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1. Introduction

Phosphor-converted white light-emitting diodes (w-LEDs) have been widely used as solid-state lighting for illumination and backlight for display.^{1,2} The properties of phosphors are crucial for the performance of LEDs, such as luminance efficacy, thermal stability, correlated color temperature (CCT), colorrendering index (CRI), and color gamut.^{3,4} Generally, LED with high CRI value and suitable CCT are needed for illumination. Phosphors with wide emission bands are beneficial to improve CRI value and adjust the CCT value for w-LEDs. Activators with parity-allowed transition are beneficial to improve the emission efficiency of phosphors, which in turn improve the luminance efficacy of w-LEDs. Eu^{2+} and Ce^{3+} ions are such kinds of activators with allowed 5d–4f transition and broad-band

emission, which are usually utilized as activators for phosphors applied in w-LEDs. $2,4-6$

The 5d orbitals of Eu^{2+} and Ce^{3+} ions are greatly influenced by the coordination environment. The energy difference between the lowest 5d excited level and the 4f ground level is adjustable, which in turn results in phosphors that can be excited efficiently by near-UV or blue light and exhibit colortunable emission. Therefore, a large number of Eu^{2+} or Ce^{3+} activated phosphors with tunable emission color have been reported on the basis of different crystal structures.⁷⁻¹⁷ The modification of crystal structures through the solid solution is an efficient way to adjust the luminescent properties of Eu^{2+} - or Ce3+-activated phosphors, which contains cationic replacement, (partial) anionic replacement, both cationic and anion group replacement, etc.⁶ These types of modification usually result in a continuous shift of the emission band, due to continuous variation of coordination polyhedron. There are other types of tuning emission colors for Eu^{2+} - or Ce^{3+} activated phosphors, which exhibit discrete variation of emission wavelength, *i.e.* crystal-site engineering approach and nano-segregation (or mixing of phases in the nano-domain region).18–20

 $LisrPO₄$ that can crystallize in a hexagonal (h) phase (PDF#14-0202), a monoclinic phase (PDF#53-1238), and a new

Tuning crystal structure and luminescence of Eu²⁺-activated LiSr_{1-x}Ba_xPO₄ solid solution for white light-emitting diodes†

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Emission spectral tuning is an important issue for phosphors. Herein, we report a Eu²⁺-activated solidsolution LiSr_{1-x}Ba_xPO₄ based on a recently discovered monoclinic LiSrPO₄, which has only one Sr²⁺ crystallographic site. Structural refinement indicates that the crystal structure can be maintained in the monoclinic phase from LiSr_{0.995}PO₄:0.005Eu²⁺ (x = 0) to LiSr_{0.095}Ba_{0.9}PO₄:0.005Eu²⁺ (x = 0.9). LiBa_{0.995}PO₄:0.005Eu²⁺ crystallizes in a trigonal phase, with different orientations of LiO₄ and PO₄ tetrahedrons from the monoclinic one. Both Sr^{2+} and Ba²⁺ are situated in the P₆Li₆ cage that is composed of six PO₄ and six LiO₄ tetrahedrons. However, different orientations of LiO₄ and PO₄ tetrahedrons results in different coordination situations for Sr^{2+} and Ba^{2+} , which leads to two different emission bands at 420 and 470 nm for Eu²⁺ activation. It is interesting that the change of Sr^{2+}/Ba^{2+} polyhedral size through solid solution does not result in a continuous shift of emission band from 420 to 470 nm, but the variation of intensity of two bands individually, which leads to the observation of both emission bands on LiSr_{0.695}Ba_{0.3}PO₄:0.005Eu²⁺. The emission intensity of the phosphors at 125 °C remains 75% of the value at room temperature, suggesting good thermal stability. The performance of phosphor-converted light-emitting diodes (pc-LEDs) indicates that phosphors can be used for near-UV chip-based white pc-LEDs. PAPER
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[†] Electronic supplementary information (ESI) available: Rietveld refinement, cell and atomic parameters, Sr/Ba/Eu–O distances, PL, XRD, spectra for QE, CIE versus current. See DOI: 10.1039/d1ma00950h

hexagonal phase (ICSD#258641) has been reported. Eu^{2+} -Activated phosphors with related structures were reported. $21-23$ Recently, we discovered a new monoclinic (m-)LiSrPO₄ with a higher symmetry than the old one.²⁴ A different Eu^{2+} -related luminescence was also observed. The crystal structure of the new m-LiSrPO₄ was refined based on the crystal structure of m-LiBaPO₄ reported by Kim et al.²⁵ The difference between crystal structures of new m-LiSrPO₄ and h-LiSrPO₄ is caused by the different orientations of LiO₄ and PO₄ tetrahedrons.²⁴ The crystal structure of LiBaPO₄ underwent monoclinic (m) \rightarrow trigonal (t) \rightarrow hexagonal (h) \rightarrow orthorhombic phase transitions upon heating from 298 to 1373 K, where the m \rightarrow t \rightarrow h transitions are displacive by the movement of polyhedrons without breaking chemical bonds.²⁵ Previous works on Eu²⁺-activated LiBaPO₄ are all based on the hexagonal phase.²⁶⁻³⁰

Since both LiSrPO₄ and LiBaPO₄ can crystallize in a monoclinic phase, a solid solution between them is expected to be obtained, which would influence the luminescence of Eu^{2+} activation. To the best of our knowledge, this is the first time to report on the evolution of Eu^{2+} -related emission on monoclinic $Li(Sr, Ba)PO₄$ solid solution. In the present work, solid solution within monoclinic phase can be realized from $Lisr_0$. $(PO_4):0.005Eu^{2+}$ to $Lisr_{0.095}Ba_{0.9}(PO_4):0.005Eu^{2+}$. While trigonal phase forms for the end compound $LIBa_{0.995}(PO_4):0.005Eu^{2+}$ without Sr^{2+} . Although there is only one crystallographic site for $Eu²⁺$ in these hosts, no continuous spectral shift is observed but a variation of two distinct emission bands upon increasing the $Ba²⁺$ content. The structure and luminescence of phosphors are studied in detail. Nanosegregation is proposed to explain the existence and intensity variation of the two emission bands.

2. Experimental

2.1. Materials and synthesis

 $\text{LiSr}_{0.995-x}\text{Ba}_x\text{PO}_4$:0.005Eu²⁺ phosphors were synthesized by using a conventional high-temperature solid-state reaction. Stoichiometric amounts of the raw materials $S₂$ (A.R.), BaCO₃ (A.R.), Li₂CO₃ (A.R.), NH₄H₂PO₄ (A.R.), and Eu₂O₃ (99.99%) were thoroughly mixed in an agate mortar. The wellmixed raw materials were sintered at 1000–1300 \degree C for 4 hours in a reductive atmosphere (a mixture containing 5% H₂ and 95% N_2 in volume percent). The synthetic process is similar to the previous work.²⁴

2.2. Measurements and characterization

X-Ray powder diffraction (XRD) patterns were measured using a Rigaku Ultima IV diffractometer (Cu Ka radiation (40 kV, 40 mA)). The XRD data for Rietveld refinements were collected under a slow scan speed in the range of 10° –120[°] by using a PANalytical X'pert Pro diffractometer equipped with a Cu K α radiation (40 kV, 40 mA). The refinements were achieved on a GSAS program. 31 The photoluminescence excitation (PLE), emission (PL) spectra, and temperature-dependent PL spectra were measured on a Hitachi F-4500 spectrophotometer with a TAP-02 temperature controller (Orient KOJI). The absolute quantum efficiency (QE), decay curves were measured using an Edinburg FLS980 fluorescence spectrometer equipped with a 450 W Xe lamp, a 320 nm-pulsed laser, and an integrating sphere. The pc-LEDs were fabricated based on a 1 watt 365 nm chip. The electroluminescent (EL) spectra of the pc-LEDs were collected on a highly accurate array spectrometer (HSP6000, HOPOO).

3. Results and discussion

XRD patterns of $Lisr_{0.995-x}Ba_xPO_4:0.005Eu^{2+}$ (x = 0-0.995) synthesized at 1200 °C are shown in Fig. 1. LiSr_{0.995}PO₄: $0.005Eu^{2+}$ (x = 0) contains the main phase that crystallizes in a new monoclinic (m-) phase as reported before, 24 and an impurity phase that can be indexed to m-LiSrPO₄ phase (PDF#53-1238) with a lower symmetry than the main phase. It should be noted that pure monoclinic $Lisr_{0.995}PO_4:0.005Eu^{2+}$ can be achieved under sintering in 1000 and 1100 \degree C. The diffraction peaks exhibit an obvious shift to the small angle side with the increase of Ba content (x) . The impurity phase disappears with the introduction of Ba. The magnified diffraction peaks at around $2\theta = 30^\circ$ (Fig. 1b) show not only the shift of peaks but also the tendency of getting closer and finally becoming a single peak for $x = 0.995$ without Sr atoms. This phenomenon suggests the variation of cell parameters and interplanar spacing and the change of crystal structure at $x = 0.995$. Kim *et al.* reported that a slowly-cooled (-5 K h^{-1}) sample crystallized in a monoclinic LiBaPO₄, while the quenched sample is trigonal (t-)LiBaPO₄.²⁵ Our sample was furnace cooled, which had a much higher speed than $5 K h^{-1}$. Therefore, t-LiBa_{0.995}PO₄:0.005Eu²⁺ was reasonable. On the other hand, the t-LiSrPO₄ phase is not reported till now. The existence of Sr^{2+} may have great influence on the phase, therefore, all ${\rm LiSr_{0.995-x}Ba_xPO_4:0.005Eu^{2+}}$ $(x = 0.1-0.9)$ samples synthesized at 1200 \degree C have the same phase as monoclinic $LiSr_{0.995}PO₄:0.005Eu²⁺.$ Puper

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Fig. 1 (a) XRD patterns of LiSr_{0.995-x}Ba_xPO₄:0.005Eu²⁺ synthesized at 1200 °C with different Ba²⁺ concentrations (x), (b) magnified XRD patterns in the 28° –31° range. Peaks assigned by arrows: impurity m-LiSrPO₄ phase (PDF#53-1238).

Fig. 2 Rietveld refinement of LiSr_{0.995-x}Ba_xPO₄:0.005Eu²⁺ synthesized at 1200 °C. (a) x = 0.3, (b) x = 0.7, (c) x = 0.995, (d) cell parameters versus Ba content.

Rietveld refinements were conducted to further study the variation of crystal structure in detail. The crystal data of monoclinic LiBaPO₄ reported by Kim et al. were used as a starting model for refinement of samples with x equaling 0–0.9.²⁵ While the refinement for $x = 0.995$ was based on the crystal structure of trigonal (t-)LiBaPO₄.²⁵ Fig. 2a–c illustrate refinement results of representative samples, namely $x = 0.3$, 0.7, and 0.995 that were obtained at 1200 $^{\circ}$ C. The refinement result for $x = 0$ can be found in our previous work,²⁴ and the results of others are shown in the ESI† as Fig. S1. The calculated values match well with the observed data. Fig. 2d shows the cell parameters versus Ba content (x) . The cell lengths and cell volume tend to increase with Ba content. It should be noted that sample $x = 0.995$ crystallizes in a different phase compared to others, and has a half number of chemical formulas in a unit cell than others. Therefore, the two times of cell volume for $x = 0.995$ is larger than the volume of others. Detailed refinement results and crystallographic data are listed in Tables S1– S8 in the ESI.† Fig. 3 illustrates the crystal structures of representative samples viewed along the c axis. Three PO₄ and three $LiO₄$ tetrahedrons connect alternately and form a six-member ring. Two parallel rings are also connected by sharing three oxygen atoms, which form a P_6Li_6 cage. A Sr, Ba, or Eu atom is situated at the center of the cage. Fig. 3 shows that the degree of dislocation between the two parallel sixmember rings decreases upon increasing the Ba content. Fig. 4 exhibits the difference in the positions of tetrahedrons. Samples with $x = 0$ and 0.1 have similar relative positions of PO₄ and LiO₄, while those with $x = 0.3 - 0.995$ have similar relative positions. The difference in the relative positions of $PO₄$ and $LiO₄$ tetrahedrons will result in the variation of coordination polyhedrons for Sr, Ba, and Eu. Table S9 (ESI†) lists Sr/Ba/Eu–O

Fig. 3 Crystal structure of (a) m-LiSr_{0.995}PO₄:0.005Eu²⁺, (b) m-LiSr_{0.695}Ba_{0.3}PO₄: $0.005Eu^{2+}$, (c) m-LiSr_{0.295}Ba_{0.7}PO₄:0.005Eu²⁺, and (d) t-LiBa_{0.995}PO₄ 0.005Eu^{2+} from the refinement results.

and P–O distances obtained from refinement. The average distance of the nine-coordinated polyhedrons tends to increase with Ba content, which is in accordance with the cell volume. However, it should be noted that there is an abnormally long distance for Sr-rich samples as pointed out in Fig. 4, which is due to the relatively smaller cationic size of the P_6Li_6 cage.

The relative energy of 5d levels of Eu^{2+} depends on the coordination environment. Therefore, PLE and PL spectra are expected to be tunable by changing the x value in $LiSr_{0.995-x}Ba_xPO₄:0.005Eu²⁺$. Fig. 5a and b show PLE and PL

Fig. 4 Comparison of MO₉, PO₄, and LiO₄ polyhedrons in the crystal structures. (a) m-LiSr_{0.995}PO₄:0.005Eu²⁺, (b) m-LiSr_{0.695}Ba_{0.3}PO₄:0.005Eu²⁺, (c) m-LiSr_{0.295}Ba_{0.7}PO₄:0.005Eu²⁺, and (d) t-LiBa_{0.995}PO₄:0.005Eu²⁺ from the refinement results.

spectra of $Lisr_{0.995-x}Ba_xPO_4:0.005Eu^{2+}$ synthesized at 1200 °C. The PLE spectra of all the phosphors cover the 250–420 nm region with a maximum at around 365 nm, which indicates that all of the phosphors can be excited by near UV light. The profiles of PLE spectra for $x = 0$ and 0.1 are different from those for $x = 0.3-0.995$, which relate to the difference in coordination environment for Eu^{2+} . Under excitation at 365 nm, phosphors with $x = 0$ and 0.1 exhibit a PL band at 420 nm, while those with $x = 0.5 - 0.995$ have an asymmetric PL band at 470 nm. Phosphor with $x = 0.3$ possesses both PL bands at 420 and 470 nm. Fig. 5c shows the PL intensity of the two bands versus Ba content (x) , which shows a rapid decrease for the 420 nm band from $x = 0$ to 0.3 and a slight increase of the 470 nm band from $x = 0.3$ to 0.995. Fig. S2 and S3 (ESI†) show PL spectra and XRD patterns of $Lisr_{0.995-x}Ba_{x}PO_{4}:0.005Eu^{2+}$ synthesized at 1000-1300 $^{\circ}$ C, respectively. Upon changing the Ba content, similar variations in PL bands and XRD peaks compared to those obtained at 1200 \degree C were observed. The different PL profile for LiSr_{0.995}PO₄:0.005Eu²⁺ (x = 0) obtained at 1300 \degree C is due to the formation of the different crystal structure (m-LiSrPO4, PDF#53-1238), which has been discussed in our previous work.²⁴

Generally, there are two types of variation for the PL band of Eu^{2+}/Ce^{3+} -activated solid solutions, where the cations occupied by an activator(s) change. The first one is a continuous shift of the PL band if the solid solution leads to the variation of a same crystallographic site occupied by Eu^{2+} or Ce^{3+} , e.g. $Ba_{2-x}Sr_{x}$ SiO_4 : $\text{Ce}^{3+13}\text{Sr}_{1-x}\text{Ba}_x\text{Si}_2\text{O}_2\text{N}_2$: Eu^{2+} (x = 0-0.75),³² Ca_{1-x}Sr_xHf₄ $(PO₄)₆: Eu²⁺,³³ etc. The second one is a discontinuous shift of$ the PL band if the solid solution leads to the variation of occupancy for Eu^{2+} or Ce^{3+} on more than one crystallographic sites, e.g. $\text{Li}_4\text{Sr}_{1+x}\text{Ca}_{1-x}(\text{SiO}_4)_2:\text{Ce}^{3+}$, 18 CsKNa_{2-y}Li_y(Li₃SiO₄)₄:Eu²⁺ $(y = 0-1)$,³⁴ etc. It is interesting that no continuous shift of PL band is observed in $Lisr_{0.995-x}Ba_xPO_4:0.005Eu^{2+}$ series by changing the Ba/Sr ratio, where there is only one Ba/Sr crystallographic site as indicated by the refinement result. It is speculated that environments of $SrO₉$ and Ba $O₉$ in $\text{LiSr}_{0.995-x}\text{Ba}_x\text{PO}_4$:0.005Eu²⁺ (x = 0.1-0.9) are similar to those in LiSr_{0.995}PO₄:0.005Eu²⁺ (x = 0) and LiBa_{0.995}PO₄:0.005Eu²⁺ $(x = 0.995)$, respectively, based on the variation of PL bands **Publish Article article is article in the common article is article in the state of the state of 1/27/2023 120 AM. The state of t**

Fig. 5 PLE and PL spectra of LiSr_{0.995-x}Ba_xPO₄:0.005Eu²⁺ synthesized at 1200 °C, (a) PLE monitored at 420 or 470 nm, (b) PL excited at 365 nm, (c) PL intensity versus Ba content, (d) normalized PLE spectra monitored at two different wavelengths for $x = 0.3$ and 0.995

and relative PL intensities, although refinement suggests only one Ba/Sr crystallographic site. $SrO₉$ is dominant in samples with $x = 0$ and 0.1, which results in a 420 nm PL band after Eu²⁺ substitution. While Eu^{2+} tends to occupy the BaO₉ site for samples with $x = 0.5 - 0.995$, which results in a 470 nm PL band. Eu^{2+} ions situate on both SrO₉ and BaO₉ in the sample with $x = 0.3$; therefore, two PL bands are observed. This phenomenon is similar to that in Eu²⁺-doped $(CaMg)_x(NaSc)_{1-x}Si_2O_6$ solid solution, where nano-segregation occurred and allowed predictive control of the relative PL intensity of the two bands with the same profile and wavelength center to the end compounds.¹⁹ Therefore, the variation of PL spectra indicates that nano-segregation also happens in the $Lisr_{0.995-x}Ba_{x-x}$ $PO_{\scriptscriptstyle{4}}\textrm{:}0.005 \textrm{Eu}^{2+}$ solid solution. Materials Advances

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The relationship between emission peak and site coordination environment can be discussed by comparing the crystal field strength (calculation of crystal field splitting), centroid shift of 5d levels for Eu^{2+} , and Stokes shift for Eu^{2+} on Sr^{2+} and Ba²⁺ sites. The average $\text{Sr}^{2+}/\text{Ba}^{2+}/\text{Eu}^{2+}-\text{O}$ distances (R_{av}) are 2.751 and 2.899 Å in $Lisr_{0.995}PO_4:0.005Eu^{2+}$ and $Liba_{0.995}$ $PO₄:0.005Eu²⁺$, respectively, according to Table S9 (ESI†). The crystal field splitting $(\varepsilon_{\text{cfs}}/eV)$ can be calculated according to the following equation, 35,36

$$
\varepsilon_{\rm cfs} = \beta_{\rm poly}^Q R_{\rm av}^{-2} \tag{1}
$$

Both Sr^{2+} and Ba^{2+} are coordinated by nine O atoms, therefore, Eu^{2+} on these two sites should have the same β_{poly}^Q value. Eu^{2+} on the Sr²⁺ site with smaller R_{av} has a higher ε_{cfs} value than that on the Ba²⁺ site. The centroid shift (ε_c /eV) of 5d levels for Eu²⁺ can be obtained using the following equation, $35,36$

$$
\varepsilon_{\rm c} = 1.79 \times 10^{13} \sum_{i=1}^{N} \frac{\alpha_{\rm sp}^{i}}{(R_i - 0.6\Delta R)^6}
$$
 (2)

Most Ba–O bonds are longer than those of Sr–O according to Table S9 (ESI†). The calculated ε_c for Eu²⁺ on Sr²⁺ site is larger than that on Ba²⁺ site. The combination of ε_c and ε_{cfs} suggests that Eu^{2+} on the Sr^{2+} site should have a smaller energy difference between the lowest 5d level and 4f ground state than Eu^{2+} on the Ba^{2+} site, which corresponds to a longer excitation wavelength of the lowest PLE band for Eu^{2+} on the Sr^{2+} site. Fig. 5a and d show the excitation spectra monitored at 420 and 470 nm, which is in accordance with the calculation results. The Stokes shift is defined as the energy difference between the peak values of PLE and PL bands. Therefore, it is obvious that the Stokes shift of Eu^{2+} on the Ba^{2+} site is much higher than that on the Sr^{2+} site. Therefore, a huge different Stokes shift of Eu^{2+} on Sr^{2+} and Ba^{2+} sites is the main reason for the difference of emission peaks for Eu^{2+} on the two sites. Fig. 6 illustrates the difference for Eu²⁺ ions on Sr²⁺ and Ba²⁺ related sites in the configurational coordinate diagram. For Ba polyhedrons related emission, the excited state of Eu^{2+} suffers a larger offset (ΔR) of the parabola, therefore, resulting in a larger Stokes shift and longer emission wavelength. The PLE bands monitored at 470 and 500 nm for $x = 0.995$ also almost overlap, which is consistent, with that, there is only one Ba site in trigonal

Fig. 6 Configuration coordinate diagram for Eu^{2+} ions on (a) Sr and (b) Ba polyhedrons, illustrating the origin of similar PLE bands but different PL bands.

 $LiBaPO₄$. The m-, t-, and even h-phase differ in the orientation of LiO₄ and PO₄, which results in the variation of $Sr/BaO₉$ polyhedrons. The change of Ba/Sr ratio is easy to cause the variation of Sr/BaO₉ polyhedrons. Therefore, the unusual asymmetric PL band at 470 nm may originate from Eu^{2+} ion on Ba^{2+} sites with a slight difference among themselves. Fig. 7 plots the decay curves of samples monitored at 420 and 470 nm. The different decay behaviors for 420 and 470 nm related emissions for $x = 0.3$ again suggest the different coordinate environments. The decay monitored at 420 nm slows down with the increase of Ba²⁺ content from $x = 0$ to 0.3, which suggests that the change of chemical composition and the decrease of Eu^{2+} occupancy on Sr^{2+} -related site lead to the variation in the nonradiative transition rate.

The internal quantum efficiency (QE) of $\rm{Lisr_{0.995-x}Ba_{x}PO_{4}}$ $0.005Eu^{2+}$ obtained at 1200 °C was estimated to be about 76.08%, 60.10%, 68.47% and 75.34% for x = 0, 0.3, 0.9, and 0.994, respectively. The QE values of the end compounds are higher than those of solid-solution compounds. Related spectra for QE measurements are shown in Fig. S4 (ESI†). The relationship between QE and decay time can be expressed as 13,37

$$
QE = \tau/\tau_0 \tag{3}
$$

where τ and τ_0 are the decay times with and without nonradiative transition, respectively. The increase of lifetime for 470 nm emission as indicated in Fig. 7b is consistent with the increase of QE value from $x = 0.3$ to 0.995, which suggests the decrease of nonradiative rate.

Fig. 7 Decay curves of LSBPO:0.005Eu²⁺, (a) monitored at 420 nm for $x = 0 - 0.1$ and at both 420 and 470 nm for $x = 0.3$, (b) monitored at 470 nm for $x = 0.3 - 0.995$

Fig. 8 Temperature-dependent PL spectra of LSBPO:0.005Eu²⁺, (a) $x = 0$, (b) $x = 0.5$, (c) $x = 0.995$, and (d) PL intensity versus temperature with fitted E_a values.

Thermal stability is also an important parameter for phosphors used in pc-LEDs. The temperature-dependent PL spectra of representative samples are shown in Fig. 8a–c. The PL intensity tends to decrease with increasing temperature from room temperature to 200 °C for all samples. Fig. 8d plots the PL intensity versus temperature. It is obvious that the PL intensity of sample $x = 0.995$ drops slowly from 25 to 100 °C and then faster with a further increase in temperature. While the PL

intensity of samples $x = 0$ and 0.5 decreases smoother than that of $x = 0.995$ at higher temperatures. The PL intensity of the samples at 125 \degree C remains at 75% of the initial value at room temperature. The fitted curves in Fig. 8d are achieved by using the Arrhenius formula,³⁸

$$
\frac{I_T}{I_0} = \left[1 + D \exp\left(\frac{-E_a}{kT}\right)\right]^{-1} \tag{4}
$$

Fig. 9 EL spectra of white pc-LEDs based on LSBPO:0.005Eu²⁺ phosphors combined with 365 nm chip and commercial green and red phosphors, (a) $x = 0.3$, (b) $x = 0.9$, (c) $x = 0.995$, and (d) CIE coordinates of the pc-LEDs under different currents.

where I_T and I_0 are the PL intensity at temperature T (in K) and 0 K, respectively. E_a is the activation energy for thermal quenching. D is a constant depending on phosphor, and k is the Boltzmann constant. The fitted E_a values for $x = 0$, 0.5, and 0.995 are 0.17, 0.32, and 0.53 eV, respectively. A higher E_a value means that it needs more energy for the excited electrons to return to the ground state nonradiatively. The calculated E_a values are in accordance with the temperature-dependent PL intensity below 125 \degree C. Furthermore, the PL peak of the samples change little upon increasing temperature. The stability of both PL intensity and peak suggests that LSBPO:0.005Eu²⁺ phosphors could be suitable for pc-LEDs.

A series of white-emitting pc-LEDs were fabricated by combining LSBPO:0.005Eu²⁺ phosphors and commercial greenemitting $(Ba, Sr)_2SiO_4:Eu^{2+}$, red-emitting CaSiAlN₃:Eu²⁺ with a 365 nm n-UV chip. The electroluminescent (EL) spectra of the pc-LEDs under 50–350 mA forward bias currents are shown in Fig. 9a–c. The performance of the pc-LED based on $LisrPO₄$: $0.005Eu^{2+}$ (x = 0) can be found in our previous work, which contains a gap around the cyan-emission region. However, pc-LEDs based on samples $x = 0.3, 0.9$, and 0.995 can overcome this drawback as these phosphors show emission bands covering the cyan region. All the pc-LEDs have a color-rendering index (R_a) value higher than 90 and a warm white emission with correlated color temperature (CCT) at 3000–4000 K. The EL intensities of pc-LEDs tends to increase with working current, and the EL profile changes little. Luminance efficacies (LE) of the pc-LEDs at 50 mA are shown in Fig. 9. It is believed that the LE values can be improved by optimizing the parameters for pc-LED fabrication, for example, the weight ratio of phosphor to silica gel. Fig. 9d shows the CIE coordinates of the pc-LEDs under different current, detailed CIE values versus current are plotted in Fig. S5 (ESI†). These results suggest that the LSBPO:0.005Eu²⁺ phosphors can be candidates as blueemitting component for pc-LEDs based on the n-UV chip. Materials Advances

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Conclusions

In summary, emission-tunable Eu^{2+} -activated $Lisr_{1-x}Ba_xPO_4$ solid solution was obtained by a high-temperature solid-state reaction. The crystal structure of a recently refined monoclinic phase can be maintained in $\textrm{Lisr}_{0.995-x}\textrm{Ba}_x\textrm{PO}_4:0.005\textrm{Eu}^{2+}$ for x in 0–0.9 range. While, it crystallizes in a trigonal phase for $LiBa_{0.995}PO₄:0.005Eu²⁺$ ($x = 0.995$). The end compounds $LiSr_{0.995}PO_{4}:0.005Eu^{2+}$ and $LiBa_{0.995}PO_{4}:0.005Eu^{2+}$ exhibit broad emission bands at 420 and 470 nm, respectively. However, no continuous shift of the emission band is observed in the solid solution, although there is only one crystallographic Sr or Ba site in the crystal structure. The formation of nanosegregation is proposed to explain the variation of emission upon changing Ba/Sr ratio in $\textrm{Lisr}_{0.995-x}\textrm{Ba}_{x}\textrm{PO}_{4}:0.005\textrm{Eu}^{2+}.$ The work provides an example of controlled photoluminescence tuning with a discontinuous shift of the emission band. Thermal stability and performance of pc-LEDs suggest that the

phosphors can be a candidate as phosphors for solid-state lighting based on n-UV LED.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

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