Optical Characterization of Nano- and Microcrystals of EuPO₄ Created by One-Step Synthesis of Antimony-Germanate-Silicate Glass Modified by P₂O₅

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Abstract: Technology of active glass-ceramics (GC) is an important part of luminescent materials engineering. The classic method to obtain GC is based on annealing of parent glass in proper temperature and different time periods. Generally, only the bulk materials are investigated as a starting host for further applications. However, the effect of an additional heat-treatment process on emission and structural properties during GC processing is omitted. Here, we focus on the possibility of obtaining transparent glass-ceramic doped with europium ions directly with a melt-quenching method. The influence of phosphate concentration (up to 10 mol %) on the inversion symmetry of local environment of Eu³⁺ ions in antimony-germanate-silicate (SGS) glass has been investigated. The Stark splitting of luminescence spectra and the local asymmetry ratio estimated by relation of \((5D_0 \rightarrow 7F_2)/(5D_0 \rightarrow 7F_1)\) transitions in fabricated glass confirms higher local symmetry around Eu³⁺ ions. Based on XRD and SEM/EDX measurements, the EuPO₄ nano- and microcrystals with monoclinic geometry were determined. Therefore, in our experiment, we confirmed possibility of one-step approach to fabricate crystalline structures (glass-ceramic) in Eu-doped SGS glass without additional annealing process.

Keywords: glass-ceramics; europium oxide; luminescence properties; Stark splitting; EuPO₄ nanocrystals; antimony-germanate-silicate glass

1. Introduction

In photonic material technology, transparent glass-ceramic (GC) with active nanocrystals is still required [1–4]. Specific optical properties of active GC materials resulting from two-phase structure (amorphous and crystalline) are found in the vicinity of lanthanide ions doped glassy host. The main benefits of a crystalline environment surrounding the rare earth ions are the high absorption and emission cross sections, lower phonon energy, and optimization of the ion–ion interaction [5]. The europium (Eu³⁺) ion is one of the interesting to analyze luminescence properties because: (i) it is characterized by relatively high quantum efficiency since the energy gap between the \(^5D_0\) emitting level and the underlying \(^7F_j\) multiplet is approx. 12,000 cm⁻¹, which means that the non-radiative decay is less likely to occur even in hosts with high phonon energies such as phosphate and silicate glasses.
(1100–1200 cm$^{-1}$); (ii) the intensity ratio between electrical-dipole (ED: $^5D_0 \rightarrow ^7F_2$) to magnetic-dipole (MD: $^5D_0 \rightarrow ^7F_1$) transitions can be used as a probe for site symmetry which is helpful to control ceramization process; and (iii) the Eu$^{3+}$ ions-doped optical material can be used as an efficient phosphor for solid state light sources [6–10]. It is also well known that the intensities of emission bands of europium ion in glasses depend on its concentration and glass composition [11–14]. In order to obtain optimum emission characteristics for glass-ceramic applications, the influence of glass composition as well as concentration dependence studies of Eu$^{3+}$ ions are essential. To date, a lot of photonic glasses after ceramization are presented in original papers [5,15–20]. Our idea is to modify thermally stable antimony-germanate glass with phosphorous oxide. Based on our earlier investigations on luminescent properties of antimony-germanate glasses doped with lanthanide ions we confirmed good thermal stability of glassy matrix. We also observed that the combination of two various glass-forming elements is particularly important in reduction of unfavorable non-radiative transitions between energy levels of rare-earth ions [21–23]. Thus, in case of glass-ceramic fabrication we decided to use P$_2$O$_5$ as a precursor of crystallization in antimony-germanate-silicate glasses doped with Eu$_2$O$_3$.

In the view of above, we demonstrated one-step method to obtain glass-ceramic material through the modification of antimony-germanate-silicate glass with a small amount of P$_2$O$_5$. The local symmetry in vicinity of Eu$^{3+}$ ions and EuPO$_4$ crystals creation were characterized by UV–vis spectrometry, X-ray diffraction (XRD), Scanning Electron Microscopy (SEM), and Energy Dispersive X-ray Spectrometry (EDS). It has been noticed, that the glass with 5 mol % of P$_2$O$_5$ shows band partition to emission sub-wavelengths of Eu$^{3+}$ ions, which is characteristic effect in glass-ceramic materials. Also, the partial crystallization has been observed by naked eye directly after the melting process.

2. Results and Discussion

2.1. Analysis of Excitation and Emission Spectra

Figure 1 shows excitation spectra of europium ions in the fabricated glasses, which have been monitored at the wavelength of 616 nm ($^5D_0 \rightarrow ^7F_2$ transition). All glass samples are labeled due to phosphate oxide concentration as follows: SGS05P, SGS1P, SGS3P, SGS5P, SGS7P, and SGS10P. In analyzed spectral range (350–500 nm) five bands centered at the wavelengths 362 nm ($^7F_0 \rightarrow ^5D_4$), 382 nm ($^7F_0 \rightarrow ^5L_7$), 395 nm ($^7F_0 \rightarrow ^5L_6$), 416 nm ($^7F_0 \rightarrow ^5D_3$) and 464 nm ($^7F_0 \rightarrow ^5D_2$) were observed. The band at 395 nm is characterized by most intense transition in UV–vis spectral range, hence it is suitable for the effective excitation of the Eu$^{3+}$ ions by laser radiation at 394 nm. It worth to notice that intensity of second important excitation band at 464 nm in glass sample doped with 0.5 mol % P$_2$O$_5$ is more prominent. This band is called as the “hypersensitive transition” and strongly depends on structural changes of glassy matrix [24]. In our experiment, the increase of phosphate content in antimony-germanate glass leads to the strong reduction of “hypersensitive transition” intensity. Thus, the intensity change of $^7F_0 \rightarrow ^5D_2$ transition observed in fabricated glasses confirms the structural distortion of the glass network in the vicinity of Eu$^{3+}$ ions.

![Figure 1](image-url) Figure 1. The excitation spectra of SGS05P, SGS1P, SGS3P, SGS5P, SGS7P, and SGS10P glass samples monitored at 616 nm.
The luminescence spectra of fabricated antimony-germanate-silicate glasses doped with Eu\(^{3+}\) ions with different content of P\(_2\)O\(_5\) under excitation by laser radiation with \(\lambda_{\text{exc}} = 394\) nm were shown in Figure 2. In the range of 525–725 nm five characteristic emission bands have been observed at the wavelengths of 580, 594, 612, 653, and 703 nm originating from transitions of \(^5D_0 \rightarrow ^7F_0, ^7F_1, ^7F_2, ^7F_3\) and \(^7F_4\), respectively. All characteristics were scaled (integrated) to \(^5D_0 \rightarrow ^7F_1\) transition due to their having independent features.

![Figure 2. The luminescence spectra of SGSxP glasses doped with 0.5 mol % Eu\(_2\)O\(_3\).](image)

The changes in luminescence shape at 594 nm \(^5D_0 \rightarrow ^7F_1\) and 616 nm \(^5D_0 \rightarrow ^7F_2\) suggest that incorporation of P\(_2\)O\(_5\) into glass matrix leads to the local structure modification of fabricated glasses. It worth to notice, that especially luminescence spectrum for SGS5P glass shows stark splitting for \(^5D_0 \rightarrow ^7F_1\) and \(^5D_0 \rightarrow ^7F_2\) transitions. Based on the Eu\(^{3+}\) ions luminescent properties and group-theoretical arguments it is possible to analyzed the structure of local environment in the vicinity of the ion [25,26]. In case of \(^5D_0 \rightarrow ^7F_1\) transition three emission peaks resulting from the total removal of crystal field degeneracies were observed. The intensity of the \(^5D_0 \rightarrow ^7F_2\) transition is higher than intensity of \(^5D_0 \rightarrow ^7F_1\) transition in all samples except SGS5P, where dominant is transition \(^5D_0 \rightarrow ^7F_1\). This result confirms that the local symmetry of Eu\(^{3+}\) ion is without an inversion center for SGS05P, SGS1P, SGS3P, SGS7P, and SGS10P samples and with an inversion center for SGS5P sample [27]. Additionally, the \(^5D_0 \rightarrow ^7F_2\) transition is a “hypsersensitive transition” which means that it strongly depends on the local symmetry and environmental effects in the surroundings of the Eu\(^{3+}\) ions [28]. Among the \(^5D_0 \rightarrow ^7F_1\) transitions, the emission line at 594 nm \((^5D_0 \rightarrow ^7F_1)\) is a magnetic dipole transition (MD), while the emission line at 616 nm \((^5D_0 \rightarrow ^7F_2)\) is an electric dipole transition (ED). The ratio between integrated intensity of \((^5D_0 \rightarrow ^7F_2)/(^5D_0 \rightarrow ^7F_1)\) transitions, gives a factor of the distortion degree from the inversion symmetry of local environment of the europium ions. This value is less than 1.0 for symmetric and is greater than 1.0 for non-centrosymmetric surroundings [14]. In our experiment, the ED/MD transition ratio decreases rapidly from 4.24 to 0.92 in function of P\(_2\)O\(_5\) (up to 5 mol %), then starts to increase to approximately 3 in glasses with 7 mol % and 10 mol % of P\(_2\)O\(_5\) (Table 1). The relatively lower values of the ED/MD ratio for SGS5P suggest incorporation of Eu\(^{3+}\) ions into more symmetric environment with less covalent character [27]. Therefore, the ED/MD value is related to the enhancement in the symmetry of the ligand field around Eu\(^{3+}\) ions [29]. The variation in the asymmetric ratio and stark splitting for \(^5D_0 \rightarrow ^7F_1\), \(^5D_0 \rightarrow ^7F_2\) and \(^5D_0 \rightarrow ^7F_4\) transitions in SGS5P sample may be due to the presence of the crystalline features [30]. Figure 3 presents the decay curves for all SGS glass samples excited at 394 nm, and the lifetime values obtained by fitting the curves with exponential functions are listed in Table 1. The luminescence decay curves of \(^5D_0\) energy level of
Eu$^{3+}$ in the glasses from SGS05P to SGS7P were fitted to the double-exponential fitting functions with a short decay ($\tau_1$) and a long decay ($\tau_2$). However, the luminescence decay curves of $^5D_0$ state of Eu$^{3+}$ in the SGS10P glass was best fitted to a single-exponential fitting function, which indicated a 1.99 ms lifetime. The luminescence intensity $I(t)$ of the Eu$^{3+}$ in glasses could be described by the sum of two exponential decay components from

$$I(t) = A_1 \exp\left(-\frac{t}{\tau_1}\right) + A_2 \exp\left(-\frac{t}{\tau_2}\right)$$  \hspace{1cm} (1)

where $\tau_1$ and $\tau_2$ were short- and long-decay components, respectively. Parameters $A_1$ and $A_2$ were fitting constants. According to Equation (1), the average lifetime $<\tau_{avg}>$ was given by

$$<\tau_{avg}> = \frac{A_1 \tau_1^2 + A_2 \tau_2^2}{A_1 \tau_1 + A_2 \tau_2}$$  \hspace{1cm} (2)

where weight factors $A_1$ and $A_2$ were introduced. According to Equation (2), the average lifetimes of $^5D_0$ energy level of Eu$^{3+}$ in the fabricated antimony-germanate glasses were calculated and presented in Table 1. It is well known that the luminescence of rare earth ions depended more or less efficiently on the molecular structure of the host. The luminescence decay rate was the sum of the radiative decay and multiphonon relaxation rates [31]. The non-radiative relaxation between various $J$ states might occur by interaction of the electronic levels of rare earth ions with suitable vibrational modes of the environment [32]. Therefore, the double-exponential decay curves indicated that there were two different surroundings of the europium ions in the fabricated glass: some in the glass and the others in the nanocrystals. However, in the case of glasses with highest concentration of P$_2$O$_5$ (10 mol %) the luminescence decay has single-exponential character, which is a rather unexpected result and needs further structural investigations.

![Figure 3](image-url)  
**Figure 3.** The luminescence decay of $^5D_0$ energy level of europium ions in fabricated SGSxP glasses.

**Table 1.** The ED/MD transition ratio, lifetimes ($\tau_1$ and $\tau_2$) and average lifetime $<\tau_{avg}>$ of $^5D_0$ level of Eu$^{3+}$ ions.

| Glass Sample | ED/MD Ratio | $<\tau_{avg}>$ [ms] |
|--------------|-------------|---------------------|
| SGS05P       | 4.24        | 1.29                |
| SGS1P        | 2.02        | 1.29                |
| SGS3P        | 1.27        | 1.86                |
| SGS5P        | 0.92        | 0.83                |
| SGS7P        | 2.66        | 1.72                |
| SGS10P       | 3.03        | 1.99                |
As the \( P_2O_5 \) concentration increases from 0.5 to 7 mol \%, there is a slight increase of average \( ^5D_0 \) lifetime from 1.29 to 1.86 ms. In fact, the variation of lifetime is not monotonic and the shortest average lifetime of \( ^5D_0 \) level was obtained with the SGS5P glass sample. This effect could be related to the division of the excitation energy between Stark’s sub-levels. It is also well known in literature that the double-exponential decay indicates the energy transfer between Eu\(^{3+} \) ions by cross-relaxation process. Mainly in glasses with high concentration of Eu\(^{3+} \) ions. However, an interesting effect is that the SGS10P glass is characterized by longest lifetime (1.99 ms) than others glass samples. Based on these results, we confirmed that in fabricated glasses double-exponential decay only depends on concentration of \( P_2O_5 \).

2.2. X-ray Diffraction (XRD), Scanning Electron Microscopy (SEM), and Energy Dispersive X-ray Spectrometry (EDS)

In order to confirm the presence of a crystalline phase or phases, the XRD measurements for SGS5P (lowest asymmetry ratio) sample was performed in the range from 15 to 60 degrees of diffraction angle (Figure 4). Broad halo effect between 2\( \theta \) = 20°–35° suggests the structural disorder and confirms dominance the amorphous nature of the SGS glass. However, in the diffraction pattern, three weak reflections peaks in the range of 2\( \theta \) from 28 to 31 degrees were also observed. After the reference analysis of peak positions calculated reflections are in good accordance with monoclinic EuPO\(_4\) crystalline phase (ICSD: 01-083-0656). From the peak width of the XRD pattern, the size of EuPO\(_4\) nanocrystals was estimated by the Scherrer formula

\[
D = \frac{\lambda K}{\beta \cos \theta}
\]

where \( D \) is the crystal size at the vertical direction, \( \lambda \) is the wavelength of the X-ray, \( \theta \) is the angle of diffraction, \( \beta \) is the full width at half maximum (FWHM) of the diffraction peak and the instrument constant \( K = 0.9 \) [33]. The estimated average size of nanocrystals is about 30 nm. Figure 5 shows scanning electron microscope (SEM) images of SGS3P and SGS5P samples. As can be seen, in both SEM images there are visible crystals of a size range approx. 90–120 nm for the sample labeled as SGS3P (Figure 5a) and a few microns for the SGS5P sample (Figure 5b).

![XRD patterns of SGS5P glass sample.](image)

In the case of the SGS5P sample, the analysis of the chemical composition in two different areas (EDS for amorphous—1; and for crystalline—2) shows that the microparticles observed in the SEM image originate from crystals of europium phosphate (supplementary file—Figure S1).
In the case of SGS3 sample, due to the nanometric size of the particles is not possible to obtain an EDS spectrum separately for crystalline and the amorphous area. However, due to very similar chemical composition of the analyzed glass samples, it can be assumed that EuPO$_4$ crystals have been also observed.

3. Materials and Methods

3.1. Glass Preparation

Antimony-germanate-silicate (SGS) glasses have been investigated in our earlier works due to good thermal stability, which is required in the optical fiber fabrication process [22,23]. Here, we proposed the modification of SGS glass by small amount of P$_2$O$_5$ from 0.5 mol % to 10 mol %. The glass with molar composition 25Sb$_2$O$_3$–25Ge$_2$O$_3$–10Al$_2$O$_3$–5Na$_2$O–(35-x)SiO$_2$–xP$_2$O$_5$–0.5Eu$_2$O$_3$ was synthesized by standard melt-quenching method. The Eu$_2$O$_3$ concentration was fixed on the level of 0.5 mol % and glasses are labeled as: SGS05P, SGS1P, SGS3P, SGS5P, SGS7P, and SGS10P. All the raw materials were analytical grade reagents (99.99%). Europium ions were used as a local environment probe, to obtain valuable information on the site symmetry of active ions in a fabricated host directly by luminescent measurements. A homogenized set was placed in a platinum crucible and melted in an electric furnace at 1450 °C for 60 min in oxygen atmosphere. Next, the glass melt was poured into a brass plate at the room temperature (RT) and then annealed at 400 °C for 12 h to release the internal stress from the quench. Next, glasses were cooled down to room temperature and polished carefully in order to meet the requirements for optical measurements. After this, in a few glass samples, the partial-crystallization (opaque) phenomenon and agglomeration of crystals were observed by naked eye.

3.2. Structural and Spectroscopic Analysis

The XRD patterns of fabricated glasses were measured in the range from 10° to 80° using an XPert Pro diffractometer (PANalytical, Eindhoven, Netherlands). The Cu X-ray tube with K$_\alpha$ radiation was used. The morphology of prepared samples was examined by FEI Company (Hillsboro, OR, USA) Nova Nano SEM 200 scanning electron microscope with an attachment for chemical analysis with energy dispersive X-ray spectroscopy (EDX, EDAX). The analyses were carried out in secondary electron mode (SE). Prior to analyses, the samples were covered with carbon layer. The excitation and luminescence spectra of the glasses in a range of 350–750 nm were measured using a JobinYvon Fluoromax4 spectrophotometer (Horiba, Kyoto, Japan). A system PTI QuantaMaster QM40 coupled with tunable pulsed optical parametric oscillator (OPO), pumped by a third harmonic of a Nd:YAG laser (OpotekOpolette 355 LD, Carlsbad, CA, USA) was used for luminescence decay measurements.
The laser system was equipped with a double 200 mm monochromator, a multimode UV–vis PMT (R928) (Hamamatsu, Japan) and H10330B-75 detectors (Hamamatsu, Japan) controlled by a computer. Luminescence decay curves were recorded and stored by a PTI ASOC-10 (USB-2500) oscilloscope with an accuracy of ±1 µs.

4. Conclusions

As a result of our experiment we synthesized the antimony-germanate-silicate glass modified by different content of P2O5. Incorporation of phosphate oxide into host enables to create monoclinic crystal phase EuPO4 directly in melt-quenching process (without additional heat-treatment). Also, the effect of partially crystallization leads to prominent Stark splitting of luminescence bands at 594 nm and 616 nm, which suggests that Eu3+ ions are surrounded by crystalline phase. According to SEM/EDS measurements, the formation of europium phosphate crystals with nano- and micrometric size have been also observed.

Supplementary Materials: The following are available online at http://www.mdpi.com/1996-1944/10/9/1059/s1, Figure S1: EDS spectra of SGS5P glass for amorphous (1) and crystalline (2) areas.

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Conflicts of Interest: The authors declare no conflict of interest.

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