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Modification of crystal structure of \( \text{Sr}_{2-\chi}\text{Ba}_\chi\text{Si}(\text{O,N})_4 \): Eu\(^{2+}\) phosphors to improve luminescence properties

Xiaojun Li, Youjie Hua, Hongping Ma, Degang Deng and Shiqing Xu

Abstract: A series of \( \text{Sr}_{1.86}\text{Ba}_\chi\text{Si}(\text{O,N})_4 \) \((0.0 \leq \chi \leq 0.5)\) phosphors were synthesized by conventional solid state reaction method. The Sr-N and Si-N bond could be observed in FT-IR spectra. The XRD refinement results indicated that N\(^3\) would substitute for O\(^2-\) site and form Si-NO3 tetrahedrons during the process of forming \( \text{Sr}_{1.86}\text{Ba}_\chi\text{Si}(\text{O,N})_4 \). \( \text{Eu}^{2+}\) \((\text{SSON : Eu}^{2+})\) substitional solid solution. The lattice constants of \( \text{Sr}_{1.86}\text{Ba}_\chi\text{Si}(\text{O,N})_4 \) \((0.0 \leq \chi \leq 0.5)\) were expanded due to the longer bond length of Ba-O. Compared with \( \text{Sr}_2\text{SiO}_4 \), the SSON: Eu\(^{2+}\) showed a remarkable red-shift, which is due to the nephelauxetic effect and the strong crystal field splitting originated from stronger covalent bonding effect of Eu-N bond. The SSON: Eu\(^{2+}\) present \( \beta \) phase structure before the introduction of Ba\(^{2+}\), whereas the \( \alpha' \)-SSBON phase was obtained by the substitution of Ba\(^{2+}\) for Sr\(^{2+}\). Ba\(^{2+}\) doping led to an obvious blue-shift under 375 nm and 460 nm excitation. The 5d orbital of Ba\(^{2+}\) coupled with the 5d orbital of Eu\(^{2+}\) on the higher energy level position in the host crystal. Under 375 nm excitation, the PL intensity gradually increased with the increase of Ba\(^{2+}\) content. Under 460 nm excitation, the PL intensity gradually declined with the increase of Ba\(^{2+}\) content. The thermostability of \( \alpha' \)-SSBON: Eu\(^{2+}\) was significantly improved compared with \( \beta \)-SSO: Eu\(^{2+}\) and \( \beta \)-SSON: Eu\(^{2+}\). On the base of the adjustable emission wavelength, enhanced PL intensity and excellent thermostability in SSBON: Eu\(^{2+}\), we anticipate that these materials can be used as green or red phosphor in white light emitting diodes.

1 Introduction

Recently, a new package type that combined with color-tunable phosphors and n-UV or blue chips has been developed rapidly. As a series of important silicate materials, the RE-doped \( \text{M}_x\text{SiO}_4 \) \((\text{M}=\text{Ca}, \text{Sr}, \text{Ba}, \text{Mg})\) phosphors can easily achieved colour tuning through the chemical composition and structural modification. Such as the \( \text{Sr}_x\text{Ba}_{2-\chi}\text{Si}_\chi\text{SiO}_4 \): Eu\(^{2+}\) and \( \text{Ca}_{x-\chi}\text{Sr}_\chi\text{Si}_\chi\text{Al}_\chi\text{O}_4 \): yCa\(^{2+}\), yLi\(^{2+}\) phosphors with enhanced luminescence properties by doped ions,\(^{1,2}\) the white emitting phosphors of single materials \((\text{Ba},\text{Ca})\text{Si}_4\text{O}_9\) \():\text{Eu}^{2+}\), Mn\(^{2+}\) and Li\(_2\text{SrSiO}_4\): Eu\(^{2+}\), Ce\(^{3+}\), \( \text{Li}^{+}\), \( \text{Eu}^{2+}\) and BaMgSiO\(_4\): Eu\(^{2+}\).\(^{6,7}\) These studies suggest that the colour and crystal structure tunable RE-doped \( \text{M}_x\text{SiO}_4 \) \((\text{M}=\text{Ca}, \text{Sr}, \text{Ba}, \text{Mg})\) phosphors have enormous potential for the application of phosphor-converted white light emitting diodes (pc-LEDs).

The \( \text{Sr}_2\text{SiO}_4 \): Eu\(^{2+}\) phosphors have been extensively studied due to their special structural features and potential applications in developing white light-emitting-diodes.\(^{8-11}\) The orthosilicate phosphor of \( \text{Sr}_2\text{SiO}_4 \): Eu\(^{2+}\) has two crystallographic phases: high temperature \( \alpha' \)-Sr\(_2\text{SiO}_4 \) (orthorhombic) phase and low temperature \( \beta \)-Sr\(_2\text{SiO}_4 \) (monoclinic) phase.\(^{12-14}\) Both the \( \alpha' \) and \( \beta \) phase have two luminescence center: Eu(I) and Eu(II). Eu(I) is ten-coordinated and Eu(II) is nine-coordinated by oxygen atoms within a limited range.\(^{15-17}\) Recently, due to the remarkable emission in the red wavelength, the \( \text{Sr}_2\text{SiO}_4 \): Eu\(^{2+}\) with N\(^3\) substitution \((\text{SSON : Eu}^{2+})\) has been reported in some literatures.\(^{18-20}\) These research mainly included the following five aspects: (a) the detailed crystal structure;\(^{19}\) (b) the replacement of \( \text{M}^{3+} \) cations \((\text{M}=\text{Ca}, \text{Ba}, \text{Mg})\);\(^{20}\) (c) the distinction from Eu(I) and Eu(II) due to nitridation;\(^{21}\) (d) the detail of coordination environment of Eu\(^{2+}\) and the interaction mechanism of nitrogen on red-shift emission;\(^{22}\) (e) the effects of \( \text{N}^3 \) ions on the crystal structure and luminescent properties of \( \alpha'\)-Sr\(_2\text{SiO}_4\text{N}_x\text{O}_{4-x} \): Eu\(^{2+}\) \((1.33 \leq x \leq 2.4)\) by varying N content.\(^{23}\) However, the phase transformation of SSON: Eu\(^{2+}\) and the change of PL intensity with varying substitution content of cations have not been effectively researched. It is necessary to control the phase compositions of the SSON: Eu\(^{2+}\) phosphors and study the change of crystal structure and photoluminescence properties depending on the substitution of \( \text{N}^3\) and Ba\(^{2+}\).

In the case of \( \text{Sr}_2\text{SiO}_4 \), the \( \beta \) (monoclinic) \( \leftrightarrow \) \( \alpha' \) (orthorhombic) phase transformation occurs at 358K.\(^{18}\) The phase transformation occurs through a short-range rearrangement of the coordination structure without breaking coordination bonds (displacive transformation).\(^{18-24}\) High calcination temperature promoted the phase transformation from \( \alpha'\)-Sr\(_2\text{SiO}_4 \) (orthorhombic) to \( \beta\)-Sr\(_2\text{SiO}_4 \) (monoclinic), while the doping of Eu or Ba ions could stabilize \( \alpha' \)-Sr\(_2\text{SiO}_4 \) phase due to their long bond length with oxygen.\(^{10,25}\) In this paper, the \( \text{Sr}_{1.86}\text{Ba}_\chi\text{Si}(\text{O,N})_4 \) \((0.0 \leq \chi \leq 0.5)\) phosphors were successfully synthesized by the conventional solid-state reaction method. We investigated the effects of Ba\(^{2+}\) substitution on the crystal structural and luminescence properties
of α′-SSON: Eu²⁺ by varying Ba²⁺ content. The Sr-N and Si-N bond were confirmed by crystal structure and FT-IR spectra and the specific site of N substituting for O was determined. The specific reasons of PL intensity change and phase transformation were explained. The crystal of SSON: Eu²⁺ could be effectively controlled by Ba²⁺ doping and could achieve higher PL intensity of SSON: Eu²⁺ based on modification of the crystal structure.

2 Experimental

2.1 Sample preparation

The Sr₁₋ₓBaₓSi(O,N)₁₋ₓ: 0.02Eu²⁺ (0 ≤ x ≤ 0.5) was prepared by solid state reactions in a horizontal tube furnace using starting materials of SrCO₃ (AR), BaCO₃ (AR), α-Si₃N₄ (Alfa 99.9%), SiO₂ (AR) and Eu₂O₃ (99.9%).

All the raw materials were mixed in an agate mortar and then filled into BN crucibles. The powder mixtures were preheated at 1050 °C for 2 h, and then fired at 1450 °C (with a heating rate of 5 °C/min) for 7 h, followed by cooling down to 300 °C at a rate of 5 °C/min and down to room temperature spontaneously in the furnace with the power switched off. In order to prevent samples from being oxidized, all heating and cooling processes were conducted in a flowing reductive atmosphere of 5:95 (volume) H₂/N₂.

2.2 Characterization

The phase composition and crystallinity of the synthesized samples were investigated by powder X-ray diffraction (Bruker AXS D2 PHAREX diffractometer) with Cu Kα radiation (λ = 1.5405 Å) over the angular range 10° ≤ 2θ ≤ 80°. The excitation and emission spectra of the phosphors were measured at room temperature on a PL3-211-P spectrometer (HORIBA JOBIN YVON, America) and a 450W xenon lamp was used as the excitation source. The PL decay-curves were measured on the same spectrometer, which was combined with a Time-Correlated Single-Photon Counting (TCSPC) system. The crystallographic data including lattice parameters, unit cell volume, and phase ratios were estimated by the TOPAS package based on the XRD data, using the Sr₃SiO₄ structure as a starting model. The nitrogen and oxygen contents (Atomic %) were measured by energy dispersive spectrometry system (TEAM Apollo XL EDS, EDAX, America). Fourier-transform infrared spectra (FT-IR) were measured on a BRUKER TENSOR 27 spectrophotometer in the range of 400–4000 cm⁻¹ using the KBr pellet (~2wt %) method. The absorbance spectra of the samples were measured by an ultraviolet-visible-near infrared spectrophotometer (UV3600) using BaSO₄ as a reference in the range of 200–800 nm. All the above measurements were performed at room temperature.

3 Results and discussion,

3.1 X-ray diffraction analysis and crystal structure

Fig. 1 (a) shows the XRD patterns of β-Sr₁₋ₓBaₓSi(O,N)₁₋ₓ: 0.02Eu²⁺ and Sr₁₋ₓBaₓSi(O,N)₁₋ₓ: 0.02Eu²⁺ (0 ≤ x ≤ 0.5) with different x values. As shown in Fig. 1, the Sr₁₋ₓSi(O,N)₁₋ₓ: 0.02Eu²⁺ (x=0, SSON: 0.02Eu²⁺) matched well with the β-Sr₃SiO₄ (β-SSO, JCPDS no.38-0271) phase. In comparison with the β-Sr₃SiO₄, the degree of crystallinity of β-SSON became worse. However, with the increase of x value from 0.1 to 0.5, all the samples of the Sr₁₋ₓBaₓSi(O,N)₁₋ₓ: 0.02Eu²⁺ (0 ≤ x ≤ 0.5, SBSON: 0.02Eu²⁺) matched well with the α′-Sr₃SiO₄ (α′-SSO, JCPDS no.39-1256) phase. All the α′-SSON phases have similar XRD patterns because α′-Sr₁₋ₓBaₓSiO₄ had the same structure with α′-Sr₃SiO₄ where divalent Ba²⁺ ions occupy the sites of Sr²⁺. The degree of crystallinity of α′-SSON gradually decreased with the increase of Ba content. Besides, there were no Ba₃SiO₄ phases in SBSON when the Ba²⁺ ions substitution content increased from 0.1 to 0.5. These results indicate that the β-SSON transformed from β to α′ phase because Ba²⁺ ions substituted for Sr²⁺ ions. Fig. 1 (b) shows the local amplificatory XRD patterns of SBSON: 0.02Eu²⁺ in the range of 26.25°–32°. As shown in Fig. 1 (b), Ba²⁺ doping led to the results that XRD diffraction peaks shifted in the direction of smaller angle. This is because Ba²⁺ doping led to the lattice expansion of the α′ phase depending on the larger ionic radius of Ba²⁺ (1.49 Å for 9CN, 1.52 Å for 10CN) than that of Sr²⁺ (1.30 Å for 10CN).

Fig. 1 XRD patterns of Sr₁₋ₓBaₓSi(O,N)₁₋ₓ: 0.02Eu²⁺ (0 ≤ x ≤ 0.5) with various Ba²⁺ content and β-Sr₃SiO₄: 0.02Eu²⁺.
The crystal structure of α′-Sr$_{1.9}$Ba$_{0.1}$Si(O,N)$_4$: 0.02Eu$^{2+}$ was analyzed by TOPAS package on the basis of the XRD data, using the α′-Sr$_2$SiO$_4$ (ICSD: 35667), β-Sr$_2$SiO$_4$ (ICSD: 36041) structure as starting models. Fig. 2 shows the observed (black line), calculated (red line) and difference (green line) XRD profiles for the XRD Rietveld refinement of α′-Sr$_{1.9}$Ba$_{0.1}$Si(O,N)$_4$: 0.02Eu$^{2+}$. The peaks (2θ=27.83° and 32.81°) of β phase was marked as β symbol in Fig. 2.

These results indicate that a predominant pure α′-SBSON phase have been formed accompanied by a small portion (2.26%) of β phase.

Table 1 lists the XRD refinement data of the α′-Sr$_{1.9}$Ba$_{0.1}$Si(O,N)$_4$. These refinement data include crystallographic parameters, reliability factors, wyckoff site, atomic coordinates, occupancy, and beq. The α′-Sr$_{1.9}$Ba$_{0.1}$SiO$_4$ crystallizes in an orthorhombic unit cell belonging to a space group of Pmnb: ba-c (no. 62). The refinement finally converged to $R_{wp}$= 5.08, $R_{p}$=6.37, $R_{p}$=4.99 and GOF =1.25, as shown in Table 2. There are obvious differences of the atomic positions and cell volume between the α′-Sr$_{1.9}$Ba$_{0.1}$SiO$_4$: Eu$^{2+}$ (391.81 Å$^3$) and α′-Sr$_{1.9}$Ba$_{0.1}$Si(O,N)$_4$: Eu$^{2+}$ (395.8783 Å$^3$) according to ref. 8 and 12.

Table 1 Rietveld refinement and crystal data for α′-Sr$_{1.9}$Ba$_{0.1}$Si(O,N)$_4$

<table>
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<tr>
<th>Site</th>
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<th>x</th>
<th>y</th>
<th>z</th>
<th>Atom</th>
<th>Occ</th>
<th>Beq</th>
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<tbody>
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<td>Sr1</td>
<td>4c</td>
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<td>0.34014</td>
<td>0.58431</td>
<td>Sr$^{2+}$</td>
<td>0.95</td>
<td>0.6832</td>
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<td>4c</td>
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<td>0.34014</td>
<td>0.58431</td>
<td>Ba$^{2+}$</td>
<td>0.05</td>
<td>0.6832</td>
</tr>
<tr>
<td>Sr2</td>
<td>4c</td>
<td>0.23628</td>
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<td>0.30213</td>
<td>Sr$^{2+}$</td>
<td>0.95</td>
<td>1.913</td>
</tr>
<tr>
<td>Ba2</td>
<td>4c</td>
<td>0.23628</td>
<td>0.00114</td>
<td>0.30213</td>
<td>Ba$^{2+}$</td>
<td>0.05</td>
<td>1.913</td>
</tr>
<tr>
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<td>O2</td>
<td>8d</td>
<td>0.47287</td>
<td>0.47287</td>
<td>0.65649</td>
<td>O$^{2-}$</td>
<td>1</td>
<td>-7.837</td>
</tr>
</tbody>
</table>

Fig. 3 (a) A unit cell of the α′-Sr$_{1.9}$Ba$_{0.1}$Si(O,N)$_4$: Eu$^{2+}$ view in the [101] direction along b axis; (b) Coordination spheres of the Si and two different Sr$^{2+}$/Ba$^{2+}$ sites of the ICSD-36042: ordered α′-Sr$_{1.9}$Ba$_{0.1}$SiO$_4$; (c) Coordination spheres of the Si and two different Sr$^{2+}$/Ba$^{2+}$ sites of the α′-Sr$_{1.9}$Ba$_{0.1}$Si(O,N)$_4$. 
Fig. 3 (a) shows a unit cell of the α′-Sr1.9Ba0.1Si(O,N)4:Eu2+ view toward the [111] plane. The Si (N/O) 4 tetrahedra and Sr/Ba polyhedra showed a hexagonal pseudo-symmetry along [101] in the single unit cell, as confirmed by the c/a ratio which was close to $\sqrt{3}$. The structural type is characterized by [010] chains of Sr coordination polyhedral sharing faces, which are linked with one another forming a three-dimensional framework of polyhedra.12

Fig. 3 (b) shows the coordination spheres of the Si and two different Sr2+/Ba2+ sites of the α′-Sr1.9Ba0.1SiO4, which is obtained by Rietveld refinement data in Table 1.

Due to the change of O atoms positions and occupation, the α′-Sr3SiO4 phase has two structure models: disordered (isotropic) and ordered (anisotropic) model.13 However, the mixed-model in α′-Sr3SiO4 as follows: The O1 and O2 atoms lying on the mirror plane (x = 0.25) was split into two, in very close positions equivalent by m symmetry, with 0.5 occupation factor (o.f.) and refined isotropically (disordered). The O1′, O2′ were out of the mirror plane (x ≠ 0.25, o.f. = p) and O1″, O2″ were in the mirror plane (x = 0.25, o.f. = 0.5-p). The o.f.s of O3 and O4 were constrained to be 1-p and p, respectively. The p parameter was ranging from 0 (pure order phase) to 0.5 (pure disorder phase). As shown in Fig. 3 (a) and (c), in our experiments, the results of refinement indicate that α′-Sr1.9Ba0.1SiO4 presents mixed ordered (p=0) and disordered (the symmetric splitting sites of Si, O1 and N atoms) phase feature. However, the ordered α′-Sr2+ phase was dominate because the disordered phase has four different O atoms sites. In addition, Fig. 3 (b) and (c) indicated that the Sr/Ba2+ ions in α′-SBSO and α′-SBSON have two sites: [Sr/Ba]1 and [Sr/Ba]2. [Sr/Ba]1 is 10-fold coordinated (CN=10) whereas [Sr/Ba]2 is 9-fold coordinated (CN=9) by oxygen atoms within a limited range.

The bond length of Si-O2 increased from 1.630 Å to ~1.72 Å after N− substituting for O2−. The observed Si–N bonds in other silicon-based oxynitrides were in the range of 1.660~1.750 Å, as shown in Table 2. Therefore, it could be speculated that the N− would substitute for O2− site during the process of formation of α′-SBSON solid-solution, as shown in Fig. 3 (c). However, paradoxically, the Si–O1, Si–O3 and Si–O3′ bond lengths in α′-Sr1.9Ba0.1Si(O,N)4 deviated from the normal ranges observed in other silicon-based oxynitrides and got shorter than ordered α′-Sr1.9Ba0.1SiO4, as shown in Table 2. Thus the Si–O1, Si–O3, Si–O3′ and Si–N in α′-SBSON formed Si-NO3 tetrahedrons. The α′-Sr1.9Ba0.1Si(O,N)4 is a substitutional solid solution due to the substitutions of N− and Ba2+ ions. During the process of forming Si-NO3 tetrahedrons in substitutional solid solution α′-SBSON, the Si–O bonds were squeezed by the larger Si–N bonds due to the covalent bond effect.23,26 Hence, the Si–O1, Si–O3 and Si–O3′ bonds got shorter obviously after N− substituting for O2− site. The other silicon-based oxynitrides didn’t appear obvious squeeze effect because they are not solid-solutions. Thus the Si–O bond lengths of other silicon-based oxynitrides are within the normal ranges. These results demonstrate that N− ions have partially been incorporated into the α′-SBSO lattice, forming a solid-solution of α′-SBSON.

Table 2 The Si–(N/O) bond lengths with different tetrahedrons types

<table>
<thead>
<tr>
<th>Silicon-based oxynitrides</th>
<th>Tetrahedron types</th>
<th>Si–O bond lengths (Å)</th>
<th>Si–N bond lengths (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y2Si2O7N4(ref. 27)</td>
<td>Si–NO3</td>
<td>1.595–1.706</td>
<td>1.708, 1.731</td>
</tr>
<tr>
<td>Sr2Si2O7N4(ref. 28)</td>
<td>Si–NO3</td>
<td>1.619–1.657</td>
<td>1.660–1.732</td>
</tr>
<tr>
<td>SrSi2O7N4(ref. 29)</td>
<td>Si–N2O</td>
<td>1.603–1.608</td>
<td>1.740–1.746</td>
</tr>
<tr>
<td>ordered α′-SBSON</td>
<td>Si–O3</td>
<td>1.603–1.630</td>
<td></td>
</tr>
<tr>
<td>α′-SBSON</td>
<td>Si–(N/O)4</td>
<td>1.458–1.558</td>
<td>1.800</td>
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</table>

Fig. 4 shows the bond length change of Sr1/Ba1-O and Sr2/Ba2-O bonds in the ordered α′-Sr1.9Ba0.1SiO4 and α′-Sr1.9Ba0.1Si(O,N)4. All the bond length values were obtained from Fig. 3. To be convenience for comparison, the N and O2 in SBSON was set as O2 and O3 respectively, which was consistent with the ordered α′-Sr1.9Ba0.1SiO4.

As shown in Fig. 4, compared with the other (Sr, Ba)-O bond length, the (Sr, Ba)-O1, O2 bond length of α′-SBSON got longer than that of ordered α′-SBSO. Besides, the variation of (Sr, Ba)-O1, O2 bond length was more obvious than that of (Sr, Ba)-O2, O3 bond. On the other hand, the larger ionic radius will lead to longer ionic bond length because the ionic bond length is the sum of anion and cation radius. The ionic radius of N− (1.32 Å) is larger than that of O2− (1.24 Å). So theoretically, the N− substitution for O2− could make the bond length of Sr–O get longer. However, the Sr2-O1 bond length of SBSON got shorter obviously than that of SBSO, which should be ascribed to squeeze effect. It would lead to different spectra variation between the Eu(I) and Eu(II) sites. The results of our experiment are consistent with the theoretical analysis. Therefore, we can speculate that N− was easier to substitute for the O2 site than O3 site, which was consistent with the Si–O/N bond length analysis of Si–NO3 tetrahedrons as shown in Fig. 3.

**Table 3** shows the results of XRD Rietveld refinement data of Sr1.98Ba0.02Si(O,N)x: 0.02Eu2+ (0 ≤ x ≤ 0.5) with various Ba content. As shown in Table 3, the phase transformation of SSBN from β to α′ phase obviously occurred when 0.1 mol Ba2+ ions entered into the...
crystal lattice. With the Ba content increasing from 0.1 to 0.5, the α′ phase was the major phase and the unit cell volume gradually increased. The XRD Rietveld refinement results showed that a small amount of Eu$^{2+}$ and Ba$^{2+}$ ions could enter into the host lattice of α′-SSON without destroying the crystal structure. In fact, Ba$^{2+}$ had been reported as a stabilizer with the function of stabilizing metastable phase due to its larger ionic radius (0.152 nm). It is known that the high-temperature α′-Sr$_2$SiO$_4$ phase can be stabilized at room temperature by substituting Ba$^{2+}$ ions on the crystalline structure of α′-SSON: Eu$^{2+}$. The α′ ↔ β transition of Sr$_2$SiO$_4$ can be simply accomplished by a rearrangement of atoms without breaking coordination bonds (displace transformation). When a proper amount of Ba$^{2+}$ was entered into SSON, similar to SSO, the displacive transformation from β-SSON to α′-SSON phase occurred, as shown in Fig. 1 and Table 3.

Table 3. Lattice parameters of Sr$_{1-x}$Ba$_x$Si(O,N)$_4$: 0.02Eu$^{2+}$ (0 ≤ x ≤ 0.5) [unit: Å, degree]

<table>
<thead>
<tr>
<th>Sample</th>
<th>x</th>
<th>a</th>
<th>b</th>
<th>c</th>
<th>β</th>
<th>V (Å$^3$)</th>
<th>Rwp</th>
<th>α′-SSON</th>
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<td>0 (β)</td>
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<td>7.090</td>
<td>9.773</td>
<td>90</td>
<td>393.71</td>
<td></td>
<td></td>
</tr>
<tr>
<td>β-SSO</td>
<td></td>
<td>5.663</td>
<td>7.084</td>
<td>9.767</td>
<td>92.67</td>
<td>391.39</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fig. 5 Unit cell volume of SBO: 0.02Eu$^{2+}$ (0 ≤ x ≤ 0.5), α′-Sr$_2$SiO$_4$ and β-Sr$_2$SiO$_4$.

Fig. 5 indicates the unit cell volume of SBO: 0.02Eu$^{2+}$ (0 ≤ x ≤ 0.5), α′-SSO and β-SSO. The unit cell volume gradually increased with the increase of the doped Ba$^{2+}$ content. The ionic radius of Ba$^{2+}$ (1.49 Å for 9CN, 1.52 Å for 10CN) is larger than that of Sr$^{2+}$ (1.30 Å for 9CN, 1.35 Å for 10CN) and the ionic bond length is the sum of anion and cation radius. So the bond length of Ba-O is longer than that of Sr-O. Therefore, the expansion of the lattice constants with increasing Ba-content should be ascribed to the longer bond length of Ba-O than that of Sr-O.

3.2 N/O contents for Sr$_{1.98-x}$Ba$_x$Si(O,N)$_4$: 0.02Eu$^{2+}$ (0 ≤ x ≤ 0.5) as a function of x

The N/O contents were measured by energy dispersive spectrometry system (EDS). Table 4 shows the experimental and theoretical calculation values of N/O contents. With the increase of x from 0 to 0.5, the experimental and the theoretical values of N/O contents in SSON or SBO had a little difference due to the surface oxidation and the measuring error. When the introduction of Ba$^{2+}$ ions increased from 0 to 0.1, the nitrogen content increased from 13.70 % to 15.67 % obviously. With the further introduction of Ba$^{2+}$ ions, the nitrogen content hardly changed. This can be ascribed to the stabilization effect of Ba$^{2+}$ in metastable phase.

Table 4. The measured oxygen and nitrogen content in Sr$_{1.98-x}$Ba$_x$Si(O,N)$_4$: 0.02Eu$^{2+}$ (0 ≤ x ≤ 0.5) (Atomic %)

<table>
<thead>
<tr>
<th>Sample</th>
<th>Nitrogen</th>
<th>Oxygen</th>
<th>Theoretical value of the N contents (Atomic %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSON(x=0)</td>
<td>13.70</td>
<td>31.87</td>
<td>21.05</td>
</tr>
<tr>
<td>SBSN(x=0.1)</td>
<td>15.67</td>
<td>31.83</td>
<td>21.05</td>
</tr>
<tr>
<td>SBSN(x=0.2)</td>
<td>15.55</td>
<td>32.34</td>
<td>21.05</td>
</tr>
<tr>
<td>SBSN(x=0.3)</td>
<td>15.09</td>
<td>31.77</td>
<td>21.05</td>
</tr>
<tr>
<td>SBSN(x=0.4)</td>
<td>15.48</td>
<td>31.34</td>
<td>21.05</td>
</tr>
<tr>
<td>SBSN(x=0.5)</td>
<td>15.63</td>
<td>31.90</td>
<td>21.05</td>
</tr>
</tbody>
</table>

3.3 Comparison of FT-IR spectra for α′-SSO: Eu$^{2+}$, α′-SBSN: Eu$^{2+}$, β-SSO: Eu$^{2+}$, and β-SBSN: Eu$^{2+}$

Fig. 6 presents the FT-IR spectra of α′-SSO: Eu$^{2+}$, α′-SBSN: Eu$^{2+}$, β-SSO: Eu$^{2+}$, β-SBSN: Eu$^{2+}$, and Sr$_2$Si$_2$N$_4$: Eu$^{2+}$ phosphors. Apparently, these FT-IR spectra show remarkable differences among these phosphors. The peak of 1475 cm$^{-1}$ was assigned to Sr/Eu–O bond of α′ phase, whereas the peak of 1636 cm$^{-1}$ was assigned to Sr/Eu–O bond of β phase. This was because the monoclinic β phase ($\alpha=\gamma=90^\circ$, $\beta=92.67^\circ$) have higher energy than the orthorhombic α′ phase ($\alpha=\gamma=90^\circ$). The longer bond length would lead to the weaker bond energy. Thus the absorption energy of Sr/Eu–N bond was lower than that of Sr/Eu–O bond due to the longer bond length of Sr–N than that of Sr–O. As shown in Fig. 6, the α′-SSON: Eu$^{2+}$ contained a new peak of 1364 cm$^{-1}$ which was assigned to Sr/Eu–N bond, whereas the α′-SSO didn’t show such peak. The β-SBSO: Eu$^{2+}$ contained a new peak of 1439 cm$^{-1}$ which was also assigned to Sr/Eu–N bond. Besides, the vibration energy of Sr/Eu–N bond in α′-SBSN (1364 cm$^{-1}$) and β-SBSN: Eu$^{2+}$ (1439 cm$^{-1}$) were different from the Sr/Eu–N bond in Sr$_2$Si$_2$N$_4$ (1633 cm$^{-1}$).

As show in the shaded area of Fig. 6, the absorption bands of Si-
(N/O)$_4$ tetrahedrons with symmetric and antisymmetric stretching vibration were in the range of vibration energy (700 cm$^{-1}$-1100 cm$^{-1}$). The larger the distinction of Si-O/N bond energy, the borader the absorption band of Si-(N/O)$_4$ tetrahedrons. The longer bond length would lead to the weaker bond energy. The distinction of Si-O bond length in α′-SSO was larger than β-SSO. Thus the Si-O$_x$ absorption band of α′-SSO was borader than that of β-SSO. The quantity of Si-N bond increased simultaneously along with the absorption band of α′-SSO was borader than that of β-SSO. The phosphors.

As shown in Fig. 6, the Si-N$_x$ absorption was borader and became Si-(N/O)$_4$ absorption band during the process of formation of α′-SSON and β′-SSON solid-solutions, as shown in the shaded area of Fig. 6. Besides, the Si-N$_x$ absorption band of Sr$_x$Si$_2$N$_y$ was narrower than the Si-(N/O)$_4$ absorption of SSON due to the smaller distinction among the Si-N$_x$ bond energy. These results also confirm that nitrogen had entered the crystal lattice and formed chemical bonds with surrounding Sr$^{2+}$/Eu$^{2+}$ ions and Si atoms in the α′-SSBSON: Eu$^{2+}$ and β-SSON phosphors.

![Fig. 6 FT-IR spectra of α′-SSO: Eu$^{2+}$, α′-SSBSON: Eu$^{2+}$, β-SSO: Eu$^{2+}$, β-SSON: Eu$^{2+}$ and Sr$_x$Si$_2$N$_y$: Eu$^{2+}$ phosphors.](image)

**3.4 Luminescence properties of Sr$_{1.98}$Ba$_{0.02}$Si(O,N)$_4$: 0.02Eu$^{2+}$ (0 ≤ x ≤0.5)**

Fig. 7 shows the photoluminescence (PL) spectra of β-SSO: Eu$^{2+}$ and β-SSON: Eu$^{2+}$ under 375 nm and 460 nm excitation and the corresponding photoluminescence excitation (PLE) spectra. The dominant peak wavelengths (DPWs) of the Eu(I) and Eu(II) sites emissions of β-SSO is ~480nm and ~540nm respectively. The DPWs of the Eu(I) and Eu(II) sites emissions of ordered α′-SSO is ~490nm and ~560nm respectively. As shown in Fig. 7, compared with the β-SSO: Eu$^{2+}$, after N$^3$ substituting for O$^3$, under 375nm excitation, the DPWs of Eu(I) (~465nm) and Eu(II) (~536nm) sites emissions appeared a slight red-shift. Under 460nm excitation, the emission peak of Eu(I) site almost disappeared and the emission peak of the Eu(II) sites appeared an obvious red-shift, which was changed from 543nm to 612nm. However, the PL spectra of β-SSON: Eu$^{2+}$ appeared another impurity emission peak at 537nm, which should be attributable to the emission of β-SSO: Eu$^{2+}$ phosphor. In addition, Fig. 6 shows a weaker peak of Sr/Eu–N bond (1439 cm$^{-1}$) also indicated that the Sr$_{1.98}$Ba$_{0.02}$Si(O,N)$_4$ (x=0) sample of our experiments was the mixed phase of β-Sr$_{1.98}$Si(O,N)$_4$ and β-Sr$_{1.98}$SiO$_4$ phase. However, due to the similar electronic configuration between O and N, the β-SSO and β-SSON phases couldn’t be distinguished effectively by X-ray diffraction analysis. Besides, the introduction of N$^3$ would result in the decrease of PL intensity, as shown in the PL and PLE spectra in Fig. 7.

The partial incorporation of N$^3$ ions caused a strong crystal field splitting and gave rise to a nephelauxetic effect. The strong crystal field could be ascribed to the stronger covalent properties of Eu-O/N bond. The nephelauxetic effect was related to appropriate centroid shift in the RE cation d-orbitals and can be attributed to the covalency between the RE cation and the surrounding anions, as well as their polarizability (the performance of the deformation).
excitation level of the Eu\(^{2+}\) ions shifted toward the ground level at the Eu(II) site. Hence, the 5d energy levels of Eu\(^{2+}\) ions is declined, which led to long wavelength emission (~610nm) suitable for low energy blue LED excitation rather than n-UV excitation. And there appeared a red-shift in β-SSO: Eu\(^{3+}\) after N substituting for O. Besides, the Eu(I)(CN=10) sites was not significantly affected by nitridation. This was because Eu(I) (CN=9) sites had smaller coordination number (CN) and the bond length of Eu(II)-N was shorter than that of Eu(I)-N. Unfortunately, as shown in Fig. 3 (b), with the introduction of N\(^{3+}\) ions, an asymmetric Si-N\(_3\)O\(_3\) tetrahedron was observed in local scale, which undermined the crystal symmetry. At the same time, the degree of crystallinity for β-SSO: Eu\(^{3+}\) became worse due to the increased proportion of Si-N and (Sr/Eu)-N bonds, as shown in Fig. 1. Hence, the PL intensity of β-SSO: Eu\(^{3+}\) obviously decreased with the N entered into the lattice, as shown in Fig. 7.

Fig. 8 shows the photoluminescence (PL) spectra of Sr\(_{1.98-x}\)Ba\(_x\)Si\(_3\)O\(_4\)N\(_x\): 0.02Eu\(^{3+}\) (0 ≤ x ≤ 0.5) with varying Ba content under 375nm and 460 nm excitation and the corresponding photoluminescence excitation (PLE) spectra. On the basis of the occupation of sites by Eu\(^{3+}\) in domains of different coordination environment, a color point tuning in single-phase host can be realized, as shown in Fig. 8. In order to ensure the precision of the PL intensity and emission wavelengths, the tests of the PL and PLE spectra were ran 5 times. It shows that the emissio n wavelengths, the tests of the PL and PLE were realized, as shown in Fig. 8. In order to ensure the precision of the PL intensity and emission wavelengths, the tests of the PL and PLE spectra were ran 5 times. It shows that the emissio n wavelengths, the tests of the PL and PLE were ran 5 times. It shows that the emission wavelengths hardly changed while the PL intensity changed slightly, as shown in the inset of Fig. 8. Due to the complexity of the PL spectra, the discussion would be divided into two parts: (1) the PL spectra with the increase of Ba\(^{2+}\) content from 0 to 0.1; (2) the PL spectra with the increase of Ba\(^{2+}\) content from 0.1 to 0.5.

With the substitution of 0.1 mol Ba\(^{2+}\) for Sr\(^{2+}\), under 375nm excitation, the DPWs of the Eu(I) sites changed from 470nm to 490nm and that of the Eu(II) sites changed from 543nm to 579nm. Both the PL intensity of Eu(I) and Eu(II) sites declined. Under 460nm excitation, the DPWs of the Eu(I) sites of β-SSO: Eu\(^{3+}\) (537nm) disappeared and that of β-SSON: Eu\(^{3+}\) (612nm) was hardly changed. Besides, the PL intensity of 612nm was significantly improved. The change of emission peak sites was due to the phase transformation from β phase to α′-SSSON phase as shown in Fig. 1 and Fig.2. The decrease of PL intensity of Eu(I) and Eu(II) sites upon 375 nm excitation was because the phase transformation from β to α′ led to a red-shift. A short wavelength ultraviolet light excited to induce a longer emission wavelength, which need to later stokes shift and energy. The significant improvement of the PL intensity of red emission at Eu(II) sites under 460 nm excitation was because the proper Ba\(^{2+}\) content (x=0.1) could obtain a pure α′ phase (as shown in Fig. 1 and Fig. 2). Whereas the sample of undoped Ba\(^{2+}\) was the mixture phase of β-SSO: Eu\(^{2+}\) and β-SSON: Eu\(^{3+}\).

With the increase of Ba\(^{2+}\) content from 0.1 to 0.5, upon 375nm excitation, the DPWs of the Eu(I) and Eu(II) sites emissions appeared a obvious blue-shift, the PL intensity of Eu(I) and Eu(II) sites gradually raised. Unpon 460nm excitation, the DPWs of Eu(I) sites emissions had hardly changed with a small amount replacement of Ba\(^{2+}\) (0.1 ≤ x ≤ 0.2) and appeared a obvious blue-shift with a large amount replacement of Ba\(^{2+}\) (0.3 ≤ x ≤ 0.5). The PL intensity of Eu(II) sites gradually declined with the increase of Ba content from 0.1 to 0.5. In fact, with the increase of Ba\(^{2+}\) content from 0.1 to 0.5, the PL intensity of SBSNO: Eu\(^{3+}\) gradually increased upon the 375 nm excitation while gradually decreased upon 460 nm excitation. This opposite phenomenon is difficult to be ascribed to the measuring error or external environment (such as temperature, impurity phase, atmosphere and so on). Base on this point, this phenomenon only can be caused by the incorporation of Ba\(^{2+}\). Generally, the ultraviolet light excitation (λ\(_{ex}\)=375nm) made short wavelength emission more efficient, whereas the blue light excitation (λ\(_{ex}\)=460nm) made long wavelength emission more efficient. This could be ascribed to the short wavelength emission at higher energy site and long wavelength emission at lower energy site.\(^{25}\) As shown in Fig. 8, the PLE spectra indicated that the ultraviolet light excitation (λ\(_{ex}\) = 350-400 nm) was more beneficial to short wavelength emission (540nm), whereas the blue light excitation (λ\(_{ex}\)=420-460 nm) was more beneficial to long wavelength emission (610nm). The further interpretations of these phenomena of Fig. 8 will be discussed in detail in the following paragraphs.

Fig. 8 PLE and PL spectra of Sr\(_{1.98-x}\)Ba\(_x\)Si\(_3\)O\(_4\)N\(_x\): 0.02Eu\(^{3+}\) (0 ≤ x ≤ 0.5) with varying Ba content: (a) 375nm excitation and (b) 460nm excitation.
Fig. 9 PLE spectra of Sr$_{1.88}$Ba$_{0.1}$Si(O,N)$_4$:0.02Eu$^{2+}$ corresponded to 490nm, 580nm, 610nm emission, respectively.

Fig. 9 shows the excitation spectra of the α'-Sr$_{1.88}$Ba$_{0.1}$Si(O,N)$_4$:0.02Eu$^{2+}$ monitored at 490nm, 580nm, 610nm emission peaks. As shown in Fig. 9, the emission of Eu(I) (490nm) site only could be excited by UV light. The blue light (λ$_{em}$=460nm) excitation led to efficient longer red light emission (610nm) of Eu(II) site than UV-light excitation. The UV excitation (λ$_{ex}$=375nm) is weighted against the red emission (610nm) of Eu(II) site but make for yellow emission (580nm) of Eu(II) site. The Sr$_{1.88}$Ba$_{0.1}$Si(O,N)$_4$:0.02Eu$^{2+}$ can realize an intense red emission at ~610nm under 400–500 nm excitation.

Fig. 10 PL decay curves of Sr$_{1.88}$Ba$_{0.1}$Si(O,N)$_4$:0.02Eu$^{2+}$ ($\lambda_{em}$=375nm, $\lambda_{ex}$=490nm; $\lambda_{em}$=580nm, $\lambda_{ex}$=375nm; $\lambda_{em}$=610nm, $\lambda_{ex}$=375nm).

The PL decay curves of the Eu$^{2+}$ ions in Sr$_{1.88}$Ba$_{0.1}$Si(O,N)$_4$:0.02Eu$^{2+}$ phosphors were obtained with excitation at 375 nm, monitored at 490 nm and 580 nm respectively, as shown in Fig. 10. The varying decay curves of the Eu$^{2+}$ ions can be fitted well to a typical double-exponential decay curve using the following equation:

$$I(t) = A_1 \exp\left(-\frac{t}{\tau_1}\right) + A_2 \exp\left(-\frac{t}{\tau_2}\right)$$  (2)

where $I$ represents the luminescent intensity, $A_1$ and $A_2$ are constants, $t$ is time, $\tau_1$ and $\tau_2$ are the decay times for the exponential components. The average decay time (τ*) can be calculated using the following equation:

$$\tau^* = \frac{\int_0^\infty I(t)dt}{\int_0^\infty I(t)dt}$$  (3)

where $I(t)$ is the intensity at time $t$. The average lifetime of the Eu$^{2+}$ ions is determined to be 0.3569 μs and 1.0203 μs for α'-SBSON:0.02Eu$^{2+}$ with the monitored wavelength at 490 nm and 580 nm respectively. In general, the emission decay times are similar when active ions occupy the same site. Fig. 8(a) presents the emission decay of Eu$^{2+}$ at 490 nm in Eu(I) site and 580 nm in Eu(II) site upon 375nm excitation. It is clear that the lifetime of Eu$^{2+}$ in two sites above is different, which can demonstrate the existence of two sites in α'-SBSON:0.02Eu$^{2+}$.

Fig. 11 shows the diffuse reflection spectra of β-Sr$_{1.98}$SiO$_4$:0.02Eu$^{2+}$ and Sr$_{1.98-x}$Ba$_x$Si(O,N)$_4$:0.02Eu$^{2+}$ ($x=0$, 0.1 and 0.5). Compared with the β-Sr$_{1.98}$SiO$_4$:0.02Eu$^{2+}$, the absorption band of Sr$_{1.98-x}$Ba$_x$Si(O,N)$_4$:0.02Eu$^{2+}$ appeared a remarkable red-shift, as shown in the right shaded area of Fig. 11. It could be ascribed to the decline of 5d level of Eu$^{2+}$ and larger crystal field splitting depending on the N$^3-$ substitution. With the increase of Ba$^{2+}$ content from 0 to 0.1, the absorption in 400-500 nm increased significantly, while the absorption in 300-400 nm increased slightly. This was because of the phase transformation from β-SSON to α'-SBSON phase. With the increase of Ba$^{2+}$ content from 0.1 to 0.5, the absorption in 400-500 nm hardly changed, while the absorption in 300-400 nm increased significantly. It could be attributed to the enhanced absorption of host lattice due to the large amount of Ba$^{2+}$ introduction.

Fig. 11 Diffuse reflection spectra of β-Sr$_{1.98}$SiO$_4$:0.02Eu$^{2+}$ and Sr$_{1.98-x}$Ba$_x$Si(O,N)$_4$:0.02Eu$^{2+}$ ($x=0$, 0.1 and 0.5).
Fig. 12 shows the diffuse reflection spectra (DRS) of Sr$_2$Si(O,N)$_6$, Sr$_{1.48}$Ba$_0.5$Si(O,N)$_4$ host materials and the Eu$^{2+}$ doped Sr$_{1.48}$Ba$_0.5$Si(O,N)$_4$: 0.02Eu$^{2+}$ phosphor. As shown in the shaded area of Fig. 12, the substitution of Ba$^{2+}$ led to an obvious red-shift of the absorption band. The absorption band of Sr$_{1.5}$Si(O,N)$_4$ was located in ultraviolet region, which was range from ~265nm (4.7eV) to ~341nm (3.6eV). As a contrast, the doping of Eu$^{2+}$ significantly enhanced the absorption from the ultraviolet to blue light region which was caused by the typical 4f$^7$-4f$^6$5d$^1$ transition. These results indicated that Ba$^{2+}$ only could enhance the absorption in the ultraviolet region.

![Fig. 12](image)

**Fig. 12** Diffuse reflection spectra of Sr$_2$Si(O,N)$_6$, Sr$_{1.48}$Ba$_0.5$Si(O,N)$_4$ and Sr$_{1.48}$Ba$_0.5$Si(O,N)$_4$: Eu$^{2+}$.

Fig. 13 shows the energy level diagram of Ba$^{2+}$ and Eu$^{2+}$ in SBSON: 0.02Eu$^{2+}$. As shown in Fig. 8, whether the SBSON: Eu$^{2+}$ was excited by UV light or blue light, a large amount replacement of Ba$^{2+}$ (0.3 ≤ x ≤ 0.5) could lead to an obvious blue-shift phenomenon. Because the similar ionic radii between Sr$^{2+}$ (1.31 Å, 9CN; 1.36 Å, 10CN) and Eu$^{2+}$ (1.30 Å, 9CN; 1.35 Å, 10CN), and the Ba$^{2+}$ (1.49 Å, 9CN, 1.52 Å, 10CN) had larger ionic radii, the Eu$^{2+}$ would preferentially occupy Sr$^{2+}$ sites in SBSON. Thus, a small amount replacement of Ba$^{2+}$ (0.1 ≤ x ≤ 0.2) hardly led to an obvious blue-shift phenomenon. The Eu$^{2+}$ would gradually occupy Ba$^{2+}$ sites in SBSON host when a large amount of Ba$^{2+}$ (0.3 ≤ x ≤ 0.5) substituted for Sr$^{2+}$, which led to a obvious blue-shift. The blue-shift was related to the influence of Ba-content on the crystal-field strength and the covalence state. The crystal field splitting (Dq) can be determined by the following equation:

$$Dq = \frac{1}{6} Ze^2 \frac{r^4}{R^5}$$

where Dq is a measure of the energy level separation, Z is the anion charge, e is the electron charge, r is the radius of the d wavefunction, and R is the bond length. When larger Ba$^{2+}$ ions partially substituted for smaller Sr$^{2+}$ ions in the host lattice, bond length increased, which led to the lattice constants expansion of α'-SSON, as shown in Table 4. It caused weakening of crystal field strength. In addition, the electronegativity of Ba$^{2+}$ (0.89) is lower than that of Sr$^{2+}$ (0.95). With the increase of Ba$^{2+}$ content, the difference of electronegativity between cation and anion (electronegativity: N is 3.04, O is 3.44) would increase, which result in the covalency of the host lattice decreased. Thus, the crystal field strength became weak, which lead to an obvious blue-shift in the SBSON: Eu$^{2+}$. On the other hand, the alkaline earth ions form chains along the c axis in orthorhombic form of Sr$_2$SiO$_6$. The Eu$^{2+}$ randomly occupied Sr1 and Sr2 sites and form two luminescence centers Eu(1) and Eu(II) respectively. The positive charges can orient one d-orbital preferentially. As the length of the
c axis increases, the effect of preferential orientation of a d-orbital in the chain direction decreases so that the Eu$^{2+}$ emission shifts to shorter wavelength. Therefore, a blue-shift of the 5d-4f transition of Eu$^{2+}$ should occur.

Besides, according to the eqn(4) and Fig.3 (b) and (c), compared with the α′-SSO, the N$^{3-}$ doping lead to the shorter Sr2-O1 (3.105 Å vs. 2.881 Å, 3.105 Å vs 2.744 Å) bond length. Similarly with the Si-(N/O)$_4$ tetrahedron, the Sr2-O1 bond might be squeezed by the longer Sr-N bond (2.879 Å, 2.972 Å) because the N$^{3-}$ ions was simultaneously connected to Sr1 and Sr2 sites. The R value of Eu(ii) sites ligand get smaller. Therefore, the crystal-field strength is more influential on emission of the 9-oxygen coordinated Eu(II) sites than that of Eu(I) sites.31

For application in high power LEDs, the thermal stability of phosphors is one of the important issues to be considered. The temperature quenching performances of β-SSO: Eu$^{2+}$, β-SSON: Eu$^{2+}$ and α′-SBSON: Eu$^{2+}$ phosphors are shown in Fig. 14. As shown in Fig. 14, it could be observed that the thermal stability of α′-SBSON: Eu$^{2+}$ is superior to that of β-SSO: Eu$^{2+}$ and β-SSON: Eu$^{2+}$. In fact, according the previous discussion, the pure β-SSON: Eu$^{2+}$ phase wasn’t obtained in our experiments and it had some β-SSO: Eu$^{2+}$ impurity phases. This thermal phenomenon could be ascribed to the improvement of stability of α′ phase depending on the Ba$^{2+}$. The introduction of Ba$^{2+}$ led to a greater rigid lattice, which in turn results in the highest luminescence efficiency for intermediate compositions at elevated temperatures. As the Ba$^{2+}$ substitution for the Sr$^{2+}$ site would cause a longer and looser Ba-O compared with the Sr-O, Ba$^{2+}$ ions should preferentially coordinate the O$^{2-}$ and N$^{3-}$ in SBSON: Eu$^{2+}$. The partial Eu$^{2+}$ substituted for Ba$^{2+}$, which led to the result that above-average numbers of N$^{3-}$ and O$^{2-}$ was clustered surrounding Eu$^{2+}$. Therefore, the binding strength of Eu-O/N is slightly improved as partial Ba$^{2+}$ replacing Sr$^{2+}$, resulting in a large force constant and a low thermal quenching.54 Besides, the introduction of N could enhance the thermal stability. Compared with O, N have less lone pair electron and could form more chemical bond, which could lead to the result that more complex structure and the better thermal stability.

To determine the activation energy for thermal quenching, the Arrhenius equation was fitted to the thermal quenching data.67, 55, 56

\[
I(T) = \frac{I_0}{1 + c \exp(\frac{-E}{kT})}
\]

Where \(I_0\) is the initial intensity, \(I(T)\) is the intensity at a given temperature \(T\), \(c\) is a constant, \(E\) is the activation energy for thermal quenching, and \(k\) is Boltzmann’s constant. The curves in Fig. 14 were obtained by fitting according to the equation 5. The fitting results give a comparable activation energy \(E\) of 0.231 eV for α′-SSO: Eu$^{2+}$, 0.200 eV for β-SSON: Eu$^{2+}$ and 0.172 eV for β-SSO: Eu$^{2+}$. By increasing the temperature up to 150 °C at which the white LEDs usually work, the emission intensity of the α′-SSON: Eu$^{2+}$ phosphor remained at 78% of that measured at room temperature.

4 Conclusions

The Sr$_{1−x}$Ba$_x$Si(O,N)$_4$: 0.02Eu$^{2+}$ (0 ≤ x ≤ 0.5) phosphors were prepared by conventional solid state reaction method. The Sr-N and Si-N bond were confirmed by FT-IR spectra and crystal structure. Specially, compared with the Sr$_2$SiO$_4$: Eu$^{2+}$, the XRD refinement results indicated that N$^{3-}$ would substitute for O$^{2-}$ site and form Si-N-O3 tetrahedrons during the process of forming SSON: Eu$^{2+}$ substitutional solid solution. The SSON: Eu$^{2+}$ phosphors exhibited broad excitation spectra and a strong red emission ~ 610nm at Eu(II) sites. Ba$^{2+}$ doping could achieve the phase transformation from the β phase to α′-SBSON phase. The lattice constants were expanded with the introduction of Ba$^{2+}$, which could be attributed to the longer bond length of Ba-O. On the premise that the red emission peaks were not changed under 460nm excitation, the substitution of 0.1 mol Ba$^{2+}$ ions could enhance the PL intensity of SSON: Eu$^{2+}$. Whereas the substitution with 0.5 mol Ba$^{2+}$ ions could enhance the PL intensity of green light emission in SSON: Eu$^{2+}$. Both Ba$^{2+}$ and N$^{3-}$ doping could improve the thermal stability of Sr$_2$SiO$_4$: Eu$^{2+}$ phosphor. For the adjustable emission properties from green to red and the enhanced PL intensity by Ba$^{2+}$ doping in the SSON: Eu$^{2+}$ phosphor, it could be a good candidate for use in white light emitting diodes.

Acknowledgements

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Notes and references
graphical abstract
48x26mm (300 x 300 DPI)