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Cation substitution induced novel gehlenite Ca₂GaAlSiO₇:Eu²⁺/Ce³⁺ phosphor with green/blue emission for UV-WLEDs†

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By partially substituting Ga^{3+} ions at Al1(2a) and Al2/Si1(4e) sites in gehlenite $Ca_2Al_2SiO_7$, novel $Ca_2GaAlSiO_7$:Eu²⁺/Ce³⁺ phosphors with efficient green and blue emissions have been prepared for the first time. Structure and luminescence properties of these newly obtained samples have been investigated in detail together with the effect of Ga^{3+} substitution which has caused a decrement of Eu–O bond length and covalency. The excitation spectra suggested that both the Eu²⁺ and Ce^{3+} doped $Ca_2GaAlSiO_7$ samples had broad excitation bands in the UV range, which are favourable properties for application as UV-WLED phosphors. Under 350 nm irradiation, the $Ca_2GaAlSiO_7$:Eu²⁺ had intense emission peaking at 510 nm, while $Ca_2GaAlSiO_7$:Ce³⁺ emitted bright blue light. The energy migration mechanism of the Eu²⁺ ions was confirmed to be electric dipole–dipole interaction in terms of the experimental results and analysis. The thermal quenching mechanism, and quantum efficiency of the composition-optimized samples have been investigated. Moreover, a white LED device has been obtained by coating our prepared phosphors with $CaAlSiN_3$ red phosphor on a UV chip. The experimental results suggested that the $Ca_2GaAlSiO_7$:Eu²⁺ and $Ca_2GaAlSiO_7$:Ce³⁺ samples might have potential value in serving as green and blue phosphors for UV-WLEDs, respectively.

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1. Introduction

In recent decades, the environmental and energy issues have become more and more severe, which make the population of WLEDs (white light emitting diodes) become urgent since they have many advantages over traditional lighting, for example, good reliability, high brightness and long service time. 1-5 Currently, a phosphor-converted WLED is the main approach to realize white light, in which the phosphor serves as the key material for governing the colour characteristics.⁶⁻⁹ In the commercial market, the most popular fabrication of WLEDs is InGaN blue-emitting chips, coating with yellow-emitting YAG:Ce. However, this combination faces the problems of low colour rendering index (CRI < 75) and high correlated colour temperature (CCT \approx 6000 K) resulting from the shortage of emission in the red region, which has a restriction on application in medical and indoor lighting. Researchers have made many efforts to solve this problem, such as improving the red component of YAG:Ce phosphor10,11 and developing novel red

Novel emission phosphors can be sought in a variety ways, *e.g.*, exploring new host through single-particle-diagnosis approach, tutilizing energy transfer between different dopants, designing isostructural solid solutions, and adjusting the metal-ligand hybridization and crystal field splitting by cationic/anionic substitutions. The ionic substitution of host frequently employed in Ce³⁺ and Eu²⁺ ions doped phosphors since the energy levels of outmost 5d orbits are easily affected by the coordination environment. Moreover, the spinand orbit-allowed electron transition results in intense luminescence intensity which is one of the key factors for application. Li *et al.* had obtained different emission colours with Ba²⁺ substituting for Sr²⁺ in Sr₂Si₂O₂N₂:Eu.²¹ Won Bin Im had reported green phosphors with different CIE coordinates by

phosphors (Sr[Mg₃SiN₄]:Eu²⁺,¹² Sr[LiAl₃N₄]:Eu²⁺,¹³ K₂SiF₆:Mn⁴⁺ (ref. 14)) which can mix with YAG:Ce. Nowadays, with the development of chips, the realization of white light by UV-A (315–400 nm) chips coating with tri-colour phosphors was expected to dominate the market since the improved CRI can make the light be warmer and more similar to natural light.^{15,16} Moreover, since the UV light is invisible to naked eye, the performance of WLEDs in this combination strongly depends on the emission spectra of phosphors, making the luminescence properties easy to control. In this way, it is urgently needed to search for new phosphors matching well with the commercial UV chips to meet the requirement of warm white light.

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substituting Ba²⁺ for Sr²⁺ ion in Sr₃Ce_{0.025}AlO₄F.²² Among many kinds of inorganic materials, aluminosilicate-based phosphors have been investigated due to their relatively easy preparation, structural diversity, and good thermal stability. In 1982, M. Kimata investigated the structure of Ca2Al2SiO7 gehlenite in detail.23 Qiu and Wang et al. reported luminescence property of Eu²⁺ and Ce³⁺/Tb³⁺ ion in this host, respectively. ^{24,25} Wang et al. obtained yellowish green emission by substituting Sr²⁺ for Ca²⁺ ions Ca₂Al₂SiO₇:Eu²⁺.26 Xia et al. obtained solid solution by substituting Mg²⁺-Si⁴⁺ for Al³⁺-Al³⁺ in Ca₂Al₂SiO₇.²⁷ As we know that the Ga³⁺ ion has similar property with Al³⁺ due to their same valence states and outmost electron configurations. The Ga³⁺ ions can be used to tune the emission wavelength and improve the luminescence performance of phosphors, for example, the commercial green Y₃(Al,Ga)₅O₁₂:Ce³⁺ and Y₃-(Al,Ga)₅O₁₂:Tb³⁺ phosphors. Moreover, the gallate based oxides had been proved to be good host materials for phosphor. 28,29 In this work, by partially substituting Ga^{3+} for Al^{3+} ions, we firstly obtained the Ca₂GaAlSiO₇ which can be expressed as Ca₂Al_{0.5}-Ga_{0.5}(Al_{0.5}Ga_{0.5}Si)O₇ according to their wyckoff positions. The structure and luminescence properties of Eu2+ and Ce3+ ions in the host had been investigated as well as the effect of Ga³⁺ substitution. The performance of our prepared phosphors has great attraction for UV-WLEDs.

2. Experimental section

2.1 Starting materials and synthesis

Series of $Ca_2GaAlSiO_7:xEu^{2^+}/yCe^{3^+},yLi^+$ (CGASO: $xEu^{2^+}/yCe^{3^+},yLi^+$) samples had been prepared via common solid-state reactions. The starting materials $CaCO_3$ (A.R.), Ga_2O_3 (A.R.), Al_2O_3 (A.R.), SiO_2 (A.R.), $Li_2CO_3(A.R.)$, CeO_2 (99.99%) and Eu_2O_3 (99.99%) were firstly weighed on analytical balance in accordance with the molar ratio of each element in the formula. Since the valence states between Ca^{2^+} and Ce^{3^+} are different, the addition of charge compensators becomes necessary. Thus, the Li_2CO_3 providing Li^+ ions were added during the preparation. Raw materials with proper ratio were well mixed by thoroughly grinding for 15 min. After that, the mixtures were sintered at 1300 °C for 3 h under reductive atmosphere (20% H_2 + 80% N_2) to form the as-prepared phosphors.

2.2 Characterization techniques

Phase identification of our as-prepared phosphors was confirmed by powder XRD analysis operated on AXS D8 (Bruker) with radiation of CuK α ($\lambda=0.15405$ nm) at 40 kV and 40 mA. The photoluminescence emission and excitation spectra were collected on Hitachi fluorescence spectrophotometer (F-7000) with excitation source being 150 W xenon lamp. Diffuse reflectance spectra measurement was performed on Hitachi spectrophotometer (U-4100-UV/Vis) using BaSO₄ as standard reference. For the luminescence dynamic measurement, the signals were collected on digital oscilloscope (Lecroy Wave Runner 6100, 1 GHz) with tunable laser serving as excitation source (pulse width: 4 ns, gate: 50 ns). Photoluminescence quantum yield (QY) of our prepared samples was collected on

C9920-02 absolute quantum yield system (Hamamatsu Photonics K.K., Japan). The spectra at different temperatures were operated on Edinburgh Instrument spectrophotometer (FLS 920) equipped with temperature controller. The EL spectrum, CIE coordinates, colour rendering index (CRI), and colour temperature (CCT) of the fabricated LED device were measured by Starspec SSP6612.

Results and discussion

3.1 Phase analysis

Phase purity of our prepared phosphors is checked by powder Xray diffraction and the typical patterns have been shown in Fig. 1. The diffraction peaks of Ca₂GaAlSiO₇ host and rare earth ions doped samples match well with the standard data of Ca₂-Al₂SiO₇ (JCPDS#35-0755) which is shown as a comparison. The results reveal that the as-prepared samples are single phase and have same structure with Ca2Al2SiO7. Since the ionic radius of Ga^{3+} (r = 0.47 Å, CN = 4) is much larger than that of Al^{3+} ion (r = 0.47 Å, CN = 4) is much larger than that of Al^{3+} ion (r = 0.47 Å, CN = 4) is much larger than that of Al^{3+} ion (r = 0.47 Å, CN = 4) is much larger than that of Al^{3+} ion (r = 0.47 Å, CN = 4) is much larger than that of Al^{3+} ion (R = 0.47 Å). 0.39 Å, CN = 4), the substitution of Ga^{3+} has caused the diffraction peaks shift to lower angle as can be seen in Fig. S1 (ESI†). Considering the similar ionic radii, the doped rare earth ions are thought to occupy the site of Ca²⁺ ions. Due to the different valence between Ce³⁺ and Ca²⁺, the doping of Ce³⁺ will lead to positive charges in the host, which can affect stability and luminescence intensity of the phosphor. To solve this problem, the Li⁺ ions serving as charge compensators were added along with Ce3+ ions. In the host, one Ce3+ ion and one Li⁺ ion are considered to replace two Ca²⁺ ions, which can be demonstrated as $2V_{Ca}''+Ce^{3+}+Li^{+}\!\rightarrow\!Ce_{Ca}^{\textstyle \cdot}+Li_{Ca}'.$ From Fig. 1, we can see the Ca²⁺ sites in the lattice are substituted by the doped ions without causing any impurity.

Rietveld structure refinement of the $Ca_2GaAlSiO_7$ host has been conducted using the crystal structure parameters of Ca_2 - Al_2SiO_7 as initial structure model. The refinement results are listed in Table 1 and the refined XRD profiles together with

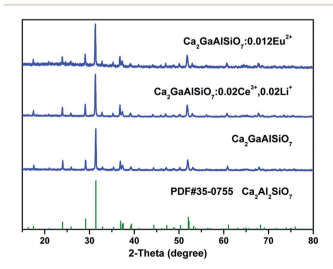


Fig. 1 Powder X-ray diffraction patterns of typical prepared samples. The standard data of $Ca_2Al_2SiO_7$ (JCPDS#35-0755) are shown as references.

Table 1 Final refined structure parameters of CGASO host derived from the Rietveld refinement of X-ray diffraction data^a

Atom	Site	x	у	z	Frac	Uiso
Ca1	4e	0.3397(8)	0.1602(2)	0.5092(5)	1.00	0.0119(7)
Ga1	2a	0.00000	0.00000	0.0000	0.50	0.02500
Ga2	4e	0.14310	0.35690	0.95630	0.25	0.02500
Al1	2a	0.00000	0.00000	0.00000	0.5	0.02500
Al2	4e	0.14310	0.35690	0.95630	0.25	0.02500
Si1	4e	0.14310	0.35690	0.95630	0.5	0.02500
O1	2c	0.50000	0.00000	0.17590	1.00	0.02500
O2	4e	0.1477(7)	0.3523(7)	0.2863(9)	1.00	0.0164(2)
О3	8f	0.0881(6)	0.1670(6)	0.7823(8)	1.00	0.0121(1)

^a Cell parameters: $a = 7.7441(2) \text{ Å}, b = 7.7441(2) \text{ Å}, c = 5.1039(2) \text{ Å}, \alpha =$ $\beta=\gamma=90^\circ,\,V=306.08(1)\,\text{Å}^3;\,Z=2;\,\text{space group:}\,P\bar{4}2_1m$ (113); reliability factors: $\chi^2 = 2.649$, $R_{\rm wp} = 7.35\%$, $R_{\rm p} = 5.63\%$ Average {Ca}-O bond length: 2.548(6) Å.

crystal structures are demonstrates Fig. 2. The Ca₂GaAlSiO₇ host possesses tetrahedral crystal system and $P\bar{4}2_1m$ (113) space group. In the lattice, the Ca²⁺ site is coordinated by eight oxygen ions forming polyhedron, which can be substituted by Eu²⁺/Ce³⁺ ion forming emission centre. The Ga³⁺, Al³⁺ and Si⁴⁺ cations are 4-fold coordinated and form [GaO₄], [AlO₄] and [SiO₄] tetrahedrons, respectively. These tetrahedrons are further connected by sharing oxygen atoms. The 2a sites are occupied by Ga³⁺ and Al³⁺ ions with the ratio of 1:1, while the 4e sites are occupied with $Ga^{3+}:Al^{3+}:Si^{4+}=1:1:2$. The fit quality of this refinement is pretty good which can be reflected from reliability results $\chi^2 = 2.649$, $R_{\rm wp} = 7.35\%$, $R_{\rm p} = 5.63\%$. The cell parameters of our prepared $Ca_2GaAlSiO_7$ are determined to be a = b =

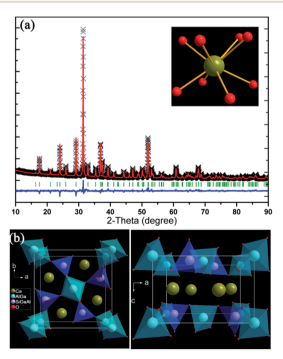


Fig. 2 (a) Rietveld refinement pattern of CGASO host and the coordinate environment of Ca²⁺ ion. (b) Unit cell structure of CGASO viewing along c and b direction.

7.7441(2) Å, c = 5.1039(2) Å and V = 306.08(1) Å³, while the standard lattice parameters of Ca₂Al₂SiO₇ reported by M. Kimata are a = b = 7.6770(3) Å, c = 5.0594(3) Å and V =298.18(2) Å³. The expansion of cell volume for as-prepared Ca₂GaAlSiO₇ can be ascribed to large radius of Ga³⁺ than Al³⁺ ions. Moreover, the average length of Ca-O bond was determined to be 2.548(6) Å in Ca₂GaAlSiO₇ host, which is shorter than that in Ca₂Al₂SiO₇ (Ca-O: 2.563 Å). This phenomenon can be ascribed to that the substitution of Ga³⁺ for Al³⁺ ions will increase the lattice strain due to the cation size mismatch. And thus, the anion clusters in the host will reduce this strain by shorting the cation-anion (Ca-O) distance.30

3.2 Photoluminescence properties of CGASO:Eu²⁺ and CGASO:Ce3+,Li+ phosphors

Fig. 3 depicts reflection spectra of our typical prepared samples. For the CGASO host, there is a sharp drop under 230 nm caused by electron transition from valence band (VB) to conduction band (CB) of the host. For rare earth ions doped samples, the reflection spectra exhibit not only the host absorption but also absorption bands arising from $4f \rightarrow 5d$ electronic transition of Eu²⁺ and Ce³⁺ ions. The absorption band of CGASO:0.012Eu²⁺ covers the UV and blue region (250-480 nm), while the absorption band of CGASO:0.02Ce3+,0.02Li+ is in the range of 250-400 nm with several peaks. By utilizing the following equation, an approximate band gap value can be obtained:31

$$[F(R_{\infty})h\nu]^n = A(h\nu - E_{\mathfrak{g}}) \tag{1}$$

where $h\nu$ represents the photon energy; $E_{\rm g}$ is the optical band gap value; n = 1/2 and 2 corresponds for indirect and direct transition, respectively; A stands for a constant; $F(R_{\infty})$ stands for Kubelka-Munk function reported by Kubelka and Munk in 1931, which can be written as32

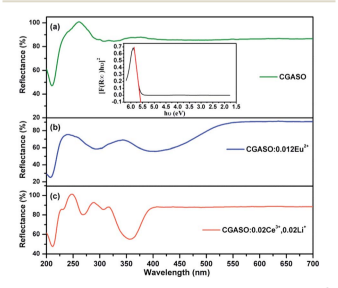


Fig. 3 Diffuse reflection spectra of (a) CGASO, (b) CGASO:0.012Eu²⁺ and (c) CGASO:0.02Ce3+,0.02Li+ samples. The inset picture shows the relationship between $[F(R_{\infty})hv]^2$ and hv of the CGASO host.

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 $F(R_{\infty}) = \frac{(1-R)^2}{2R} = \frac{K}{S}$

$$F(R_{\infty}) = \frac{(1-R)^2}{2R} = \frac{K}{S}$$
 (2)

in which S, K and R represent for coefficients of scattering, absorption and reflection of light energy, respectively. By making $[F(R_{\infty})h\nu]^2$ plot against energy $(h\nu)$ and extrapolating to $[F(R_{\infty})hv]^2 = 0$, an intersection between the straight line and the energy axis is obtained and the value at this point is the band gap value. For CGASO host, E_g was calculated to be 5.37 eV.

The photoluminescence emission (PL) and excitation (PLE) spectra of as-prepared Eu²⁺ doped phosphors have been depicted in Fig. 4. The excitation spectrum of CGASO:0.012Eu²⁺ consists a main band peaking at 350 nm and a shoulder band peaking at 290 nm, resulting from the spin- and orbit-allowed electron transition from 4f⁷ ground state to 4f⁶5d¹ excited state. At the irradiation of 350 nm, the phosphor exhibits a broad emission spectrum with maximum at 508 nm. According to the literatures, energy difference between emission peak wavelength and zero phonon line equals to half of the Stokes shift approximately. Moreover, the energy of zero-phonon line can be estimated from the intersection of the emission and excitation spectra empirically.33-35 For our prepared CGASO:0.012Eu2+ sample, the lowest excited state of ⁷F_I multiplet of the 4f⁶ configuration (zero-phonon line) is determined to be 2.85 eV utilizing the interaction point (435 nm) of excitation and emission curves. In this case, energy difference between emission peak and zero-phone line is calculated to be 0.42 eV. Correspondingly, the Stokes shift value for CGASO:Eu²⁺ sample is estimated to be 0.84 eV. The emission spectra of Ca₂(Al,Ga)₂SiO₇ with different Ga³⁺ substitution content had been shown in Fig. S2,† from which a blue shift can be observed. As discussed previously, the Ga³⁺ substitution can make the Eu-O bond length become short since the Eu²⁺ ions will occupy the Ca²⁺ site in the lattice. Thus, crystal field splitting of 5d energy level of Eu²⁺ should become larger since $D_q \propto 1/R^5$. In this way, a red shift is expected other than a blue shift. However, the emission of Eu²⁺ is also affected by nephelauxetic effect arising from the covalency between cation and ligands. The electronegativity of Ga (1.81) is higher than that of Al (1.61), which will enlarge the electron-cloud overlap of Ga-O. In this case, the covalency of Eu-O connected in Eu-O-Ga will decrease and finally cause the emission shift to a short wavelength. Thus, the blue shift is the result of both crystal field splitting and nephelauxetic effect due to the decrement of bond length and covalency of Eu-O.

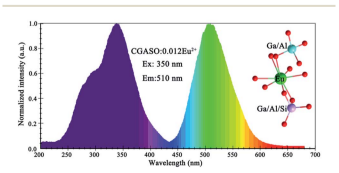


Fig. 4 Excitation and emission spectra of CGASO:0.012Eu²⁺ phosphor.

In order to obtain the optimal doping concentration of Eu²⁺ ions, CGASO: xEu^{2+} phosphors with x varying from 0.003 to 0.021 have been prepared and measured. From Fig. 5(a) we can see that the emission intensity value of CGASO:Eu²⁺ experiences a parabolic change in the whole doping range of Eu²⁺ ions and the maximum emission intensity appears at x = 0.012. The decrement of emission intensity with further increment of Eu2+ content is resulting from concentration quenching effect. The relationship between emission intensity and doping content of phosphors has been depicted in Fig. 5(b) for having an intrusive view. On the basis of Dexter's energy transfer theory, energy migration among the activators appears at high doping content and further results in the concentration quenching. Utilizing the following equation given by G. Blass, critical distance (R_c) for the dopants can be estimated:36

$$R_{\rm c} \approx 2 \left[\frac{3V}{4\pi x_{\rm c} N} \right]^{1/3} \tag{3}$$

In the above formula, x_c stands for critical concentration, Vrepresents the unit cell volume, the N is the number of cation sites that activators can substitute in each unit cell. For CGASO host, N = 4, $V = 306.08 \text{ Å}^3$ and $x_c = 0.012$. Thus, the value of R_c is obtained as 23.01 Å. For oxide phosphors, multipole-multipole interaction and exchange interaction among the activators are usually responsible for non-radiative energy transfer. However, the exchange interaction can be excluded in our discussion since it only fits the energy transfer of forbidden transitions and occurs with critical distance of activators smaller than 5 Å. As a result, multipole-multipole interaction is the main reason for concentration quenching of the Eu²⁺ in our prepared phosphor. According to the report of Van Uiter in 1971, the PL intensity (I) per Eu²⁺ ion in CGASO:Eu²⁺ phosphor satisfy the equation^{37,38}

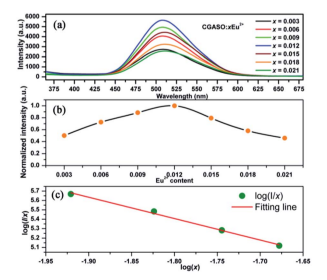


Fig. 5 (a) Emission spectra of CGASO: xEu^{2+} (x = 0.003-0.021) samples. (b) Dependence of relative emission intensity of CGA- $SO:xEu^{2+}$ phosphors on Eu^{2+} content. (c) Relationship between log(I/x)and $\log x$.

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$$\frac{I}{x} = k \left[1 + \beta(x)^{\theta/3} \right]^{-1} \tag{4}$$

in which β and k are constants in a given condition; $\theta=6$, 8 and 10 stands for electric dipole–dipole (d–d), dipole–quadrupole (d–q), and quadrupole–quadrupole (q–q) interactions, respectively; x is the concentration of Eu^{2+} ions. To get a precise value of θ , the relationship between $\log(I/x)$ and $\log(x)$ is plotted after taking logarithm on eqn (4). Correspondingly, the value of θ obtained from the slope of the fitting line as shown in Fig. 5(c) is found to be 6.63, revealing that the concentration quenching of Eu^{2+} ions is mainly resulted from electric dipole–dipole interaction.

The PL and PLE spectra of CGASO:0.02Ce³⁺,0.02Li⁺ phosphor have been shown in Fig. 6. The PLE spectrum with monitor wavelength being 410 nm covers a range from 225 to 375 nm containing three peaks at 254, 293 and 350 nm. This broad excitation band is resulted from spin- and orbit-allowed $4f^1 \rightarrow$ 4f⁰5d¹ electronic transition of the Ce³⁺ ions in crystal field. The emission peak of CGASO:0.02Ce³⁺,0.02Li⁺ locates at 410 nm. Similar with the Eu²⁺ doped samples, the substitution of Ga³⁺ for Al³⁺ ions in Ca₂Al₂SiO₇ host has also caused blue shift of the emission spectra as shown in Fig. S3.† Generally, the ground state of the Ce³⁺ ions can spilt into two energy levels (²F_{5/2} and ²F_{7/2}) owing to spin-orbit interaction. As a result, two symmetric bands with energy difference being $\sim 2000 \text{ cm}^{-1}$ can be obtained by Gaussian decomposition of the emission band. 39,40 For our prepared CGASO:0.02Ce3+,0.02Li+ phosphor, wavelength numbers of the two well-separated Gaussian components are 23 180 cm⁻¹ and 25 007 cm⁻¹ corresponding to 4f⁰5d¹ \rightarrow $^2F_{7/2}$ and $4f^05d^1 \rightarrow ^2F_{5/2}$ transitions, respectively. The energy difference determined to be 1827 cm⁻¹ agrees well with the theoretical value.

The Ce^{3+} doped samples with different doping contents had been synthesized and investigated. As shown in Fig. 7, the optimal doping concentration is y = 0.02 for the Ce^{3+} doped samples. When the value of y is more than 0.02, the energy migration among Ce^{3+} ions has a negative effect on the emission intensity. The relation between Ce^{3+} luminescence intensity and its doping concentration has also been illustrated in Fig. 7 for a clearly view. One can also see that the emission spectra have a small red shift with the increment of Ce^{3+} content, resulting from the variation of the crystal field splitting

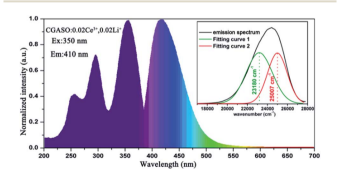


Fig. 6 Photoluminescence spectra of CGASO:0.02Ce³⁺,0.02Li⁺ sample. The inset shows the Gaussian peaks of the emission spectrum.

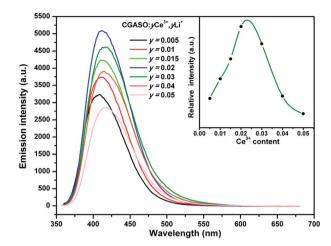


Fig. 7 Emission spectra of CGASO: yCe^{3+} , yLi^+ (y=0.005-0.05) phosphors. The inset shows the relationship between emission intensity and doping content of Ce^{3+} ion.

around Ce³⁺ ions. Since the Li⁺ ions codoped with Ce³⁺ ions have smaller radius than Ca²⁺ ions, the doping of the Li⁺ ions can shrinks the CGASO crystal lattice. The variation on crystal volume makes the crystal field strength around the Ce³⁺ ions enhance and the 5d energy levels of Ce³⁺ split largely. As a result, the Ce³⁺ emission shift to longer wavelength. The luminescence spectra indicate that the Ce³⁺ doped CGASO sample is a potential UV excited blue emission phosphor.

3.3 Luminescence dynamic, thermal stability and CIE coordinates of CGASO:Eu²⁺ and CGASO:Ce³⁺,Li⁺ phosphors

To have an understanding on the luminescence dynamic of the phosphors, the decay curves of CGASO: xEu^{2+} ($\lambda_{ex}=350$ nm, $\lambda_{em}=508$ nm) and CGASO: yCe^{3+} , yLi^+ ($\lambda_{ex}=350$ nm, $\lambda_{em}=410$ nm) samples have been measured and illustrated in Fig. 8. The average lifetimes of the samples can be determined by the following equation:

$$\tau_{\text{avg}} = \int_0^\infty t I(t) dt / \int_0^\infty I(t) dt$$
 (5)

where I(t) represents the emission intensity of activators at time t. The lifetimes for CGASO:xEu $^{2+}$ phosphors with x being 0.003, 0.006, 0.009, 0.012, 0.015, and 0.021 were determined to be 0.793, 0.733, 0.685, 0.659, 0.594, and 0.522 μ s, respectively. The lifetimes of Ce $^{3+}$ ions were calculated to be 39.4, 38.4, 36.0 33.5, 31.9, and 28.6 ns for CGASO:yCe $^{3+}$,yLi $^{+}$ samples with y=0.005, 0.01, 0.015, 0.02, 0.03, and 0.04, respectively. In this case, both the increment of Eu $^{2+}$ and Ce $^{3+}$ content have decreased the decay lifetime of phosphors, which can be ascribed to the energy transfer among activators.

Since the working temperature of WLEDs can reach 150 °C, thermal stability comes to be an important characteristic for phosphors and is always addressed. Fig. 9 shows the normalized emission intensity of CGASO:0.012Eu²⁺ and CGASO:0.02-Ce³⁺,0.02Li⁺ phosphors with temperature ranging from 25 to 200 °C (λ = 350 nm). With the increment of temperature, the luminescence intensity of Eu²⁺ and Ce³⁺ both decreased

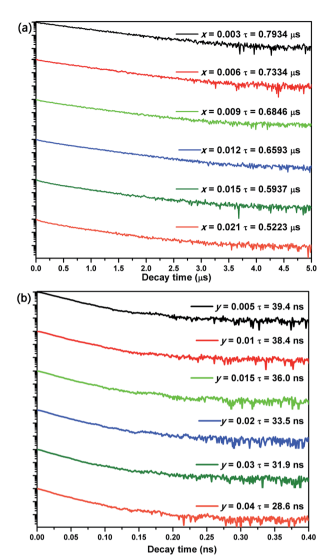


Fig. 8 Decay curves of (a) CGASO:xEu²⁺ and (b) CGASO:yCe³⁺,yLi⁺ phosphors with different doping concentrations.

gradually owing to thermal quenching. At 150 °C, the luminescence intensity of CGASO:0.012Eu2+ and CGASO:0.02-Ce³⁺,0.02Li decreased to 50.2% and 71.8% of their initial intensity at 25 °C. Generally, there are two models to explain the reason of thermal quenching. One is the autoionization model proposed by Dorenbos,41 the other is configurational diagram as depicted in inset of Fig. 9. The autoionization model has some limitations and is only suitable for the phosphors that the 5d energy level of rare earth ions is close to the conduction band minimum well before the 4f-5d crossing point. The configurational diagram depicts relationship between energy and ionic degrees of freedom for both ground and excited state of a material and is extremely successful in demonstrating various luminescence processes. Here, the latter one is employed to demonstrating the thermal quenching of prepared phosphors. Under UV irradiation, the electrons of activators can be excited from ground state (point a) to excited state (point b). The excited electrons will modify the forces inside the crystal and alter the

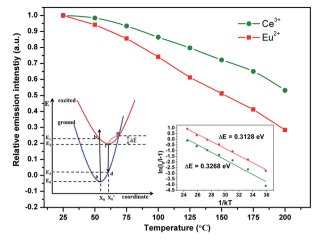


Fig. 9 Relative emission intensities of CGASO: $0.012Eu^{2+}$ and CGASO: $0.02Ce^{3+}$, $0.02Li^{+}$ samples at different temperatures. The inset depict the configuration coordinate diagram and plots of $\ln[(I_0/I_7)-1]$ versus 1/kT for CGASO: $0.012Eu^{2+}$ and CGASO: $0.02Ce^{3+}$, $0.02Li^{+}$ samples.

average atomic positions from X_0 to X_0^* , leading to the crossing of the 5d and 4f energy curves. At low temperature, the excited electrons will go back to ground state with light emission after dissipation some energy through phonon relaxation and the crystal will also returns to its original ground state geometry. However, at high temperature the excited electrons can absorb the phonon energy and be thermally activated to the crossing point. Finally, the electrons at crossing point decay non-radiatively to the ground state, making the thermal quenching appear. This phenomenon will be more severe with temperature increasing. Energy difference between point e and c is called activation energy (ΔE) which affects the thermal stability of phosphors directly. The value of ΔE can be determined by the following modify Arrhenius equation:

$$I_T = I_0 \left[1 + A \exp\left(-\frac{\Delta E}{kT}\right) \right]^{-1}$$
 (6)

where I_0 the initial emission intensity, I_T stands for the emission intensity at different temperatures, A is a constant for a given host, k is the Boltzmann constant (8.629 × 10⁻⁵ eV K⁻¹). The inset of Fig. 9 depicted the dependence of $\ln[(I_0/I_T) - 1]$ on 1/kT. According to eqn (6), the values of ΔE were calculated to be 0.3128 and 0.3268 eV for CGASO:0.012Eu²⁺ and CGASO:0.02-Ce³⁺,0.02Li⁺ samples, respectively.

Quantum yield (Φ) as an important parameter for the prepared phosphor has been measured by integrated sphere method. Under 350 nm irradiation, quantum yields for CGASO:0.012Eu²⁺ and CGASO:0.02Ce³⁺,0.02Li⁺ samples are 65.0% and 65.2%, respectively. Since the value of quantum yield can be easily affected by the morphology, crystallinity and size of the phosphor, the Φ can be further improved by optimize the preparation condition and phosphor composition. Fig. 10(a) illustrates the Commission International del'Eclairage (CIE) chromaticity coordinates of typical CGASO:0.012Eu²⁺ and CGASO:0.02Ce³⁺,0.02Li⁺ determined by their corresponding PL

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Fig. 10 (a) CIE coordinates and digital photographs of CGA-SO: $0.012Eu^{2+}$ and CGASO: $0.02Ce^{3+}$, $0.02Li^{+}$ samples. (b) EL spectrum of a white LED device fabricated by "CGASO: $0.012Eu^{2+}$ + CGA-SO: $0.02Ce^{3+}$, $0.02Li^{+}$ + red phosphor + 365 nm LED chip". The inset shows the corresponding LED device photograph.

spectra. For the Eu²⁺ doped phosphor, the CIE coordinates located in the green region with the value being (0.193, 0.504), which is consistent with its emission colour as can be seen from the digital photograph. The CIE coordinates of CGASO:0.02- Ce^{3+} , 0.02Li was determined to be (0.156, 0.037). The luminescence properties indicate that our prepared CGASO:0.012Eu2+ and CGASO:0.02Ce³⁺,0.02Li⁺ phosphors are potential UV light excited green and blue phosphors, respectively. It is known to us that white light can be obtained when mix blue, green and red light at appropriate intensity. To demonstrate the application of prepared phosphors, a white LED device has been fabricated by coating phosphors (CGASO:0.012Eu²⁺, CGASO:0.02Ce³⁺,0.02Li⁺, and CaAlSiN₃:Eu²⁺) on a 365 nm UV chip. The corresponding EL spectrum and digital photograph has been depicted in Fig. 10(b). The Ra and CCT values for the fabricated white LED device with CIE coordinates being (0.359, 0.320) are determined to be 82.1 and 4232 K, respectively.

4. Conclusions

To summarize, we have obtained the UV efficiently excited Eu²⁺ and Ce³⁺ doped Ca₂GaAlSiO₇ phosphors firstly by cation substitution. Structure refinement by GSAS indicates that Ca₂-GaAlSiO₇ own tetrahedral crystal system and $P\bar{4}2_1m$ (113) space group. The Ca₂GaAlSiO₇:0.012Eu²⁺ sample emits intense green light with CIE coordinates being (0.193, 0.504) and quantum efficiency being 65.0%, while the emission of Ca₂-GaAlSiO₇:0.02Ce³⁺,0.02Li⁺ locates in blue region with CIE coordinates being (0.156, 0.037) and quantum efficiency value being 65.2%. The substitution of Ga³⁺ for Al³⁺ has caused a blue shift of emission due to the decrement of cation-ligand distance and covalency. At 150 °C, the emission intensity of CGASO:0.012Eu²⁺ and CGASO:0.02Ce³⁺,0.02Li⁺ decrease to 50.2% and 71.8% of their initial values, respectively. Moreover, a white LED device with $R_a = 82.1$ and CCT = 4232 K had been fabricated by 365 nm UV chip with our prepared phosphors and commercial red phosphor. The investigation suggests that the Eu²⁺ and Ce³⁺ activated samples may be potential green and blue phosphors for UV WLEDs.

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