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Introduction

Wide visible-range fluorescence of Eu³⁺ located in the macroscopic bi-layer ceramic/glass composite

Haifeng Shi,^a Jiaxin Yang,^a Zhimin Yu,^a Yu Song,^a Edwin Yue Bun Pun,^b Xin Zhao^{*a} and Hai Lin^b *^{ab}

The Eu³⁺ doped fluoride bi-layer ceramic/glass composite (GC_{ZBL}-Eu) was prepared by a one-step method and the effective wide visible-range fluorescence was recorded. The de-population rates of the ⁵D₀, ⁵D₁, ⁵D₂, and ⁵D₃ multi-levels in the glass layer (G_{ZBL}-Eu) were estimated to be 214, 746, 1163, and 680 s⁻¹, respectively, and that in the ceramic layer (C_{ZBL}-Eu) were 211, 730, 1075, and 654 s⁻¹, which implies multi-channel radiative transitions due to the non-radiative relaxation limitation of low OH content and low phonon energy. Simultaneously, the quantum efficiencies of the ⁵D₀ levels in G_{ZBL}-Eu and C_{ZBL}-Eu were as high as 98.5% and 94.8%, respectively, thus demonstrating the effectiveness of the radiative transition emissions from Eu³⁺. Besides, GC_{ZBL}-Eu with the glass forming layer increases the emission intensity by 24% compared to C_{ZBL}-Eu, which is attributed to the multiple-cycle reflection in the composite structure of the glass–ceramic transition region, and the color coordinates of C_{ZBL}-Eu (0.483, 0.385) and GC_{ZBL}-Eu (0.469, 0.389) show that they can release yellowish-white light. The heterostructured GC_{ZBL}-Eu provides a new approach for laser lighting, fluorescent display, and up-conversion applications.

White light-emitting diodes (WLEDs) have recently attracted considerable attention as next generation light sources and as alternatives for conventional incandescent and fluorescent lamps.¹⁻⁴ One of the most popular ways to obtain WLEDs is to combine near UV-LEDs with a glass or crystal phosphor emitting in the blue, green, and red spectral regions.⁵⁻⁷ For this purpose, multiple luminescence centers in the phosphors are widely adopted to achieve high brightness and high color rendering index of white light emission.⁸⁻¹² However, rare earth (RE) co-doping leads to complex cross relaxation processes, which cannot release photons efficiently. In view of the high fluorescence quantum efficiency in visible white light illumination, the exploration focusing on the compound material with single RE³⁺ ions doping and multi-channel emission becomes urgent.

The emission spectrum of Eu³⁺ located in the hosts, characterized by the existence of high energy phonons such as the oxides, is strong only for the case of the bands produced by the radiative relaxations from ${}^{5}D_{0}$.¹³⁻¹⁹ The emissions from ${}^{5}D_{0}$ are favored mostly because the energy difference between the ${}^{5}D_{3}$, ${}^{5}D_{2}$, ${}^{5}D_{1}$, and ${}^{5}D_{0}$ levels of the Eu³⁺ cation can be covered by

non-radiative transitions from the higher levels to ${}^{5}D_{0}$. The resulting emission profile, with intense bands mostly confined to a narrow range, is not suitable for the preparation of whitelight phosphors if the intention is to use only a single RE³⁺ dopant. However, in hosts having weak phonons, such as fluorides, the non-radiative transitions from the higher electronic levels to ⁵D₀ become difficult and the electronic transitions originating in those levels acquire higher intensity than in the oxide hosts. The result is a fluorescence spectrum exhibiting bands of significant intensity (albeit emission at wavelength higher than 500 nm remains strongest) at various positions of the visible range; such a spectral profile makes Eu^{3+} , in fluorides, a possible candidate for the fabrication of white light phosphors based on a single RE³⁺. In this work, for the first time, it has been examined whether the emission of Eu³⁺, introduced in a Zr–Ba–La–F host (macroscopic ceramic/ glass bi-layer), can be used to obtain such phosphors. In addition, the transitions ${}^5D_0 \rightarrow {}^7F_1$ and ${}^5D_0 \rightarrow {}^7F_2$ have different dependences on the crystal field environment belonging to the magnetic dipole and electric dipole transitions.²⁰⁻²⁵ Also, Eu³⁺ ions have similar ionic radius and crystalline properties as other RE³⁺ ions, which can be used as a probe of the local structure to explore the influence of the micro-environment of RE3+ ions on the luminescence of materials.

In this work, Eu³⁺-doped fluoride ceramics with a glass layer (GC_{ZBL}-Eu) have been prepared and intense yellowish-white fluorescence, attributed to multi-peak emission, was observed.

[&]quot;School of Textile and Material Engineering, Dalian Polytechnic University, Dalian 116034, P. R. China. E-mail: zhaoxin@dlpu.edu.cn; lhai8686@yahoo.com

^bDepartment of Electronical Engineering and State Key Laboratory of Terahertz and Millimeter Waves, City University of Hong Kong, Tat Chee Avenue, Kowloon, Hong Kong, P. R. China

With the formation of the thin glass layer on the ceramic surface, the GC_{ZBL}-Eu increases its emission intensity by 24% compared to C_{ZBL}-Eu, which is ascribed to multiple-cycle reflection in the complex structure of the glass-ceramic transition region. The micro-environment symmetry of G_{ZBL}-Eu and C_{ZBL}-Eu is judged according to the value of the J-O parameter Ω_2 with different emission intensity ratios between ${}^5D_0 \rightarrow {}^7F_2$ and ${}^5D_0 \rightarrow {}^7F_1$. The differential fluorescence characteristics, such as the radiative lifetimes of the multi-levels and the quantum efficiency of the 5D_0 levels from Eu³⁺ in G_{ZBL}-Eu and C_{ZBL}-Eu and GC_{ZBL}-Eu are located in the yellowish-white lighting region, which indicates that the hetero-structured materials can be well applied to optical devices.

Preparation and measurements of GC_{ZBL} -Eu

The fluoride bi-layer ceramic/glass composites were prepared according to the molar host composition of 60ZrF₄-30BaF₂-10LaF₃ (ZBL) via the melt-quench method and 0.2 wt% EuF₃ was introduced into the ZBL matrix as the dopant. The well-mixed high-purity fluoride raw materials were melted at 900 °C for 5 min in a platinum crucible using an electric furnace at the rate of 5 °C min⁻¹ from room temperature to 900 °C. In addition, NH₄F on the raw materials provides a reducing atmosphere and thus, the molten glass was quenched onto an aluminum plate. Herein, the bottom molten glass, when in contact with the aluminum plate, takes away a lot of heat rapidly and forms an ultrathin glass layer owing to the process of efficient heat conduction. Correspondingly, the upper liquid starts to crystallize at the interface with air because air cannot take away a lot of heat. Subsequently, the molded samples were annealed at 260 °C for 2 h and then cooled down slowly to room temperature. The schematic diagram of the systematic preparation procedure is exhibited in Fig. 1. Relevant tests were performed on the glass surface of the bi-layer ceramic/glass composite, the ceramic layer, and the glass layer, which were named as GC_{ZBL}-

Eu, C_{ZBL} -Eu, and G_{ZBL} -Eu, respectively (C_{ZBL} -Eu and G_{ZBL} -Eu samples were ground to the glass or ceramic layer by the GC_{ZBL} -Eu composite).

The differential thermal analysis (DTA) scan was carried out on a WCR-2D differential thermal analyzer at the rate of 10 °C min⁻¹ from room temperature to 900 °C. X-ray diffraction (XRD) measurements for the powders of G_{ZBL} -Eu and C_{ZBL} -Eu were carried out on a Shimadzu XRD-7000 diffractometer (40 kV, 30 mA). The morphological behavior of the section of GC_{ZBL} -Eu was observed by a field-emission scanning electron microscope (SEM instrument, JEOL JSM-7800F). The glass layer thickness of



Fig. 2 (a) The DTA curve of G_{ZBL} -Eu. Inset: the cross-section morphology of the GC_{ZBL} -Eu sample under an optical microscope. (b) The XRD patterns of G_{ZBL} -Eu and C_{ZBL} -Eu.



Fig. 1 The schematic diagram of the systematic preparation procedure.

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Fig. 3 The schematic illustration of the crystal structure of $BaZrF_6$.

 GC_{ZBL} -Eu was measured by a fluorescence microscope (Imaging system CK-500). The refractive index of G_{ZBL} -Eu was measured by using the Metricon 2010 prism coupler. The visible fluorescence spectra and the fluorescence decay curves were obtained using a Hitachi F-7000 fluorescence spectrophotometer equipped with an R928 photomultiplier tube (PMT) as the detector and a commercial Xe-lamp as the excitation source.

Results and discussion

The transition temperature $T_g = 305$ °C, the onset crystallization temperature $T_x = 364$ °C, and the peak temperature $T_c =$ 379 °C of the DTA curve are shown in Fig. 1(a). The parameters including the temperature difference value $\Delta T = T_x - T_g = 59$ °C, the thermal stability parameter $H = \Delta T/T_g = 0.19$, and the Saad–Poulain criterion $S = \Delta T(T_c - T_x)/T_g = 2.9$ °C demonstrate that this sample is more effortless to crystallize than other oxide glasses or oxyfluoride glasses, and is suitable for the preparation of the fluoride ceramic-based composite glass. The inset of Fig. 2(a) shows the cross-sectional morphology of GC_{ZBL}-Eu under an optical microscope. The shiny part of this picture is the glass layer, whose thickness was identified to be ~0.81 mm, the dark part is ceramic-based, and the intersection part of the two layers is called the GC transition region.

The typical XRD pattern of G_{ZBL} -Eu in Fig. 2(b) exhibits a broad diffuse scattering at lower angles without the narrow diffraction peaks of the crystal phase and the amorphous state of the prepared glass layer is well-identified. At the same time, by comparing the sharp diffraction peaks in the XRD spectrum of C_{ZBL} -Eu, the crystal phase of C_{ZBL} -Eu is identified to be pure BaZrF₆. The slight differences in the cell parameters between C_{ZBL} -Eu (a = 7.744 Å, b = 11.691 Å, c = 5.404 Å, $\alpha = \beta = \gamma = 90^{\circ}$) and the standard BaZrF₆ phase (a = 7.681 Å, b = 11.357 Å, c =5.511 Å, $\alpha = \beta = \gamma = 90^{\circ}$) indicate the possibility of lattice deformation caused by the introduction of Eu³⁺. The cell volumes of orthorhombic C_{ZBL} -Eu and BaZrF₆ judged by the cell parameters were calculated to be 489.25 and 480.74 Å³ by the equation $V = a \times b \times c$, indicating that the doping of Eu³⁺ has little effect on the cell volume before and after substitution. In



Fig. 4 (a and b) SEM images of the cross-section of GC_{ZBL} -Eu. (c and d) EDS spectra and the elemental content of G_{ZBL} -Eu and C_{ZBL} -Eu. (e–i) Elemental mapping of C_{ZBL} -Eu.



Fig. 5 (a) The FT-IR spectrum of the glass layer. (b and c) The excitation spectra of G_{ZBL}-Eu and C_{ZBL}-Eu monitored at 615 nm emission.

order to maintain electrical neutrality, Eu^{3+} , whose radius of 0.95 Å is between that of Zr^{4+} (0.80 Å) and Ba^{2+} (1.35 Å), is more likely to replace Zr^{4+} in the lattice and forms F^- vacancies.

Molten glass appears to be disordered owing to increased interatomic amplitudes and irregular diffusion of the molecules, and the disordered state is fixed to form a glass on sudden cooling. When molten glass cools down slowly, each atom returns to the lowest energy state and forms crystals. In this matrix, $BaZrF_6$ is a crystal with lower lattice energy and is easier to precipitate; Fig. 3 shows the schematic diagram of the crystal structure of $BaZrF_6$.

Spectrum 1 and spectrum 2 with the same element type were chosen from the SEM images of the sample cross-section in Fig. 4(a) for elemental analysis, as shown in Fig. 4(c) and (d). In comparison, the element contents in G_{ZBL} -Eu and C_{ZBL} -Eu have little difference within the allowable error range, indicating that there is no obvious element migration during the crystallization process. In addition, the absence of oxygen and other elements in the EDS spectra indicates that there is no foreign matter in the prepared G_{ZBL} -Eu and C_{ZBL} -Eu and the surface of the samples is not oxidized. The SEM image of the ceramic region displays that the BaZrF₆ crystalline grain covered by glass in

 C_{ZBL} -Eu grows in an ordered direction and arrangement, as shown in Fig. 4(b). The elemental mapping in the regions contained within the red frames of Fig. 4(e–i) exhibits that Zr and Ba are densely distributed whereas it is the opposite for La, which indicates that the red frame region represents the crosssection of the BaZrF₆ grains coated with glass.

Based on the FT-IR spectrum of the glass layer in Fig. 5(a), the molar absorption coefficient α_{OH} can be used to evaluate the residual OH content in the glass samples and was found to be 0.91 cm⁻¹ in the 75TeO₂-10ZnO-10Na₂O-5GeO₂ glasses,²⁶ while the value in this work is as low as 0.57 cm⁻¹. The low OH content of the samples is helpful in achieving the anticipated photon emission by reducing the fluorescence loss. In view of the high sensitivity of the multi-channel transition to the phonon energy, the maximum phonon energy (*E*) of the glass samples was estimated through the empirical formula *E* = 92.9 + 0.4257*R*, where *R* corresponds to the wavenumber at 10% transmittance of the infrared transmission sideband and the maximum phonon energy of G_{ZBL}-Eu is identified to be ~483 cm⁻¹.

The energy difference between the phonon sideband spectrum and the zero-phonon line is the coupled phonon energy



Fig. 6 (a) Emission spectra of G_{ZBL} -Eu and C_{ZBL} -Eu under 395 nm excitation. (b) The schematic diagram of multimodal emission energy levels of Eu³⁺ under 395 nm excitation.



Fig. 7 The relative spectral power distributions (a) and the relative emission photon distributions (b) of G_{ZBL} -Eu and C_{ZBL} -Eu under 395 nm excitation.



Fig. 8 Fluorescence decay curves from the ${}^{5}D_{0}$, ${}^{5}D_{1}$, ${}^{5}D_{2}$, and ${}^{5}D_{3}$ level of Eu $^{3+}$ in G_{ZBL}-Eu (a–d) and C_{ZBL}-Eu (e–h) monitored at 615, 535, 509, and 464 nm, respectively.

for the electron transition of the RE³⁺ ions. The phonon energies of G_{ZBL}-Eu and C_{ZBL}-Eu were estimated to ~452 and ~424 cm⁻¹ according to the fitting of the two curves shown in Fig. 5(b) and (c), respectively. The local vibrational modes with the corresponding RE³⁺ ions are generally considered to be slightly less than the maximum phonon energy of the material.²⁷ G_{ZBL}-Eu and C_{ZBL}-Eu with low phonon energies, which have been confirmed by the two methods mutually, can reduce the possibility of non-radiative transitions and facilitate fluorescence emission.

The emission spectra of G_{ZBL}-Eu and C_{ZBL}-Eu were recorded under 395 nm excitation as shown in Fig. 6(a). The emission intensity of C_{ZBL}-Eu is much higher than that of G_{ZBL}-Eu owing to the ordered arrangement of BaZrF₆ particles and another factor is that Eu³⁺ maybe be retained in C_{ZBL}-Eu more than in GZBL-Eu. Eleven emission peaks of GZBL-Eu and CZBL-Eu are located at 429, 445, 464, 488, 509, 525, 535, 554, 590, 615, and 698 nm, which originate from the ${}^{5}D_{3} \rightarrow {}^{7}F_{2,3,4}$, ${}^{5}D_{2} \rightarrow {}^{7}F_{2,3}$, ${}^{5}D_{1}$ \rightarrow $^{7}F_{0,1,2}$, and $^{5}D_{0} \rightarrow$ $^{7}F_{1,2,4}$ transitions of Eu³⁺, respectively. The energy level diagram in Fig. 6(b) shows the radiation and relaxation processes of multiple channels and the values of energy gaps among the ⁵D₀, ⁵D₁, ⁵D₂, and ⁵D₃ levels. In the G_{ZBL}-Eu and CZBL-Eu systems, the multiphonon relaxation processes from the ⁵D₃ and ⁵D₂ levels are accomplished by six-phonon bridging and that from ${}^{5}D_{1}$ level is completed by four-phonon bridging. All of them belong to high order processes (more than 3 phonons) and thus, the probability of energy loss by nonradiative transitions is lowered. Therefore, the blue and green light emission peaks from ⁵D₃, ⁵D₂, and ⁵D₁ levels cooperating with the red-light emission peaks from the ⁵D₀ level can be well applied to luminescent color tuning, thus expanding the application direction of single RE³⁺ ions.

By referring to the halogen lamp, the relative spectral power distribution is converted from the emission spectrum and is shown in Fig. 7(a). Depending on the relative spectral power distribution, photon distribution $N(\nu)$ can be deduced as $N(\nu) = \frac{\lambda^3}{hc} P(\lambda)$, where ν , λ , h, c, and $P(\lambda)$ represent the wavenumber, wavelength, Planck constant, vacuum light velocity, and spectral power distribution, respectively.²⁸⁻³⁰ Under 395 nm excitation, the emission photon number distribution curves of G_{ZBL}-Eu and C_{ZBL}-Eu are derived as shown in Fig. 7(b).

 Ω_2 , which is influenced by the local structure symmetry of Eu³⁺ in G_{ZBL}-Eu and C_{ZBL}-Eu, is usually used to describe the emission distribution of the ⁵D₀ level and can be calculated based on the emission photon number distributions instead of the general absorption spectra. In the calculation, the ordinary refractive index *n* value of G_{ZBL}-Eu is measured to be 1.555 and the value of C_{ZBL}-Eu is estimated to be 1.590.³¹ The Ω_2 value of Eu³⁺ in C_{ZBL}-Eu is 1.75 × 10⁻²⁰ cm², which is lower than 2.49 × 10⁻²⁰ cm² in G_{ZBL}-Eu. A smaller Ω_2 value indicates a higher symmetry of the local structure of Eu³⁺ ions and weaker covalency around the metal–donor interaction. Also, compared with C_{ZBL}-Eu, G_{ZBL}-Eu, and other substrates,³²⁻³⁷ a smaller Ω_2 value (higher symmetry) corresponds to lower red-light emission ratio of the ⁵D₀ \rightarrow ⁷F₂ transition. In other words, at a higher symmetry site, the intensity ratio of the ⁵D₀ \rightarrow ⁷F₂ emission is reduced and thus,

	G _{ZBL} -Eu		C _{ZBL} -Eu		
Energy level	Experimental lifetime (ms)	De-population rate (s^{-1})	Experimental lifetime (ms)	De-population rate (s^{-1})	
${}^{5}D_{0}$	4.67	214	4.74	211	
⁵ D ₁	1.34	746	1.37	730	
⁵ D ₂	0.86	1163	0.93	1075	
⁵ D ₃	1.47	680	1.53	654	

Table 1 The experimental lifetimes and de-population rates of ${}^{5}D_{0}$, ${}^{5}D_{1}$, ${}^{5}D_{2}$, and ${}^{5}D_{3}$ in G_{ZBL}-Eu and C_{ZBL}-Eu

it becomes more similar to that of the ${}^{5}D_{0} \rightarrow {}^{7}F_{1}$ transition. This means that C_{ZBL} -Eu can better limit the emission of red light and using the shorter-wavelength orange light generated in the ${}^{5}D_{0} \rightarrow {}^{7}F_{1}$ transition, it is easier to synthesize white light with blue and green light from the ${}^{5}D_{3}$, ${}^{5}D_{2}$, and ${}^{5}D_{1}$ level transitions. Such a change towards more equal intensity emission at various wavelengths is more suitable for realizing an overall white emission of the phosphor and the excitation source system.

To better express the capacity of the potential radiative transition among the levels, the experimental average lifetimes and the de-population rate of Eu³⁺ in G_{ZBL}-Eu and C_{ZBL}-Eu can be derived from the fluorescence decay curves, as shown in Fig. 8, and the related results are calculated and listed in Table 1. In the fluoride system with low phonon energy, photons can be effectively released and the low depopulation rate represents that the radiative transition occupies a major proportion. In order to evaluate the effective radiation of C_{ZBL}-Eu and G_{ZBL}-Eu, the quantum efficiencies of the ${}^{5}D_{0}$ level was calculated. The data of theoretical radiative fluorescent lifetimes (including the relevant calculation data), experimental average lifetimes, and quantum efficiencies are listed in Table 2. The quantum efficiencies of Eu³⁺ from the ⁵D₀ level in G_{ZBL}-Eu and C_{ZBL}-Eu were calculated to be 98.5% and 94.8%, which demonstrate the effectiveness of radiative transition emission from Eu^{3+} in G_{ZBL} -Eu and C_{ZBL} -Eu. The difference in the quantum efficiencies of G_{ZBL}-Eu and C_{ZBL}-Eu is consistent with the conclusion that the luminous efficiency of Eu³⁺ decreases in a more symmetrical crystal field environment. The high quantum efficiencies of G_{ZBL}-Eu and C_{ZBL}-Eu demonstrate the effectiveness of radiative transition emissions from Eu³⁺ and show the application prospects in energy conversion and energy saving.



Fig. 9 Comparison of emission intensities (a) and the relative spectral power distributions (b) between C_{ZBL} -Eu and GC_{ZBL} -Eu under 395 nm excitation.

The emission spectra and the relative spectral power distributions of C_{ZBL} -Eu and GC_{ZBL} -Eu are shown in Fig. 9(a) and (b) under 395 nm excitation, and both them show multi-channel

Table 2 Photon number ratios, spontaneous transition probabilities A_{ij} , fluorescence branching ratios β_{ij} , calculated radiative fluorescent lifetimes τ_{rad} , experimental average lifetimes τ_{exp} , and quantum efficiencies of ⁵D₀ in G_{ZBL}-Eu and C_{ZBL}-Eu

Sample	Transition	Energy (cm ⁻¹)	Photon number ratio	A_{ij} (s ⁻¹)	eta_{ij} (%)	$ au_{ m rad}$ (ms)	$ au_{ m exp}$ (ms)	Quantum efficiency (%)
G _{ZBL} -Eu	${}^{5}\mathrm{D}_{0} \rightarrow {}^{7}\mathrm{F}_{1}(N_{1})$	16 949	_	64.3	30.5	4.74	4.67	98.5
	${}^{5}\mathrm{D}_{0} \rightarrow {}^{7}\mathrm{F}_{2}(N_{2})$	16 271	$N_2/N_1 = 1.30$	83.8	39.7			
	${}^{5}\mathrm{D}_{0} \rightarrow {}^{7}\mathrm{F}_{4}(N_{4})$	14 318	$N_4/N_1 = 0.98$	63.0	29.8			
C _{ZBL} -Eu	${}^{5}\mathrm{D}_{0} \rightarrow {}^{7}\mathrm{F}_{1}(N_{1})$	16 938	_	68.7	34.4	5.00	4.74	94.8
	${}^{5}\mathrm{D}_{0} \rightarrow {}^{7}\mathrm{F}_{2}(N_{2})$	16 207	$N_2/N_1 = 0.98$	63.3	31.7			
	${}^{5}\mathrm{D}_{0} \rightarrow {}^{7}\mathrm{F}_{4}(N_{4})$	14 351	$N_4/N_1 = 1.06$	67.9	33.9			



Fig. 10 (a) Color coordinates in the CIE 1931 chromaticity diagram of C_{ZBL} -Eu and GC_{ZBL} -Eu under 395 nm excitation. (b) The schematic diagram of enhanced GC_{ZBL} -Eu emission process under 395 nm excitation. Inset: fluorescent photographs of C_{ZBL} -Eu and GC_{ZBL} -Eu under 395 nm excitation (left), and the schematic of the photos obtained (right).

radiative transitions of Eu^{3+} in C_{ZBL} -Eu and GC_{ZBL} -Eu. Herein, the CIE-1931 chromaticity coordinates for the luminescence of C_{ZBL} -Eu and GC_{ZBL} -Eu were derived to be (0.483, 0.385) and (0.469, 0.389), which fall into the yellowish-white region, as shown in Fig. 10(a). The chromaticity coordinates prove that the emission light of C_{ZBL} -Eu and GC_{ZBL} -Eu is yellowish-white, suggesting that the Eu³⁺-activated GC_{ZBL} -Eu system has application prospect as a single phosphor.

The emission intensity of GC_{ZBL} -Eu at 590 nm is 1.24 times higher than that for C_{ZBL} -Eu, where the significant improvement in the emission intensity can be attributed to the complex surface morphology of the ceramic matrix and the intense dispersion effect of the glass layer, and the schematic diagram of increasing the incident light absorption efficiency and enhancing the emission is shown in Fig. 10(b). When the incident light shines into GC_{ZBL} -Eu, the under-utilized light is rereflected to the ceramic boundary, forming a multiple-cycle effect owing to the specular reflection of the glass layer. Then, the incident light will be absorbed by the glass phase in the reflection process of the glass–ceramic transition region and is converted into other wavelengths for emission.

As shown in the inset of Fig. 10(b), intense lighting is observed in C_{ZBL} -Eu and GC_{ZBL} -Eu, and the violet lighting in the picture is illuminated by a 395 nm Xe lamp. Simultaneously, the purple source overlaps with the emitted yellowish-white light to form the pink-white light. Effective fluorescence emission enhancement facilitates better application of GC_{ZBL} -Eu with warm-white light emission of laser lighting.

Conclusion

Hetero-structured GC_{ZBL}-Eu with glass region, ceramic region, and GC transition region obtained by the one-step method exhibits yellowish-white light and the crystal phase of C_{ZBL}-Eu is identified as BaZrF₆. Multi-channel radiative transitions were discovered through multi-peak emission, long fluorescence lifetimes, and de-population rates of the ⁵D₀, ⁵D₁, ⁵D₂, and ⁵D₃

levels in G_{ZBL} -Eu and C_{ZBL} -Eu, which is because the low OH content and low phonon energy limit the non-radiative relaxation. The effective radiative transition emission of the ${}^{5}D_{0}$ levels in G_{ZBL} -Eu and C_{ZBL} -Eu is demonstrated by the high quantum efficiency of 98.5% and 94.8%, respectively. The release of yellowish-white light is indicated by the color coordinates of C_{ZBL} -Eu (0.483, 0.385) and GC_{ZBL} -Eu (0.469, 0.389), and the emission intensity of GC_{ZBL} -Eu is 1.24 times higher than that of C_{ZBL} -Eu due to multiple-cycle reflection in the complex structure of the GC transition region. The effective yellowish-white lighting proves that the GC_{ZBL} -Eu system with fluorescence enhancement has potential in the application direction of laser illumination.

Conflicts of interest

There are no conflicts to declare.

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