Title

Preparation and luminescent properties of Eu-doped transparent mica glass-ceramics

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Abstract

Eu-doped transparent mica glass-ceramics were prepared, the influence of Eu-doping on the crystallization of the parent glasses was investigated and the luminescent properties of the parent glasses and the glass-ceramics were estimated. A small additive amount of Eu element was very effective in preparing transparent mica glass-ceramics. However, the excess addition led to the coarsening of phase separation in the glass phase and the separation of unidentified crystal phases and β-eucryptite during heating of the parent glasses, which caused white opaque at lower heating temperatures. When mica crystals were separated, Eu ions entered the interlayers of mica crystals. The observed emission and excitation spectra showed that parts of Eu$^{3+}$ ions which were added as Eu$_2$O$_3$ were reduced to Eu$^{2+}$ ions during melting of the starting materials and heating the parent glasses in air and the energy transfer from Eu$^{2+}$ to Eu$^{3+}$ ions occurred.

Keyword: Optical properties; Glass; Glass ceramics; Silicate
1. Introduction

Europium ions (Eu$^{2+}$ and Eu$^{3+}$) are well known as activators in phosphors. The Eu$^{2+}$-doped phosphors emit blue light, which originated in $4f^65d \to 4f^7$ transition of Eu$^{2+}$ ions and the Eu$^{3+}$-doped phosphors emit red light, which originated in $5D_0 \to 7F_J$ transition of Eu$^{3+}$ ions. For example, BaMgAl$_{10}$O$_{17}$: Eu$^{2+}$ and (Y, Gd)BO$_3$: Eu$^{3+}$ are used as blue and red phosphors, respectively, for plasma display panels. So such phosphors activated with europium ions have been widely used in other commercial applications and also have been investigated.

Transparent glass-ceramics are interesting hosts for phosphors used for white light emitting diodes, laser materials and so on. Fluoride crystals and glasses are desirable hosts for optically active rare-earth cations because of their low phonon energies and broad transparency in the infrared region of spectrum used for amplification of light in telecommunications systems [1]. And transparent oxyfluoride glass-ceramics, based on fluoride crystals dispersed throughout a continuous silicate glass, have been shown to combine the optical advantages of rare-earth-doped fluoride crystals with the ease of forming and handling of conventional oxide glasses [1]. On the other hand, we succeeded in preparing novel transparent and machinable mica glass-ceramics from oxyfluoride glasses [2, 3]. The separated fluorine-micas had size of < 50 nm and their interlayer cations were lithium ions. The transparent fluorine-mica glass-ceramics are interesting host materials for phosphors, like transparent oxyfluoride glass-ceramics.

In this study, Eu$_2$O$_3$ was added to the starting materials of the transparent mica glass-ceramics. Eu-doped parent glasses, which were oxyfluoride glasses, were prepared by melting the starting materials in sealed platinum container and were crystallized by heating in air. Then, the influence of Eu-doping on the crystallization of the parent glasses was investigated and the luminescent properties of the parent glasses and the
obtained glass-ceramics were estimated.

2. Experimental procedure

Novel transparent and machinable mica glass-ceramic in which a large quantity of mica was separated was prepared from the parent glass with the chemical compositions corresponding to 94.9 mass% \( \text{Li}_{1.5}\text{Mg}_3\text{AlSi}_{13.25}\text{O}_{25}\text{F}_2 + 5.1 \text{mass}\% \text{MgF}_2 \) in previous study [3]. The composition of base glass in this study was the same with that of the mica glass-ceramic. 0, 0.05, 0.5, 5, 10 and 20 mol% \( \text{Eu}_2\text{O}_3 \) were added to the base glass. Such specimens were shown as E0, E005, E05, E5, E10 and E20, respectively, according to the additive amount of \( \text{Eu}_2\text{O}_3 \), in this paper. The Eu containing glasses were prepared as follows. The reagents of \( \text{MgO} \), \( \text{Al}_2\text{O}_3 \), \( \text{SiO}_2 \), \( \text{Eu}_2\text{O}_3 \), \( \text{Li}_2\text{CO}_3 \) and \( \text{MgF}_2 \) were mixed, calcined at 900 \( ^\circ\text{C} \) for 1 h, melted in a sealed platinum container at 1450 \( ^\circ\text{C} \) for 2 h, and then cooled outside of the furnace. The obtained glasses were annealed in air at about 20 \( ^\circ\text{C} \) higher temperatures than their glass-transition temperatures and cooled at 2 \( ^\circ\text{C}/\text{min} \) to eliminate strain. The parent glasses prepared by such a method were cut to dimensions of about 5 mm \( \times \) 5 mm \( \times \) 1 mm and were heated at 600-800 \( ^\circ\text{C} \) for 1 h in air to be crystallized. The heating rate was 10 \( ^\circ\text{C}/\text{min} \).

The thermally change of the parent glasses was analyzed using an X-ray diffraction (XRD) analyzer and a differential thermal analyzer (DTA). The microstructures of the parent glasses and the glass-ceramics were observed using a field emission type scanning electron microscope (FE-SEM) and a transmission electron microscope (TEM). The luminescent properties of the parent glasses and glass-ceramics with thickness of 0.4 mm were analyzed at room temperature using a fluorescence spectrophotometer.
3. Results and discussion

3.1. Influence of Eu-doping on crystallization

The photographs of the parent glasses and the heated glasses are shown in Fig. 1. Every parent glass was transparent. While the parent glasses to which 0-5 mol% Eu₂O₃ were added were colorless, those to which 10 mol% and 20 mol% Eu₂O₃ were added were light orange color. Crystals were not detected in every parent glass by XRD analysis. SEM photographs of the polished and etched surfaces of the E05 and E10 parent glasses are shown in Fig. 2. In the both glasses, droplet phase, which shows a binodal phase separation, was observed. The size of droplet phase in the E05 parent glass was 50-60 nm and that in the E10 parent glass was 20-30 nm. In every parent glass, such a binodal phase separation was observed, and the sizes of these droplet phases varied with the additive amount of Eu₂O₃.

The phase change in the E5 specimen by heating is shown in Fig. 3. Trace of chondrodite (Mg₅(SiO₄)₂F₂) was observed at 600 °C and a larger amount of chondrodite appeared at 650 °C. Next, mica was separated at 700 °C. The chondrodite acted as nuclei for mica. β-eucryptite was observed at 750 °C. (This β-eucryptite was β-eucryptite solid solution [2].) A large amount of β-eucryptite appeared at 800 °C. The diffraction peak at about 31 ° which was observed at 750 °C and 800 °C could not be identified. Probably, it originated in a compound containing Eu. The E5 specimen maintained transparency (Fig. 1(c)) though chondrodite and mica crystals were separated at 600-700 °C. However, it turned white opaque slightly with the separation of β-eucryptite at 750 °C. Such crystallization process was similar to that of the E0 specimen [3], except for the appearance of the unidentified diffraction peaks.

All specimens, except the E20 specimen, first separated chondrodite at 600 °C and the amount of the chondrodite increased at 650 °C. The E20 specimen showed only
trace of chondrodite in the XRD pattern at 650 °C, and already turned white opaque (Fig. 1 (e)). The E10 specimen also turned a little white opaque at 650 °C (Fig. 1(d)). This white opaque might be caused by the coarsening of the phase separation in the glass phase. XRD patterns of the E0-E20 specimens heated at 700 °C for 1h are shown in Fig. 4. It is obvious that Eu₂O₃ addition promoted the separation of mica crystals, compared the XRD patterns of Eu containing specimens with the XRD pattern of the E0 specimen. E005, E05 and E5 specimens separated a larger amount of mica crystals at 700 °C and maintained the transparency (Fig. 1). The E005 and E05 specimens did not separate β-eucryptite and unidentified crystal phases (particularly, the crystal phase of which diffraction peak appeared at around 31 °) at 700 °C. The E5 specimen showed only trace of their crystals in the XRD pattern and their diffraction peak intensities became stronger with an increase in Eu content. Consequently, the E10 and E20 specimens turned more white opaque at 700 °C (Fig. 1 (d) and (e)).

Above results show that a small addition of Eu₂O₃ was very effective in preparing transparent glass-ceramics in which a large amount of mica crystals was separated. The Eu₂O₃ addition might vary the composition of the droplet and matrix phases in the parent glasses and probably resulted in F- richer droplet phase and MgO- richer matrix phase, which caused the promotion of separation of mica [3]. However, the excess addition of Eu₂O₃, such as E10 and E20 specimens, led to the coarsening of phase separation in the glass phase and the separation of a larger amount of the unidentified crystal phases and β-eucryptite during heating of the parent glasses, which caused white opaque at lower heating temperatures.

TEM images and EDX spectra of the E5 specimen heated at 700 °C are shown in Fig. 5. Many mica crystals with size of <150 nm were separated. A lot of dark parts were observed in the TEM images. Then, the EDX spectra were measured on the dark
site (M) and the light site (G) which were the mica crystal and the glass phase, respectively. Eu peaks appeared together with Mg, Al, Si, O and F peaks in the EDX spectrum of M. (Cu and C peaks originated in the sample holder and the contamination, respectively.) However, no Eu peaks were observed in the EDX spectrum of G though Mg, Al, Si, O and F peaks appeared. That is, the dark parts, which were the mica crystals, contained Eu element and the light parts, which were glass phase, did not contain the Eu element. Moreover, the dark straight lines were observed in some mica crystals, which means that all layers of mica crystals did not always contain Eu element uniformly and there were some layers concentrated Eu element in a mica crystal.

The basal spacing ($c \cdot \sin \beta$; $c$ and $\beta$ are lattice constants) and lattice constant $b$ of the mica separated in the E0-E5 specimens heated at 700 °C were determined from the (003) and (060) diffraction peaks, respectively. The obtained basal spacing and lattice constant $b$ are shown in Fig. 6. In general, the basal spacing, which is the distance between basal planes in the vertical direction relative to the layer, depends mainly on the size of the interlayer cation and the electrostatic repulsion between layers. Furthermore, the lattice constant $b$ is a standard for determining whether the mica is the tetrasilicic or trisilicic type and the $b$ of tetrasilicic type mica is smaller than that of trisilicic type mica. The composition of mica separated in the E0 specimen was $\text{Li} (\text{Mg}_{2+y} \text{Li}_{1-y})(\text{Al}_y \text{Si}_{4-y})\text{O}_{10} \text{F}_2$ ($0 < y < 1$) [3]. The interlayer cations were $\text{Li}^+$ ions, the octahedral sheets contained $\text{Li}^+$ and $\text{Mg}^{2+}$ ions and the tetrahedral sheets contained $\text{Al}^{3+}$ and $\text{Si}^{4+}$ ions. However, $\text{Li}^+$ ions are too small to be placed in the interlayer of twelve-coordination. Therefore, if parts of Li ions in the interlayer are substituted by Eu element ($\text{Eu}^{2+}$ and/or $\text{Eu}^{3+}$ ions) which have larger size than the other cations in the specimen, the structure of the micas will become more stable. Actually, it is clear from Fig. 5 that the mica crystals contained Eu element. The Eu element ($\text{Eu}^{2+}$ and/or $\text{Eu}^{3+}$
ions) must be substituted for Li$^+$ ions in the interlayer of mica to stabilize mica structure. However, the basal spacing and lattice constant b of the mica separated in the Eu containing specimens were almost the same with those of the mica separated in the E0 specimen. The basal spacing should be increased by substituting larger cations for Li$^+$ ions in the interlayer. On the other hand, the number of interlayer cations was decreased by substituting divalent and/or trivalent cations for Li$^+$ ions in the interlayer, which should caused the decrease in the basal spacing. Consequently, substituting Eu$^{2+}$ and/or Eu$^{3+}$ ions for Li$^+$ ions in the interlayer did not vary not only the lattice constant b but also the basal spacing.

3.2. Luminescent properties

Emission spectra of the E5 specimen excited at 254 nm are shown in Fig. 7. The peaks at around 590 nm and 610 nm were observed for the E5 parent glass, which originated in $^5D_0 \rightarrow ^7F_J$ transition of Eu$^{3+}$ ions. In addition, the weak, broad and discontinuous peak at around 400 nm was also observed, which originated in $4f^{6}5d \rightarrow 4f^{7}$ transition of Eu$^{2+}$ ion. These results mean that parts of Eu$^{3+}$ ions which were added as Eu$_2$O$_3$ were reduced to Eu$^{2+}$ ions during melting of the starting materials. While the peaks at around 590 nm and 610 nm became weaker with an increase in heating temperature, the broad peak at around 400 nm became stronger, especially, the peak was stronger than the peaks at around 590 nm and 610 nm at 700 °C. However, all peaks became much weaker at 750 °C because the E5 specimen turned white opaque. In this way, Eu$^{3+}$ ions were reduced to Eu$^{2+}$ ions during heating in air and an amount of Eu$^{2+}$ ions was increased with an increase in heating temperature. So the E5 parent glass emitted red light by the excitation at 254 nm, and the E5 specimens heated at 700 °C and 750 °C emitted violet light. Similar luminescence properties were observed for
other Eu containing specimens in this study.

The reduction of Eu$^{3+}$ to Eu$^{2+}$ ions in air has been reported for some inorganic phosphors [11-21]. It has been observed for SrF$_3$ nanocrystals and Eu containing glass-ceramics [11], Eu-doped ZnO-B$_2$O$_3$-P$_2$O$_5$ glasses [12], CaB$_2$O$_4$: Eu [13], M$_2$B$_5$O$_9$Cl: Eu (M=Ca, Sr, Ba) [14], Sr$_4$Al$_{14}$O$_{25}$: Eu [15], BaMgSiO$_4$: Eu [16], LnBPO$_5$: Eu (Ln=Ca, Sr) [17], SrB$_5$O$_9$Cl: Eu [18], and SrB$_4$O$_7$: Eu [19]. Their reduction mechanism is explained as follows [11-19]. When three M$^{2+}$ ions such as Ca$^{2+}$, Sr$^{2+}$, Ba$^{2+}$ and Zn$^{2+}$ ions are replaced by only two Eu$^{3+}$ ions, a negative vacancy ($V_M^{''}$) is formed ($3M^{x^+}_M + 2Eu^{3+} \rightarrow 3M^{2+}_M + V_M^{''} + 2Eu^{3+}_M$). The negative charges in the vacancy ($V_M^{''}$) are transferred to the Eu$^{3+}$ ions sites (2Eu$^{3+}_M$) and reduce Eu$^{3+}$ to Eu$^{2+}$ ions ($V_M^{''} \rightarrow V_M^{x^+} + 2e$, $2Eu^{3+}_M + 2e \rightarrow 2Eu^{2+}_M$). On the other hand, the reduction is observed when the glasses with the compositions of BaB$_2$O$_3$: Eu [20] and MB$_4$O$_7$: Eu (M=Sr, Ba) [21] are crystallized. The reduction mechanism is explained as follows [20, 21]. When the non-bridging oxygens of BO$_3$ units in such glasses are connected one another to form BO$_4$ units in crystallization process of the glasses, Eu$^{3+}$ ions are reduced by evolving the resulting free oxygens and the converted Eu$^{2+}$ ions are stabilized by occupaying the Sr$^{2+}$ or Ba$^{2+}$ ions sites. Our specimens contained Mg$^{2+}$ ions as the divalent cations but the size of Mg$^{2+}$ ion was too small to be replaced by Eu$^{3+}$ ions, moreover, our parent glasses did not contain BO$_3$ units. Therefore, the reduction mechanism of Eu$^{3+}$ to Eu$^{2+}$ ions in this study should be different from above two explanations. A feature of the glasses prepared in this study is that fluorine and/or fluorides evaporated during heating. We reported that fluorine and fluorides such as AlF$_3$ evaporated from the glass with composition of KMg$_3$AlSi$_3$O$_{10}$F$_2$ (fluorophlogopite) at $> 800$ °C [22]. That is, the reduction of Eu$^{3+}$ to Eu$^{2+}$ ions in this study might be caused by evolving fluorine from the melt and glass phase. When the starting materials were melted in the sealed
platinum container, fluorine might evaporate from the melt, which resulted in the reduction of Eu\(^{3+}\) ions. Furthermore, when the parent glasses were heated in air to be crystallized, particularly, at \(\geq 700 \, ^\circ\mathrm{C}\), a larger amount of fluorine might evaporate from the glass phases and a larger amount of Eu\(^{3+}\) ions were reduced. Since the specimens in this study contained a larger amount of fluorine than the glass with composition of KMg\(_3\)AlSi\(_3\)O\(_{10}\)F\(_2\) (fluorophlogopite) [22], fluorine might evaporate at \(< 800 \, ^\circ\mathrm{C}\).

The excitation spectrum (emission spectrum: 405 nm) and emission spectrum (excitation spectrum: 289 nm) of the E5 parent glass are shown in Fig. 8. The excitation peak was observed at 289 nm, which ascribed to the 4f-5d transition of Eu\(^{2+}\) ions [12]. When the parent glass was excited at 289 nm, the broad emission peak which was similar to that shown in Fig. 7(d) appeared at around 400 nm. And it was discontinuous peak, especially at 394 nm. The excitation spectrum (emission spectrum: 610 nm) and emission spectrum (excitation spectrum: 394 nm) of the E5 parent glass are shown in Fig. 9. The some sharp excitation peaks were observed at 350-550 nm, which ascribed to the transitions between the 4f energy levels of Eu\(^{3+}\) ions [11, 12], particularly, the excitation peak at 394 nm was the strongest. When the parent glass was excited at 394 nm, the emission peak appeared at around 590 and 610 nm, which originated in Eu\(^{3+}\) ions. These results indicate that Eu\(^{3+}\) ions absorbed the parts of the emission of Eu\(^{2+}\) ions, particularly the light of 394 nm, were excited by the light adsorbed from the emission of Eu\(^{2+}\) ions and then emitted the light of around 590 nm and 610 nm. Consequently, the emission peak at around 400 nm of Eu\(^{2+}\) ions results in the discontinuous, particularly at 394 nm, and is seemed to be the broad emission containing several peaks, as shown in Fig. 7 and Fig. 8, though it is one continuous broad emission peak originally by the 4f\(^6\)5d \(\rightarrow\) 4f\(^7\) transition of Eu\(^{2+}\) ion. That is, these results mean that the energy transfer from Eu\(^{2+}\) ions to Eu\(^{3+}\) ions occurred. The energy
corresponding to the $4f^65d \rightarrow ^8S_{7/2}$ transition of Eu$^{2+}$ ions caused to excite Eu$^{3+}$ ions [23, 24], particularly, $^7F_0 \rightarrow ^5L_6$ transition which is corresponding to energy level of 394 nm.

Changes in emission intensities at 405 nm and 610 nm of the parent glasses and the specimens heated at 700 °C with an increase in Eu$_2$O$_3$ content are shown in Fig. 10. The emission intensities at 405 nm and 610 nm of the parent glasses were decreased by addition of 20 wt% Eu$_2$O$_3$. That is, the concentration quenching did not occur even for the parent glass having such high Eu$_2$O$_3$ concentration as 10 mol%. On the other hand, the emission intensities of the heated specimens were decreased by addition of 10 mol% Eu$_2$O$_3$, which might not result from concentration quenching. As the additive amount of Eu$_2$O$_3$ increased, white opaque of the specimens occurred at lower heating temperatures. That is, such decreases in the intensities resulted from the white opaque.

4. Conclusions

We fabricated the novel phosphors which were transparent and machinable mica glass-ceramics containing Eu$^{2+}$ and Eu$^{3+}$ ions. And the influence of Eu$_2$O$_3$ addition on the crystallization of the parent glasses was investigated and the luminescent properties of the parent glasses and the glass-ceramics were estimated.

A small additive amount of Eu$_2$O$_3$ was very effective in preparing transparent mica glass-ceramics. However, the excess addition led to the coarsening of phase separation in the glass phase and the separation of a larger amount of unidentified crystal phases and β-eucryptite during heating of the parent glasses, which caused white opaque at lower heating temperatures. When the mica crystals were separated, Eu ions (Eu$^{2+}$ and/or Eu$^{3+}$) were substituted for Li$^+$ ions in the interlayer of mica crystals. However, all layers of mica crystals did not always contain the Eu ions uniformly and there were some layers which concentrated the Eu ions in a mica crystal.
When the parent glasses excited at 254 nm, emission peaks appeared at around 590 nm and 610 nm, which originated in Eu\(^{3+}\) ions and the weak, broad and discontinuous emission peak appeared at around 400 nm, which originated in Eu\(^{2+}\) ion. While the former peaks became weaker by heating in air, the latter peak became stronger.

It was obvious from the emission spectra that parts of Eu\(^{3+}\) ions were reduced to Eu\(^{2+}\) ions during melting of the starting materials and during heating in air to be crystallized. This reduction of Eu\(^{3+}\) to Eu\(^{2+}\) ions might be caused by evolving fluorine from the specimens during the melting and the heating. The Eu\(^{3+}\) ions were excited by emission of Eu\(^{2+}\) ions, particularly, emission at 394 nm, and emitted the light of around 590 nm and 610 nm. That is, the energy transfer from Eu\(^{2+}\) ions to Eu\(^{3+}\) ions occurred.

The concentration quenching did not occur even for the parent glass having such high Eu\(_2\)O\(_3\) concentration as 10 mol\%. On the other hand, as the additive amount of Eu\(_2\)O\(_3\) increased, the white opaque occurred by heating at lower temperatures, which decreased the emission intensities of the glass-ceramics.
References


**Figure legends**

**Fig. 1.** Photographs of parent glasses and the glasses heated at 600 °C, 650 °C, 700 °C and 750 °C for 1h. (a) E005, (b) E05, (c) E5, (d) E10 and (e) E20 specimens.

**Fig. 2.** SEM images of polished and chemically etched surfaces of (a) E05 and (b) E10 parent glasses.

**Fig. 3.** XRD patterns of (a) E5 parent glass and E5 specimen heated at (b) 600 °C, (c) 650 °C, (d) 700 °C, (e) 750 °C and (f) 800 °C for 1h.

**Fig. 4.** XRD patterns of (a) E0, (b) E005, (c) E05, (d) E5, (e) E10 and (f) E20 specimens heated at 700 °C for 1h.

**Fig. 5.** TEM images of E5 specimen heated at 700 °C for 1h; (a) low magnification and (b) high magnification. EDX spectra of (c) dark site (M) and (d) light site (G) in the Fig. 5(a).

**Fig. 6.** Relationship between basal spacing (c·sinβ) and lattice constant (b) of mica crystals separated in (●) E0, (■) E005, (▴) E05 and (●) E5 specimens heated at 700 °C for 1h.

- Trasillic type mica: (ѵ⁴, ⚯⁵) K-phlogopite (KMg₃(AlSi₃O₁₀)F₂) and (ѵ⁹) Ca-phlogopite (Ca₀₅Mg₃(AlSi₃O₁₀)F₂).
- Tetrasillic type mica: (ѵ⁶, ⚱⁷) K-taeniolite (KMg₂Li(Si₄O₁₀)F₂), (ѵ⁸) Na-taeniolite (NaMg₂Li(Si₄O₁₀)F₂), and (ѵ⁸) K-tetrasillic mica (KMg₂₅(Si₄O₁₀)F₂).
- Brittle mica: (ѵ₁⁰) Kinoshitalite (BaMg₃(Al₂Si₂O₁₀)(OH)₂).
Fig. 7. Emission spectra of (a) E5 parent glass and E5 specimen heated at (b) 600 °C, (c) 650 °C, (d) 700 °C and (e) 750 °C for 1h; excitation spectrum, $\lambda_{ex} = 254\text{nm}$.

Fig. 8. Excitation (solid line) and emission spectra (dashed line) of E5 parent glass; emission spectrum, $\lambda_{em} = 405\text{ nm}$; excitation spectrum, $\lambda_{ex} = 289\text{ nm}$.

Fig. 9. (a) Excitation and (b) emission spectra of E5 parent glass; emission spectrum, $\lambda_{em} = 610\text{ nm}$; excitation spectrum, $\lambda_{ex} = 349\text{ nm}$.

Fig. 10. Changes in emission intensities at (a) 405 nm and (b) 610 nm of (○) parent glasses and (●) specimens heated at 700 °C for 1 h with an increase in Eu$_2$O$_3$ content; excitation spectrum, $\lambda_{ex} = 254\text{ nm}$.
Fig. 1. Photographs of parent glasses and the glasses heated at 600 °C, 650 °C, 700 °C and 750 °C for 1h. (a) E005, (b) E05, (c) E5, (d) E10 and (e) E20 specimens.
**Fig. 2.** SEM images of polished and chemically etched surfaces of (a) E05 and (b) E10 parent glasses.
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Fig. 4. XRD patterns of (a) E0, (b) E005, (c) E05, (d) E5, (e) E10 and (f) E20 specimens heated at 700 °C for 1h.
Fig. 5. TEM images of E5 specimen heated at 700 °C for 1h; (a) low magnification and (b) high magnification. EDX spectra of (c) dark site (M) and (d) light site (G) in the Fig. 5(a).
Trisilicic type mica: (▾[4], ◎[5]) K-phlogopite (K\textsubscript{Mg}\textsubscript{3}(Al\textsubscript{3}Si\textsubscript{3}O\textsubscript{10})F\textsubscript{2}) and (◎[9]) Ca-phlogopite (Ca\textsubscript{0.5}Mg\textsubscript{3}(Al\textsubscript{3}Si\textsubscript{3}O\textsubscript{10})F\textsubscript{2}).

Tetrasilicic type mica: (○[6], ◻[7]) K-taeniolite (K\textsubscript{Mg}\textsubscript{2}Li(Si\textsubscript{4}O\textsubscript{10})F\textsubscript{2}), (◎[6]) Na-taeniolite (Na\textsubscript{Mg}\textsubscript{2}Li(Si\textsubscript{4}O\textsubscript{10})F\textsubscript{2}), and (◎[8]) K-tetrasilic mica (K\textsubscript{Mg}\textsubscript{2.5}(Si\textsubscript{4}O\textsubscript{10})F\textsubscript{2}).

Brittle mica: (▵[10]) Kinoshitalite (Ba\textsubscript{Mg}\textsubscript{3}(Al\textsubscript{2}Si\textsubscript{2}O\textsubscript{10})(OH)\textsubscript{2}).
**Fig. 7.** Emission spectra of (a) E5 parent glass and E5 specimen heated at (b) 600 °C, (c) 650 °C, (d) 700 °C and (e) 750 °C for 1h; excitation spectrum, $\lambda_{ex} = 254$nm.
Fig. 8. Excitation (solid line) and emission spectra (dashed line) of E5 parent glass; emission spectrum, $\lambda_{em} = 405$ nm; excitation spectrum, $\lambda_{ex} = 289$ nm.
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Fig. 10. Changes in emission intensities at (a) 405 nm and (b) 610 nm of (○) parent glasses and (●) specimens heated at 700 °C for 1 h with an increase in Eu2O3 content; excitation spectrum, λex = 254 nm.