$_2$SiO$_4$:Eu and the luminescent properties

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We have synthesized an Eu$^{2+}$-activated Sr$_2$SiO$_4$ yellow phosphor and investigated an attempt to develop white light-emitting diodes (LEDs) by combining it with a GaN blue LED chip. Two distinct emission bands from the GaN-based LED and the Sr$_2$SiO$_4$:Eu phosphor are clearly observed at 400 nm and at around 550 nm, respectively. These two emission bands combine to give a spectrum that appears white to the naked eye. Our results showed that GaN (400-nm chip)-based Sr$_2$SiO$_4$:Eu exhibits a better luminous efficiency than that of the industrially available product InGaN (460-nm chip)-based YAG:Ce.

The white light-emitting diodes (LEDs) composed of InGaN blue LEDs and a yellow YAG:Ce phosphor have been investigated extensively due to their applications, such as back-lighting for liquid crystal displays and incandescent lamps. These applications are particularly useful for saving electric power owing to the lower energy consumption of LEDs. In addition, there have been some detailed studies on the integration of the blue InGaN LED and the yellow YAG phosphor, as this creates white light from a combination of a blue LED emission (460 nm) and a broadband yellow YAG phosphor. However, very efficient new yellow phosphors have actually been discovered, besides the YAG:Ce phosphor and the organic luminescent materials.

In the present work, we have synthesized an Eu$^{2+}$-activated Sr$_2$SiO$_4$ yellow phosphor and investigated in an attempt to develop white LEDs through the integration of the GaN blue LED chip ($\lambda_{em} = 400$ nm) and the Sr$_2$SiO$_4$:Eu phosphor into a single package. As shown in this study, the GaN (400-nm chip)-based Sr$_2$SiO$_4$:Eu white LED exhibits higher luminous efficiency compared with that of the commercially available InGaN (460-nm chip)-based YAG:Ce white LED.

In order to successfully synthesize exact stoichiometric compounds with fine particles and low agglomeration particles, a polymeric complex method using citric acid and ethylene glycol was used. The starting materials used in the polymeric complex method were Sr(NO$_3$)$_2$, tetraethyl orthosilicate (TEOS) and Eu$_2$O$_3$. Obtained from a high-purity chemicals laboratory, the materials had 99.9% purity. To prepare appropriate solutions, these raw materials were dissolved in distilled water and nitric acid, and then the required amount of each solution was collected on a quartz plate. Consequently, the mixture of citric acid and ethylene glycol (2.5 mol/10 mol ratio) was then added to the plate.

The solution was dried at 120°C until a transparent solution was obtained. It was then reheated to 200°C to initiate a condensation reaction that would produce a viscous polymeric resin. The resin was heat-treated at 350°C to form a charcoal-like porous foam; it was successively milled and then fired in a tubular furnace at 1350°C for 3 h. At this time the gas flow was adjusted to yield a mixture of 4 volumes of nitrogen to 1 volume of hydrogen. The powder samples were confirmed to be an $\alpha'$.Sr$_2$SiO$_4$ single phase with an orthorhombic structure. The luminescent characteristics of fired samples were obtained by a Perkin-Elmer LS-50 luminescence spectrometer with a flash lamp ($\Delta t = 10$ $\mu$s) as the excitation source. To compare the white luminescent LEDs between YAG:Ce and Sr$_2$SiO$_4$:Eu phosphors, white luminescence conversion LEDs were fabricated. Based on the standard LED technology, GaN (400-nm chip)-based Sr$_2$SiO$_4$:Eu white LEDs were encapsulated in a transparent epoxy resin. In the case of the commercially available YAG:Ce phosphor, however, the InGaN (460-nm chip) was used because its 460-nm blue emission was used as an optical excitation of the YAG phosphor. The relative emission spectra of a GaN-based Sr$_2$SiO$_4$:Eu and InGaN-based YAG:Ce LED under a forward bias of 20 mA was measured using a 50-cm single-grating monochrometer.

Fluorescence spectra of Sr$_2$-$x$SiO$_4$:Eu$_x$ samples with different Eu$^{2+}$ content are shown in Fig. 1. As can be seen from this figure, the emission spectra show a broadband character, and the fluorescence intensity increases with $x$, reaching a maximum at $x = 0.03$ mol. The broadness of the emission band indicates an interaction between the host and the activator, which can be attributed to the presence of an excited electron in an outer shell of the Eu$^{2+}$ ion. The emission band at about 520 nm shifts slightly to a longer wavelength with an increase in Eu$^{2+}$ concentration. This may be attributed to some changes in the crystal field around Eu$^{2+}$ that increased the Eu$^{2+}$ concentration. Although the 4f elec-

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trons of Eu$^{2+}$ are not sensitive to a lattice environment due to the shielding function of the electrons in the outer shell, the 5$d$ electrons are split by the crystal field, which may lead to a shift in its emission peak.

According to Ref. 6, the probability of an energy transfer among Eu$^{2+}$ ions increases when the Eu$^{2+}$ concentration increases. A nonradiative energy transfer from one Eu$^{2+}$ ion to another Eu$^{2+}$ ion usually occurs as a result of an exchange interaction, radiation reabsorption or a multipole–multipole interaction. In the case of the Eu$^{2+}$ ion, the $4f^7 \rightarrow 4f^6 5d^1$ transition is allowed, while the exchange interaction is responsible for the energy transfer of forbidden transitions and typical critical distance, which is about 5 Å. It is indicated that the exchange interaction plays no role in the energy transfer between Eu$^{2+}$ ions. Since the $4f \rightarrow 5d$ transition of Eu$^{2+}$ is allowed, the energy transfer in the present case will occur only as a result of an electric multipolar interaction. As an increase of Eu$^{2+}$ concentration, the distance between Eu$^{2+}$ ions becomes less, and the probability of energy transfer among Eu$^{2+}$ ions increases. In other words, the probability of Eu$^{2+}$ ions at higher levels of 5$d$, which make an energy transfer to the lower 5$d$ levels of Eu$^{2+}$ ions, increases with an increase of Eu$^{2+}$ concentration. 6 It makes it possible to shift emission peak to the longer wavelength with an increase of Eu$^{2+}$ concentration.

As already mentioned in Fig. 1, the shifting of the emission band to a longer wavelength with an increase of the SiO$_2$ content is also observed. Table I presents the main excitation peaks, the emission peaks, the Stokes shift and the crystal field splitting (CFS). The excitation spectrum of sample A (Sr/Si = 2/0.5) consists of two bands peaking around 332 and 382 nm, these two bands have almost merged and broadened so that the excitation spectrum appears as a broad single band. In addition, the emission band of sample A shows a single broad band maximizing at 523 nm, with the Stokes shift of around 7057 cm$^{-1}$, and the CFS is 3943 cm$^{-1}$. The emission band in sample D (Sr/Si = 2/1.3), on the other hand, was observed at 555 nm. This, compared to other samples, is located at a longer wavelength accompanied by increases of the Stokes shift and CFS energy. It should be noted that the shift of the emission band to a longer wavelength occurs as the SiO$_2$ content increases. The shift of the Eu$^{2+}$ emission band depends on the host lattice, covalency, and the strength of the crystal field. The increase of the SiO$_2$ content indicates an increasing degree of covalency. As covalency increases, the interaction between the electrons is reduced, so that they spread out over wider orbitals. Consequently, an increase in the covalency will reduce the energy difference between the ground state $4f^7$ and the excited state $4f^6 5d^1$, so that there will be more splitting of the $t_{2g}$ and $e_g$ levels (which means a larger CFS). This increase in covalency may lead to the shifting of the emission band to a longer wavelength.

The relative emission spectra of the white-emitting InGaN (460-nm chip)-based YAG:Ce LED and the GaN (400-nm chip)-based Sr$_2$SiO$_4$:Eu LED are shown in Fig. 2. In the case of the InGaN-based YAG:Ce LED, two distinct emission bands from the InGaN-based LED and the YAG:Ce phosphor are clearly resolved at 460 nm and at around 550 nm, respectively. On the other hand, the GaN-based Sr$_2$SiO$_4$:Eu LED shows two bands at 400 and 560 nm. The 400-nm emission band is due to a radiative recombination from a GaN active layer. This blue emission was used as an optical transition of the Sr$_2$SiO$_4$:Eu phosphor. The 560-nm emission band is ascribed to a radiative recombination of Eu$^{2+}$ impurity ions in the Sr$_2$SiO$_4$ host matrix. It is presumed that energy from the 400-nm excitation band may be trapped at the Eu$^{2+}$ ion, emitting a yellow emission at 560 nm. These two emission bands combine to give a spectrum

![FIG. 1. The Eu$^{2+}$ concentration dependence photoluminescence emission spectra of the Sr$_2$SiO$_4$ system under the 410-nm excitation wavelength.](Image)

![FIG. 2. Relative emission spectra of a white-emitting InGaN-based YAG:Ce LED and GaN-based Sr$_2$SiO$_4$:Eu LED under a 20-mA drive current.](Image)

**TABLE I. Spectral parameters of the Sr$_2$SiO$_4$:Eu for various SiO$_2$ content.**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Excitation maximum (nm)</th>
<th>Emission maximum (nm)</th>
<th>Stokes shift (cm$^{-1}$)</th>
<th>CFS (cm$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>332, 382</td>
<td>523</td>
<td>7057</td>
<td>3943</td>
</tr>
<tr>
<td>B</td>
<td>332, 384</td>
<td>527</td>
<td>7067</td>
<td>4079</td>
</tr>
<tr>
<td>C</td>
<td>332, 387</td>
<td>533</td>
<td>7078</td>
<td>4281</td>
</tr>
<tr>
<td>D</td>
<td>332, 394</td>
<td>555</td>
<td>7363</td>
<td>4730</td>
</tr>
</tbody>
</table>

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that appears white to the naked eye. The luminous efficiency of GaN-based Sr$_2$SiO$_4$:Eu measured from these results is about 3.8 lm/W, which is higher than that of the industrially available InGaN-based YAG:Ce (CRI=80). However, we have recently obtained evidence for the possibility of further improvements in the CRI by optimizing the peak wavelength of the main yellow emission.

In summary, we have synthesized an Eu$^{2+}$-activated Sr$_2$SiO$_4$ yellow phosphor, and investigated its luminescent properties in an attempt to develop white LEDs by integrating the GaN blue LED chip and the Sr$_2$SiO$_4$:Eu phosphor; the combination of a blue LED emitting (400 nm) and a broadband yellow Sr$_2$SiO$_4$:Eu phosphor creates white line. The main emission peak shifts to a longer wavelength as the SiO$_2$ content increases. This may be understood in terms of the degree of covalency and crystal field splitting of the $5d$ configuration. The white GaN-based Sr$_2$SiO$_4$:Eu developed in this work showed a higher luminous efficiency compared with the industrially available InGaN-based YAG:Ce.

A wide range of whitish colors can be realized by fabricating GaN-based LED with different amounts of Sr$_2$SiO$_4$:Eu concentration. The CIE chromaticity of GaN-based LED with different amounts of Sr$_2$SiO$_4$:Eu concentration is shown in Fig. 3. As the concentration of Sr$_2$SiO$_4$:Eu increases, the color shifts from blue to yellow. In addition, the chromaticity coordinates are close to the straight line interconnecting the points of the blue pump and the yellow phosphor. The color rendering index (CRI) of a GaN-based Sr$_2$SiO$_4$:Eu white LED packaged in a standard epoxy lamp configuration with a reflecting cup with a CIE chromaticity ($x=0.39, y=0.41$) is 68. This value is lower than that of the industrially available InGaN-based YAG:Ce (CRI=80).

FIG. 3. CIE chromaticity points of GaN-based Sr$_2$SiO$_4$:Eu LED and of the corresponding GaN LED. The chromaticity points of GaN-based Sr$_2$SiO$_4$:Eu LED (marked by square), produced by varying the Sr$_2$SiO$_4$:Eu concentration, are on the straight line connecting the chromaticity point of GaN LED (marked by a closed circle).