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Stabilization of divalent Eu\textsuperscript{2+} in fluorosilicate glass-ceramics via lattice site substitution\textsuperscript{†}

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Fluorosilicate glasses and glass-ceramics with MF\textsubscript{2} (M = Ca, Sr, Ba), ZnF\textsubscript{2} or LaF\textsubscript{3} components were investigated to host divalent Eu\textsuperscript{2+} for photoluminescence (PL) application. X-ray diffraction phase identification and a series of spectroscopic analyses were performed to reveal the relationship between microstructure and the reduction of Eu\textsuperscript{3+} → Eu\textsuperscript{2+}. The precursor glasses were believed being constituted by silicate-rich phases and fluoride-rich phases, due to the immiscibility of fluoride-and-silicate mixed glass system. After heat treatment, the fluoride-rich glass phases could transform into fluoride crystalline phase in the glass-ceramics. Europium tended to enrich in the fluoride-rich phases in the glasses or in the precipitated fluoride crystalline phases in the glass-ceramics. Small amounts of Eu\textsuperscript{3+} were reduced to Eu\textsuperscript{2+} in the glasses where the electronegativity had a crucial impact. In contrast, large amounts of Eu\textsuperscript{2+} were reduced to Eu\textsuperscript{3+} in the glass-ceramics containing MF\textsubscript{2} nanocrystals, where the reduction was determined by lattice site substitution. Using ZnAl\textsubscript{2}O\textsubscript{4} containing glass-ceramics as reference, it was evidenced that the similar and a little larger radii between sites and substitution ions are the prerequisite for Eu\textsuperscript{3+}/M\textsuperscript{2+} substitution. And using LaF\textsubscript{3} containing glass-ceramics as reference, it was certified that unbalanced charge at substitution sites induce the Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction.

Introduction

As Eu\textsuperscript{2+} possesses ideal properties for light emission such as large absorption cross section, broad band spectrum, and highly efficient photoluminescence (PL), it has been wildly adopted as active center in phosphor materials.\textsuperscript{1,2} Owing to the exposure of 5d-shell of Eu\textsuperscript{2+} to ligands, it is also facile to tune its emission from violet to red by adjusting host lattice.\textsuperscript{3} Thus, Eu\textsuperscript{2+}-doped inorganic phosphors have been widely used in commercial lighting and display devices, such as epoxy resin-or-silica mixed phosphor converted light emitting diodes (PC-LED) by mixing with resin or silica as spectrum transfer material.\textsuperscript{4} However, such PC-LED structure usually suffers from thermal deterioration because of different thermal expansions and conductivities between phosphors and epoxy resin. In order to solve this problem, researchers recently developed glass-ceramics\textsuperscript{5–9} with Eu-doped fluoride crystalline phases. It showed great advantages in thermal resistance, durability, and microstructure manipulation. Eu could enrich in fluoride phase, such as MF\textsubscript{2} (M = Ca, Sr, Ba), of glass-ceramics to adapt itself with well-behaved PL performance.

However, Eu\textsuperscript{2+} and Eu\textsuperscript{3+} always coexist in the host, because Eu cation has variable valences. It thus becomes crucial to keep divalent Eu\textsuperscript{2+} as the majority. Generally, there are several ways to transform Eu\textsuperscript{3+} into Eu\textsuperscript{2+}: (i) adding reducing agents,\textsuperscript{10–12} (ii) inputting reducing atmosphere during preparation processes,\textsuperscript{13,14} (iii) adjusting optical basicity or electronegativity of host,\textsuperscript{15,16} (iv) substituting M\textsuperscript{2+} sites in crystalline lattices.\textsuperscript{17,18} As strategies (i) and (ii) accompany drawbacks such as introducing impurities into the system or leading to infeasibility for elaborated optimization of concentration and distribution of Eu\textsuperscript{2+}, strategies (iii) and (iv) are usually considered to be more executable. Using strategy (iv), silicate,\textsuperscript{19,20} aluminosilicate,\textsuperscript{18,19,21} borophosphate\textsuperscript{22,23} and fluoride crystalline phases\textsuperscript{24–26} have been reported as good stabilizers for divalent Eu\textsuperscript{2+}. Among them, fluoride crystalline phases show most advantages owing to their ultrasmall sizes down to several nanometers, similar refractive indices with silicate glasses, ideal host to selectively enrich Eu\textsuperscript{2+}. Hence, fluorosilicate glass-ceramics containing Eu\textsuperscript{2+}-enriched fluoride nanocrystals could achieve high transparency, large Eu\textsuperscript{3+}/Eu\textsuperscript{2+} reduction ratio, enhanced PL performance and ideal physicochemical properties.

Our previous studies\textsuperscript{27–29} have evidenced that alkaline earth fluoride (MF\textsubscript{2}, where M = Ca; Sr; Ba) nanocrystals could be good hosts to precipitate and enrich Eu\textsuperscript{2+} by Eu\textsuperscript{2+}/M\textsuperscript{2+} lattice substitution. But there is still a lack of systematical dissertation summarizing such lattice substitution strategy for stabilizing...
In this study, we prepared glass and glass-ceramic samples with the composition of 50SiO$_2$–20Al$_2$O$_3$–20MF$_2$–7NaF–3EuF$_3$ (M = Ca, Sr, Ba, Zn) and 50SiO$_2$–20Al$_2$O$_3$–20LaF$_3$–7NaF–3EuF$_3$. By comparative study of the valence state evolution and spectroscopic behaviors of Eu ions, we demonstrate the feasibility of the proposed site substitution strategy to stabilize the divalent Eu$^{2+}$ in the fluorosilicate glass-ceramics containing MF$_2$ nanocrystals.

### Experimental methods

Oxyfluoride glasses (and a glass-ceramic sample) with the compositions of 50SiO$_2$–20Al$_2$O$_3$–20MF$_2$–7NaF–3EuF$_3$ in mol%, named as G1–4, GC5 (M = Ca, Sr, Ba, La, or Zn; x is the valence of ion M) was prepared by a melt-quenching method. Total weight of 50 g raw materials was weighed and mixed well in a RETSCH RM 200 mortar grinder. The batches of the raw materials were melted out on a CDR-1 diaphragm furnace and then quenched in the melt between two brass plates. The subsequent crystallization temperature of 660°C was selected between the first crystallization temperature ($T_{c1}$) and the second crystallization temperature ($T_{c2}$) for G1–4 (Fig. 1(a)). The glass-ceramics G1–4 were obtained by annealing the glasses at the above temperatures for 45 min in air. The compositions and crystallization temperatures of the groups are demonstrated below in Table 1. Note that the ZnF$_2$-based sample (GC5) loses its transparency due to an uncontrollable crystallization during the melt-crystallization process. Differential thermal analysis (DTA) measurement was carried out on a CDR-1 differential thermal analyzer with fixed specimen weight of 60 mg. XRD analysis was carried out on a PANalytical B.V. Empyrean 200895 X-ray diffractometer with Cu Kα radiation (λ = 1.54 Å), and the scan speed was 2.00 deg min$^{-1}$. The excitation and emission spectra combining quantum yields were measured by an Edinburgh Instruments fluorospectrometer FLS 920 equipped with a red-sensitive photomultiplier (Hamamatsu R928P) in a Peltier-cooled housing in the single photon counting mode and with the aid of an integration sphere. A 450 W ozone-free xenon lamp was used as the excitation source for steady-state measurements. The calibrations for quantum yields were made by the Edinburgh Instrument and the measurements were repeated three times at room temperature for each excitation scheme and then averaged overall.

### Results and discussion

#### Transition from glasses to glass-ceramics

Evidenced by differential thermal analysis (DTA, Fig. 1(a and c)) and X-ray diffraction (XRD, Fig. 1(b and d)), sample G1–4 referred to typical inorganic glasses without any crystalline phase, while sample GC5 was glass-ceramic constituted by the precipitated ZnAl$_2$O$_4$ crystalline phases and residual glass phase. Glass samples G1–G4 have two exothermic DTA peaks, where the first peaks (which appeared at lower temperature) correspond to the precipitation of fluoride nanocrystals (CaF$_2$, SrF$_2$, BaF$_2$, LaF$_3$). Such kinds of crystalline phases were revealed by XRD patterns (Fig. 1(d)) of the glass-ceramics, GC1–GC4. These samples were obtained by annealing G1–G4 at 660°C, which were located between two crystallization peaks for 45 minutes. In contrast, the as-melt sample, GC5, was already glass-ceramic with opaque appearance due to the uncontrollable crystallization of ZnAl$_2$O$_4$ during the quenching process of the melt. According to the Scherrer formula, the crystal sizes could be estimated from peak width of XRD patterns. As a result, the sizes of CaF$_2$, SrF$_2$, BaF$_2$, LaF$_3$ and ZnAl$_2$O$_4$ crystals in the glass-ceramics were evaluated as 19.3 ± 0.3 nm, 16.0 ± 0.2 nm, 17.9 ± 2.7 nm and >100 nm, respectively. GC1–4 have high transparency, but GC5 lost its transparency. This is due to much smaller size of precipitated MF$_2$ nanocrystals than the visible-near infrared wavelength, as well as similar refractive indices between MF$_2$ nanocrystals and silicate hosts. So the fluorosilicate glass-ceramics show more optical merits than the reported oxide glass-ceramics.[17–21] The volume fractions of the precipitated nanocrystals (crystallinity) could also be evaluated by the integral intensity ratio of crystalline diffraction to all the diffraction. And the crystallinities were estimated to around 20% for all the glass-ceramic samples. Therefore, the glasses were converted into the glass-ceramics containing CaF$_2$, SrF$_2$, BaF$_2$, LaF$_3$ nanocrystals except the ZnF$_2$-based GC5.

#### Spectroscopic behaviors of Eu$^{2+}$ and Eu$^{3+}$

The sharp PL peaks in Fig. 2(a and b) were assigned to the 4f–4f transitions of Eu$^{3+}$, while the broad PL bands in Fig. 2(c) could

| Sample | Composition (mol%) | $T_{c1}$ | $T_{c2}$ |
|--------|--------------------|----------|----------|
| G1     | 50SiO$_2$–20Al$_2$O$_3$–20CaF$_2$–7NaF–3EuF$_3$ | 602°C | 800°C |
| G2     | 50SiO$_2$–20Al$_2$O$_3$–20SrF$_2$–7NaF–3EuF$_3$ | 598°C | 750°C |
| G3     | 50SiO$_2$–20Al$_2$O$_3$–20BaF$_2$–7NaF–3EuF$_3$ | 597°C | 752°C |
| G4     | 50SiO$_2$–20Al$_2$O$_3$–20LaF$_3$–7NaF–3EuF$_3$ | 650°C | 882°C |
| GC5    | 50SiO$_2$–20Al$_2$O$_3$–20ZnF$_2$–7NaF–3EuF$_3$ | 815°C |         |

Fig. 1  DTA curves and XRD patterns of the G1–G4 glasses (a and b) and the GC1–GC5 glass-ceramics (c and d) with reference to the standard JCPDS cards.
be assigned to $4f^7-4f^65d^1$ transitions of Eu$^{2+}$. For Eu$^{3+}$, the PL excitation peaks of 363, 381, 393 and 414 nm were attributed to the transitions of $7F_0 \rightarrow 5D_4$, $5G_4$, $5L_6$, $5D_3$, respectively. And the PL emission peaks were ascribed to $5D_0 \rightarrow 7F_J$ where $J = 1, 2, 3$ and 4 correspond to 590, 612, 650 and 689 nm, respectively. As the 4f electron shell was shielded by the outer shells (5d and 6s), the 4f-4f transition was less influenced by coordination surroundings, which led the parabolas of 4f$^7$ excited states to be located in parallel and directly above the ground state in the configuration coordinate. Therefore, the band widths of Eu$^{3+}$ PL peaks appeared as narrow linear shapes. On the contrary, the PL bands of Eu$^{2+}$ have large band width covering more than 100 nm with the maxima at 350 and 420 nm. As d shell is exposed to ligands, there was offset between the equilibrium position of both the 4f$^6$5d$^1$ and 4f$^7$ states of Eu$^{2+}$. Therefore, the f-d transition presented large Stokes shift, and electron-phonon coupling further broadened the PL bands. Such a Stokes shift and spectral broadening are depicted in Fig. S1.† It shows that Eu$^{2+}$ has similar PL bands in the glasses and glass-ceramics. But there are large Stokes shifts on the PL excitation and emission spectra. Between the glass and glass-ceramic samples, more or less Stokes difference exists, where GC3 exhibits the largest Stokes shift (4712 cm$^{-1}$) among all the samples. These are attributed to the different coordination environments around Eu$^{2+}$ in the glass and glass-ceramics. In the glass samples, Eu$^{2+}$ is mainly surrounded by both O$^-$ and F$^-$, while Eu$^{2+}$ ions are predominantly enriched in fluoride phase in glass-ceramics. With the enrichment of Eu$^{2+}$ in different nanocrystals, the Stokes shifts of Eu$^{2+}$ also appear to have some differences due to the lattice difference of MF$_2$ (M = Ca, Sr, Ba) nanocrystals.

With configuration coordinates, Fig. S2† illustrates Stokes shift order of $\Delta$stokes(G3) < $\Delta$stokes(GC3) and $\Delta$stokes(G3) < $\Delta$stokes(GC1) < $\Delta$stokes(GC2).

All the glasses and glass-ceramics showed PL feature of both Eu$^{2+}$ and Eu$^{3+}$, except that the LaF$_3$-based samples only exhibited the PL feature of Eu$^{3+}$. For comparison, the PL intensities of each group were normalized according to glasses for Eu$^{3+}$ and glass-ceramics for Eu$^{2+}$. Compared to the glasses, most of the glass-ceramics possessed stronger PL for Eu$^{2+}$, owing to the reduction of Eu$^{3+}$ into Eu$^{2+}$ during the treatment. The Eu$^{2+}$ bands also existed in glass samples and increased by the sequence of G1 to G3 according to Fig. 2(c). The appearance of PL bands implied that reduction of Eu$^{3+}$ started in the glass forming stage. However, the PL of Eu$^{3+}$ displayed intricate changes before and after heat treatment. The Eu$^{3+}$ PL peaks of CaF$_2$ and LaF$_3$ based glass-ceramics (GC1 and GC4) strengthened after heat treatment, while the PL peaks of SrF$_2$ and BaF$_2$ based glass-ceramics (GC2 and GC3) were weaker than those of the glasses. In addition, sample G4, GC4 and GC5 almost had no intense PL bands of Eu$^{2+}$ according to Fig. 2(c), which indicates the absence of Eu$^{2+}$ in those samples. The evolution of PL intensity could be also evidenced by the integral sphere collected spectra (Fig. 3(a and b)). We believe that the PL behavior of the material was mainly influenced by: (i) the ratio diversification of Eu$^{3+}$/Eu$^{2+}$ due to the Eu$^{3+}$ → Eu$^{2+}$ reduction; (ii) the coordination of Eu$^{2+}$ into the precipitated crystalline phases. These two factors will be discussed in subsequent passages.
Electronegativity related reduction of Eu³⁺ → Eu²⁺ in the glasses

The reduction of Eu³⁺ → Eu²⁺ in glass is usually believed to relate with the optical basicity. This parameter was primarily used to describe the “electron donor ability” of oxygen as well as the concentration of non-bonding oxygen.²⁸,²⁹ From Fig. 2(c), the PL of Eu²⁺ increased by the consequence of G1 < G2 < G3. However, this trend contradicted the well-developed theory of optical basicity that the lower optical basicity (A(Ca²⁺) < A(Sr²⁺) < A(Ba²⁺)) determined the higher reduction ratio of Eu³⁺ (G1 > G2 > G3).³⁵ (Herein, due to relatively small concentration of Eu²⁺, it could be assumed to still be lower than the PL quenching concentration and thus be correlated with PL intensity.) We believed that the contradiction was primarily because the investigated glasses were fluoride and oxide mixed glass system. Oxide (silicate) glass is covalent bond majored “random tetrahedral network” and linked by sharing corners, while fluoride glass is ionic bond majored “random polyhedral packing” and linked by sharing corners, edges and planes. In the high fluoride content (50 mol%) mixed system, we have revealed that the glass was indeed constructed by separated fluoride-rich glass phases and silicate-rich glass phases.³⁰ Thus, the Eu³⁺ → Eu²⁺ reduction could not be well interpreted by optical basicity, but we tried to relate it with “electronegativity”, which was the primary origin of “optical basicity”.

Electronegativity, χ, is a chemical property that describes the tendency of an atom or ion to attract electron clouds towards itself.³¹ Due to the electronegativity differences between Eu³⁺ and other cations, Eu³⁺ tended to attract more electron density around its neighboring cations with lower electronegativity, and eventually captured an electron from fluoride. Thus, the Eu³⁺ → Eu²⁺ reduction was more likely to happen where electronegativity of neighboring cations of Eu³⁺ were lower. In the glass samples, there were indeed two kinds of separated glass phases: fluoride-rich glass phases and silicate-rich glass phases.³² It has been well evidenced that lanthanides, such as europium, could selectively enrich in fluoride rich phases.³³ According to the “random polyhedral packing” model, [AlF₆] coordination octahedra could form the skeleton of fluoride rich glass phases, and other metal ions such as Ca²⁺, Sr²⁺, Ba²⁺ and Eu²⁺/³⁺ filled the pores of [AlF₆] framework.³⁴ In G1–G3, the electronegativity decreases as per the sequence χ(Ca²⁺) > χ(Sr²⁺) > χ(Ba²⁺), Eu²⁺ ions were more likely to form in the latter groups. In G4, χ(Eu³⁺) is close to χ(La³⁺), and in GC5 χ(Zn²⁺) is much larger than χ(Eu²⁺), so only slight amounts of Eu²⁺ were found in G4 and no Eu²⁺ was detected in GC5, as shown in Fig. 2(c).

Stabilization of Eu²⁺ in the glass-ceramics via lattice site substitution

Lattice site substitution has been considered as a facile Eu²⁺ doping strategy to avoid other impurities, uncontrollable redox reaction and large aggregation. The conversion ratio of reduction through this method was mainly influenced by two crucial factors: the charge and radius of the site. In order to form stable divalent Eu²⁺, the lattice sites should meet the following criteria: (i) the sites should be originally divalent charged, (ii) the radii of the sites should be close to the radii of Eu²⁺.³⁴,³⁵

The site charge requisition of Eu²⁺ substitution could be examined by comparison between the MF₂ (M = Ca, Sr, Ba)-based (G1, G2, G3) and the LaF₃-based (G4) samples. The crystalline phases of G4 were LaF₃. The trivalent La³⁺ sites rather than divalent ones led to no transition of Eu³⁺ to Eu²⁺ and absence of the Eu²⁺ PL (Fig. 2(c)), although the radii of La³⁺ was close to Eu²⁺ (RᵥLa³⁺(1.160 Å) < RᵥEu²⁺(1.25 Å)). Thus, the divalent sites were required for reduction. As previously reported, the mechanism of the Eu³⁺ → Eu²⁺ reduction was expressed by a series of defect equations. When Eu³⁺ enters the lattice containing M²⁺ with similar radii, a cation vacancy with two negative charges is formed

\[
3M^2_+ + 2Eu^3+ → 3M^2_+ + 2Eu^{2+} + 2e^-
\]

(1)

where M^2_+ stands for metal ion which occupied its own lattice, Eu^3+ for the doping Eu³⁺ ion which occupied the lattice of metal ion with one extra positive charge, and V^−_M for the vacancy of metal ion with two negative charges.

As the vacancy act as electron donor, Eu³⁺ was reduced to Eu²⁺,

\[
2V^−_M → 2M^2_+ + 2e^−
\]

(2)

\[
2e^− + 2Eu^{2+} → 2Eu^{3+}
\]

(3)

where V^−_M stands for the vacancy of metal ion with no extra charge, Eu^3+ for the doping Eu³⁺ with no extra charge.

These substitution effects were illustrated in Fig. 4. Accordingly, the Eu³⁺ → Eu²⁺ reduction took place and alkaline earth cation vacancies formed in GC1–3 (Fig. 4(a–c)), and broad PL bands of Eu²⁺ were found. The lattice constants were calculated by Bragg’s Law, as listed in Table 2. The site substitution can be further verified according to lattice constant change. The expansion of CaF₂ lattice in GC1 corresponded with smaller radius of Ca³⁺(1.12 Å) compared to that of Eu²⁺(1.25 Å), and the shrinkage of SrF₂/BaF₂ lattice in GC2 and GC3 for larger radii of Sr²⁺(1.26 Å) and Ba²⁺(1.42 Å). Also, the Eu³⁺/La³⁺ substitution took place in LaF₃ lattice of GC4 without the Eu³⁺ → Eu²⁺ reduction. The very small deviation from standard lattice constant of LaF₃ in GC4 was due to the very similar radii of La³⁺(1.160 Å) and Eu³⁺(1.066 Å).

When the site was divalent charged, the radius difference between substitution cation pairs would play a prime role for the Eu³⁺ → Eu²⁺ reduction. By reference to ZnF₂-based sample (GC5), the influence of the radius difference on the lattice site substitution could be clearly observed. The ionic radii of with 8 coordination number were demonstrated as below: RᵥZn²⁺(0.60 Å) < RᵥEu³⁺(1.066 Å) < RᵥCa²⁺(1.12 Å) < RᵥEu²⁺(1.25 Å) < RᵥBa²⁺(1.42 Å). Due to the huge difference between doping cations (RᵥEu³⁺(1.066); RᵥEu²⁺(1.25 Å)) and lattice cations (RᵥZn²⁺(0.60 Å)), a barrier was created in the Eu³⁺/Zn²⁺ substitution. Moreover, less deviation of ZnAl₂O₄ lattice constant in GC5 also confirmed the above theory. Furthermore, the radius differences between Eu³⁺ and M²⁺ sites would determine how much Eu³⁺ would be reduced into Eu²⁺. The PL intensity of Eu²⁺ increased by the order of GC1 < GC2 < GC3, while that of Eu³⁺...
decreased by GC1 > GC2 > GC3 (Fig. 2(b)). Thus, the ratio of Eu\(^{2+}/Eu^{3+}\) should be GC1 < GC2 < GC3 assuming that the concentrations of Eu\(^{2+}\) and Eu\(^{3+}\) were lower than the PL quenching concentrations. The Eu\(^{3+} \rightarrow Eu^{2+}\) reduction was indeed driven by the radius difference between different Eu\(^{3+}/M^{2+}\) pairs. On the one hand, for Eu\(^{3+}/Ca^{2+}\) substitution with small radius difference (Ca\(^{2+}(1.12 \text{ Å})\) vs. Eu\(^{3+}(1.066 \text{ Å})\)) and Ca\(^{2+}\) sites with smaller radii than Eu\(^{2+}\) (1.25 Å), it only drove Eu\(^{3+}\) to be partially reduced to Eu\(^{2+}\), so Eu\(^{3+}/Eu^{2+}\) coexisted in GC1. The inversion of PL peak intensities at 590 nm and 612 nm for G1 (Fig. 2(a)) and GC1 (Fig. 2(b)) well verified the CaF\(_2\) crystalline coordination environment of Eu\(^{3+}\) in GC1. The PL enhancement of Eu\(^{3+}\) (Fig. 2(b)) was due to lower phonon energy and lower multi-phonon transition ratio of Eu\(^{3+}\) in CaF\(_2\) lattice of GC1. As both the Eu\(^{3+}\) and Eu\(^{2+}\) were enriched in the CaF\(_2\) nano-crystals, the strengthened cross relaxation transition between Eu\(^{2+}/Eu^{3+}\) coincidently eliminated the weakening of multi-phonon transition, so the PL lifetime of GC1 had no significant change compared to G1 (Fig. 5 and Table 3). On the other hand, Eu\(^{3+}/Ba^{2+}\) or Eu\(^{3+}/Sr^{2+}\) substitution with large radius difference (Sr\(^{2+}(1.26 \text{ Å})/Ba^{2+}(1.42 \text{ Å})\) vs. Eu\(^{3+}(1.066 \text{ Å})\)) and Sr\(^{2+}/Ba^{2+}\) sites with larger radii than Eu\(^{2+}\) (1.25 Å) drove more Eu\(^{3+}\) to be reduced to Eu\(^{2+}\). Accordingly, large amounts of Eu\(^{3+}\) remained trivalent in GC1, while small amounts remained in GC2 and GC3. No inversion of PL peak intensities at 590 nm and 612 nm for G2 and G3 (Fig. 2(a)) and GC2 and GC3 (Fig. 2(b)) implied that all Eu\(^{3+}\) in GC2 and GC3 corresponded to those remaining in the glassy phase, which exhibited much shorter PL lifetimes (Fig. 5; Table 3).

Quantum yield (QY) improvement and PL enhancement of Eu\(^{3+}\)

After heat treatment, large quantities of Eu\(^{2+}\) formed by Eu\(^{3+}/M^{2+}\) substitution in GC1–GC3, QY of the Eu\(^{2+}\) PL (Fig. 3) was improved to about 4–7 times of QY of precursor glasses. QY is dominated by two competitive processes: radiative transition (RT) and non-radiative transition (NRT). Theoretically, the probability of RT and NRT, \(W_{RT}\) and \(W_{NRT}\) have the following relationships:

\[
\eta = \frac{W_{RT}}{W_{RT} + W_{NRT}} = \tau W_{RT} \tag{5}
\]

where \(\eta\) and \(\tau\) represent QY and lifetime of PL, respectively. Accordingly, the QY improvement depended on lower NRT or longer PL lifetime. Eu\(^{3+}\) ions were mainly enriched in the separated fluoride glass phases filled with large content of quenching defects, unsaturated bonds and interfaces. Thus,

![Fig. 4](image)

**Fig. 4** Schematic model to describe site substitution of Eu\(^{2+}\) or Eu\(^{3+}\) in different primitive cells. (a–c) Eu\(^{2+}\)-doped cubic MF\(_2\) (M = Ca, Sr, Ba) cell, (d) Eu\(^{3+}\)-doped hexagonal LaF\(_3\) cell, (e) Eu\(^{3+}\)-doped cubic ZnAl\(_2\)O\(_4\) cell.
Eu\textsuperscript{2+} in the glasses had high \( W_{\text{RT}} \) and low QY (<10%), which was also evidenced by the shorter lifetimes of Eu\textsuperscript{2+} (Fig. 5(c)). On the contrary, Eu\textsuperscript{3+} in the glass-ceramics were incorporated in MF\textsubscript{2} crystalline phases, where the concentration of PL quenching centers were much lower than those in glass phases. Therefore, PL lifetimes of Eu\textsuperscript{2+} in the glass-ceramics (Fig. 5(d)) appeared much longer than those of the precursor glasses. It supports the glass-ceramics, GC1–GC3, achieved much higher QY of the Eu\textsuperscript{2+} PL.

Besides a high QY value, strong PL still requires high concentration of the active center (Eu\textsuperscript{3+}) without serious concentration quenching. The BaF\textsubscript{2}-based glass-ceramics possessed high solubility of Eu\textsuperscript{2+} rather than SrF\textsubscript{2} or CaF\textsubscript{2}-based glass-ceramics. Larger radii as well as large radius difference were found to play almost equal roles when reducing Eu\textsuperscript{3+} into Eu\textsuperscript{2+}. Therefore, the BaF\textsubscript{2}-based rather than the SrF\textsubscript{2}-based glass-ceramics got the highest QY and the strongest PL, simultaneously. It also demonstrates that the proposed MF\textsubscript{2} lattice site substitution method enabled the facile Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction and stabilization of Eu\textsuperscript{2+} in the fluorosilicate glass-ceramics. The Eu tended to enrich in fluoride-rich phases of the glasses and in the fluoride nanocrystalline phases of the glass-ceramics. In glasses, the Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction depended on the electronegativity of fluoride glass phases, while in glass-ceramics, the Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction depended on the lattice site substitution in fluoride crystalline phases. The lattice site substitution depended on two key factors: the radii and the valency difference between sites and substitution ions. The similar radii between sites and ions were the prerequisites of the doping. Too small radii of doping ions (like Zn\textsuperscript{2+}) were not preferred for Eu\textsuperscript{3+}/M\textsuperscript{2+} substitution, while comparably larger radii (like Ba\textsuperscript{2+}) were more facile for both Eu\textsuperscript{3+}/M\textsuperscript{2+} substitution and Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction. And to some extent, larger the radii, easier it was for the Eu\textsuperscript{3+}/M\textsuperscript{2+} substitution to take place. The unbalanced charge at the substitution sites drove the reduction of Eu\textsuperscript{3+} → Eu\textsuperscript{2+}. The Eu\textsuperscript{3+}/M\textsuperscript{2+} substitution was attributed to the Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction, while Eu\textsuperscript{3+}/La\textsuperscript{3+} substitution led to almost no reduction of Eu\textsuperscript{3+}. High Eu\textsuperscript{2+} doping concentration and enhanced luminescent QY of Eu\textsuperscript{2+} were simultaneously achieved in BaF\textsubscript{2}-based glass-ceramic.

**Conclusions**

The separated fluoride glass phases formed in the precursor fluorosilicate glasses transformed into fluoride nanocrystals in the glass-ceramics after heat treatment. The inclined precipitation of alkaline earth fluoride nanocrystals and the Eu\textsuperscript{3+}/M\textsuperscript{2+} lattice site substitution enabled the facile Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction and stabilization of Eu\textsuperscript{2+} in the fluorosilicate glass-ceramics. The Eu tended to enrich in fluoride-rich phases of the glasses and in the fluoride nanocrystalline phases of the glass-ceramics. In glasses, the Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction depended on the electronegativity of fluoride glass phases, while in glass-ceramics, the Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction depended on the lattice site substitution in fluoride crystalline phases. The lattice site substitution depended on two key factors: the radii and the valency difference between sites and substitution ions. The similar radii between sites and ions were the prerequisites of the doping. Too small radii of doping ions (like Zn\textsuperscript{2+}) were not preferred for Eu\textsuperscript{3+}/M\textsuperscript{2+} substitution, while comparably larger radii (like Ba\textsuperscript{2+}) were more facile for both Eu\textsuperscript{3+}/M\textsuperscript{2+} substitution and Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction. And to some extent, larger the radii, easier it was for the Eu\textsuperscript{3+}/M\textsuperscript{2+} substitution to take place. The unbalanced charge at the substitution sites drove the reduction of Eu\textsuperscript{3+} → Eu\textsuperscript{2+}. The Eu\textsuperscript{3+}/M\textsuperscript{2+} substitution was attributed to the Eu\textsuperscript{3+} → Eu\textsuperscript{2+} reduction, while Eu\textsuperscript{3+}/La\textsuperscript{3+} substitution led to almost no reduction of Eu\textsuperscript{3+}. High Eu\textsuperscript{2+} doping concentration and enhanced luminescent QY of Eu\textsuperscript{2+} were simultaneously achieved in BaF\textsubscript{2}-based glass-ceramic.

**Conflicts of interest**

There are no conflicts to declare.

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