

Low temperature afterglow from SrAl₂O₄: Eu, Dy, B containing glass

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Abstract

SrAl₂O₄: Eu, Dy, B particles were added in a phosphate glass (90NaPO₃-10NaF (in mol%)) using the direct doping method. For the first time, the composition of the particles prior to and after embedding them in the glass was analysed using EPMA analysis. Boron was found to be incorporated in already distorted surroundings creating new trapping centers in the particles which are thought to be favourable for the tunnelling process and so for the afterglow at 10K. Despite the partial decomposition of the particles, the glass exhibit afterglow at low temperature confirming to be promising materials for low temperature applications.

Keywords: persistent luminescence; phosphate glass; low temperature applications

Persistent luminescence is light emission from excited material that can last for several minutes or even hours after the termination of the excitation [1]. Luminescent materials use energy from some external source, such as sunlight for example and store it by the means of localizing charge carriers in some trapping centers. The charge carriers are released gradually from the trapping centers, followed by recombination process and emission of light. SrAl₂O₄:Eu,Dy is one of the most efficient persistent luminophores known up-to-date [2,3]. The afterglow can be observed for up to 20 hours after removing the excitation [1-3]. Due to its strong and extensive afterglow, SrAl₂O₄: Eu, Dy luminophore is widely used in many applications – emergency signs, luminous paints, luminescent coatings, in vivo imaging [2-6].

However, luminophores from Eu doped aluminate group lose 90% of their afterglow efficiency in temperatures below zero Celsius, which is a major problem for material outdoor use [7]. Recently it has been observed, that the addition of boron between 5 and 7 at% in SrAl₂O₄:Eu,Dy during the

1 synthesis led to particles with afterglow even in temperatures as low as 10K, however the reasoning
2 for this is unclear [8]. As for the other drawback, the PL (persistent luminescent) luminophores are
3 usually obtained during synthesis in the form of non-transparent powders thus limiting the excitation
4 and emission to the surface of the material and therefore wasting the material in the volume.
5 Therefore, a possible solution is to disperse these luminophores in a transparent matrix to increase the
6 intensity of the persistent luminescence emission by exciting the entire volume of the host optimizing
7 the usage of luminescent microparticles.
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12 Glasses based materials have been successfully prepared with $\text{Eu}^{2+}, \text{Dy}^{3+}$ -doped SrAl_2O_4 using the
13 “Frozen sorbet method” [9] and also using the direct doping method [10]. In the “Frozen sorbet
14 method”, the luminophore precipitates using the elements from the glass and therefore, the
15 composition of the luminophore can not be controlled. On the other hand, persistent luminescent
16 glasses can be prepared independently of their composition using the direct doping method as the
17 luminophore with target composition and spectroscopic properties are added in the glass melt.
18 However, as explained in [10], the direct doping method needs to be optimized as decomposition of
19 the particles occurs during the glass preparation. The decomposition of the particles depends on the
20 temperature at which the particles are added in the melt and also on the duration the particles are in
21 contact with the glass melt. Using optimized doping parameters, phosphate glasses with the
22 composition 90 NaPO_3 -10 NaF (in mol%) were successfully prepared with persistent luminescence by
23 adding various commercial persistent luminescent particles in the glass melt [11, 12]. This glass was
24 chosen due to its low melting temperature and high capacity for dopant particles. The addition of the
25 promising boron containing $\text{SrAl}_2\text{O}_4:\text{Eu}, \text{Dy}$ particles in this glass would allow the fabrication of a
26 promising composite for low-temperature sensing applications.
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40 In this paper, we present, for the first time, a glass-based composite that exhibits afterglow at very
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42 as well as the change in the distribution of the trapping center when embedding the luminophores in
43 the glass.
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50 In the present study, Eu, Dy, and B doped SrAl_2O_4 samples were synthesized using the sol-gel
51 method described in [8]. 1 at% Eu, 2 at% Dy SrAl_2O_4 samples were prepared with 5 and 7 at% of
52 boron (labelled as 5% B and 7% B, respectively). The $\text{SrAl}_{2-x}\text{B}_x\text{O}_4:\text{Eu}, \text{Dy}$ ($x=0.05$ and 0.07)
53 particles were then added in the glass with the composition 90 NaPO_3 -10 NaF (in mol%) using the
54 direct doping method as in [10]. The glass was prepared using $\text{Na}_6\text{O}_{18}\text{P}_6$ (Alfa-Aesar, technical
55 grade), Na_2CO_3 (Sigma-Aldrich, >99.5%) and NaF (Sigma-Aldrich, 99.99%). After melting the
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1 glasses at 750°C, the temperature of the glass melt was reduced to 575°C before adding 0.25 weight-
2 % of the PL particles. 3 min after adding the particles, the glasses were quenched and finally
3 annealed at 40 °C below their respective glass transition temperature for 6h in air. The glasses
4 prepared with the particles doped with x=0.05 and 0.07 of boron are labelled 5% glass and 7% glass,
5 respectively.
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9 An Electron Probe MicroAnalyzer (EPMA) (CAMECA, SX100) was used to determine the
10 composition of the glasses and of the particles with an accuracy of ± 0.1 at % using the Cameca
11 QUANTITOOl analytical programme. The EPMA was operated at 15 keV and 40 nA. Prior to the
12 measurement, the samples were polished and coated with a carbon layer to prevent charging.
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16 The persistent luminescence spectra were recorded using Andor Shamrock B303-I spectrometer.
17 The samples were excited by pulsed laser excitation. Excitation source was YAG:Nd laser LCS-
18 DTL-382QT (266 nm, 8 ns). The decay profiles were also recorded using the same equipment. For
19 thermally stimulated luminescence (TSL) measurements, the samples were cooled with Sumitomo
20 HC-4 closed-cycle helium cryostat, temperature range 9 – 325 K. Lake Shore 331 Temperature
21 controller was used for temperature control as well as for sample heating (6 K/min) during TSL
22 measurements up to 320 K. The excitation source for TSL measurements was X-ray tube with W
23 target (30kV, 10 mA).
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33 SrAl₂O₄: Eu, Dy particles with 5% and 7% B addition were chosen in this study due to their
34 excellent afterglow at 10K as reported in [8]. The reader is referred to this paper for a complete
35 characterization of the particles. EPMA was used to analyse the composition of the particles prior to
36 adding them in the glass.
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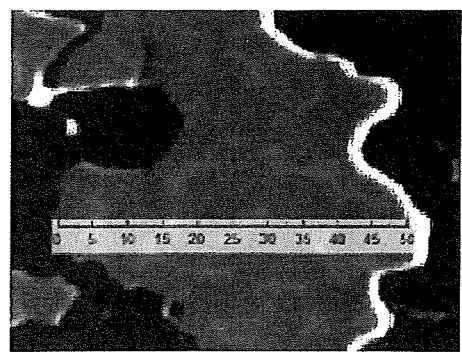
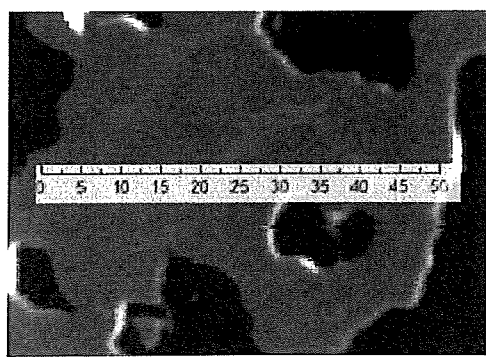
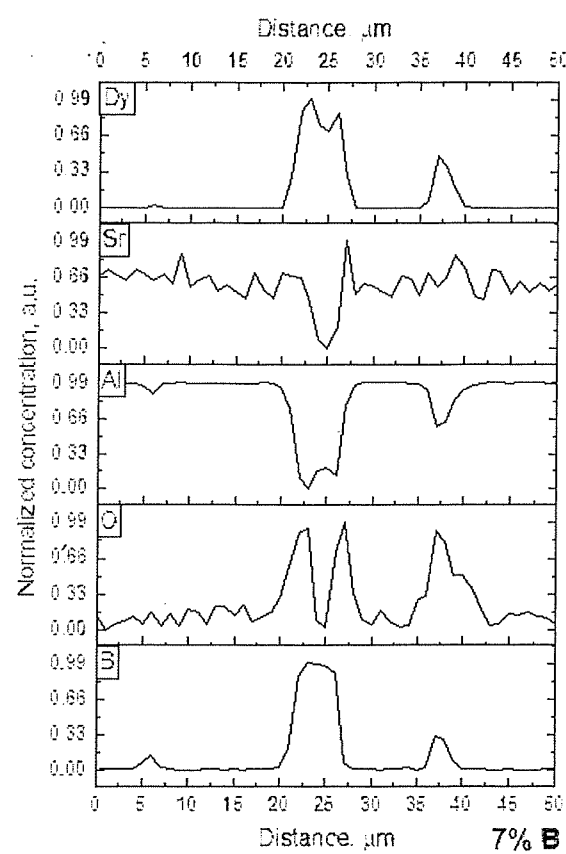
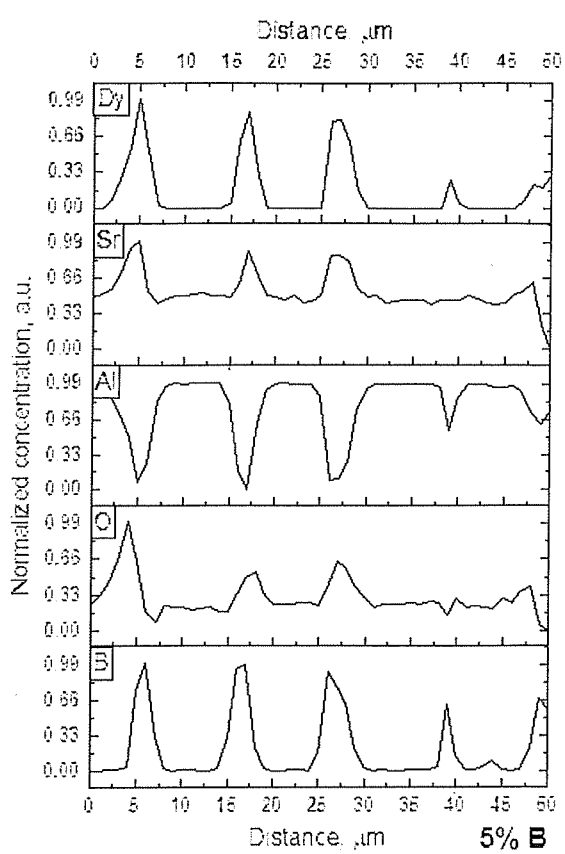


Figure 1. Composition analysis of the SrAl_2O_4 : Eu, Dy particle with 5% B (a) and 7% B (b)

1 According to the composition analysis (Fig. 1), boron seems to be unevenly distributed throughout
2 the particles. There are some doubts in the scientific community about the role of boron, when it
3 enters the lattice of SrAl₂O₄ [13, 14]. Boron is believed to create a shrinkage of lattice and a
4 distortion of the Dy³⁺ environment, but the thermally stimulated luminescence data are not consistent
5 when discussing the trapping centers in the material that are altered due to the introduction of B
6 [13,14]. The concentration of Al decreases while the concentration of B increases indicating that B is
7 incorporated in the lattice site of Al. Dy follows the same pattern, leading to the conclusion that B
8 tends to incorporate in close proximity of Dy. Recently it has been discussed that the low
9 temperature luminescence in SrAl₂O₄: Eu, Dy is due to the creation of excited Eu²⁺ via direct
10 tunnelling from some trapping centers, that can not be emptied thermally due to lack of thermal
11 energy [15]. One of the possibilities of the unknown electron trapping centers in SrAl₂O₄
12 luminophores is the Dy - distorted AlO₄ tetrahedra [3]. If B incorporates instead of Al in the crystal
13 lattice, the smaller ion size of B leads to distortion of the lattice. Previous research confirmed the
14 existence of substitutional BO₄ units in the samples when adding boron [14]. BO₄ is more ionic in
15 comparison to AlO₄ due to the higher electronegativity and smaller size of B compared to Al [14].
16 Therefore, the B addition presumably would create lattice distortions, that can lead to electron
17 trapping. Dy ions, when incorporating in the Sr lattice site, has an uncompensated charge stimulating
18 the charge trapping on BO₄ units. The EPMA analysis allow us to conclude, that B incorporates in
19 already distorted surroundings creating new trapping centers in the material which are thought to be
20 favourable for the tunnelling process.
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37 The particles were added in the glasses using the direct doping method. The glasses exhibit
38 homogeneous green PL confirming the survival of well dispersed particles in the glasses. The
39 photoluminescence spectra (Fig. 2a and b) show a broad emission band with maximum at 530 nm,
40 that can be attributed to the Eu²⁺ emission band.
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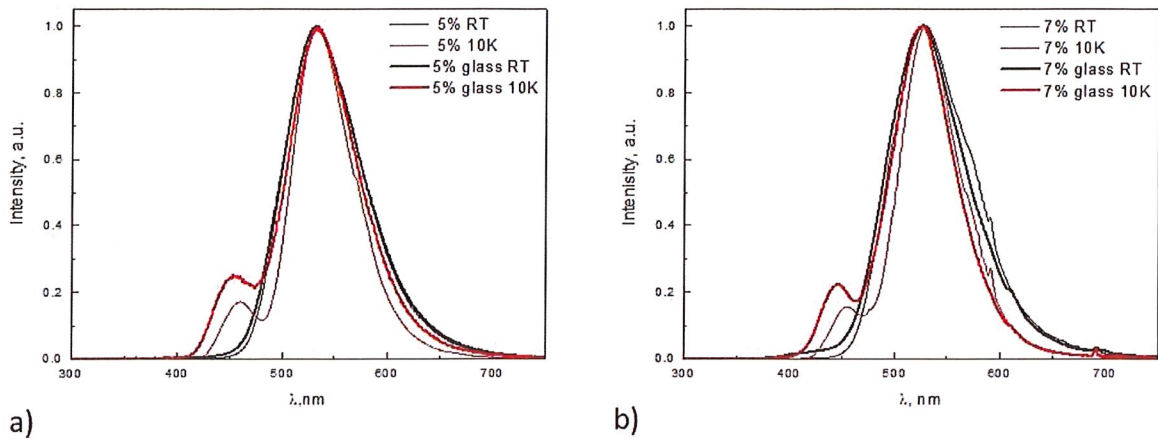
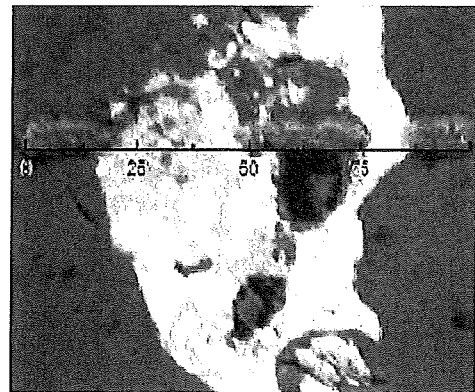
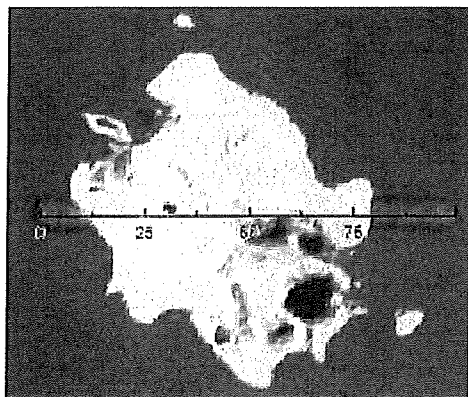
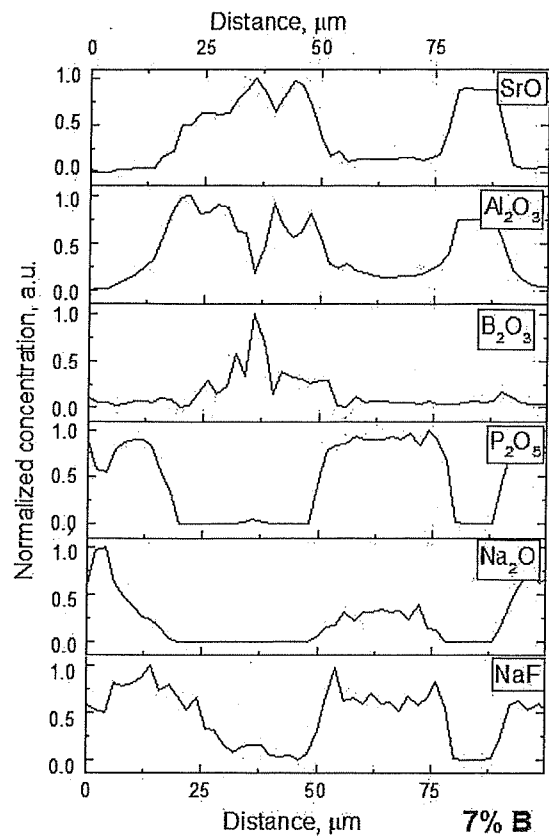
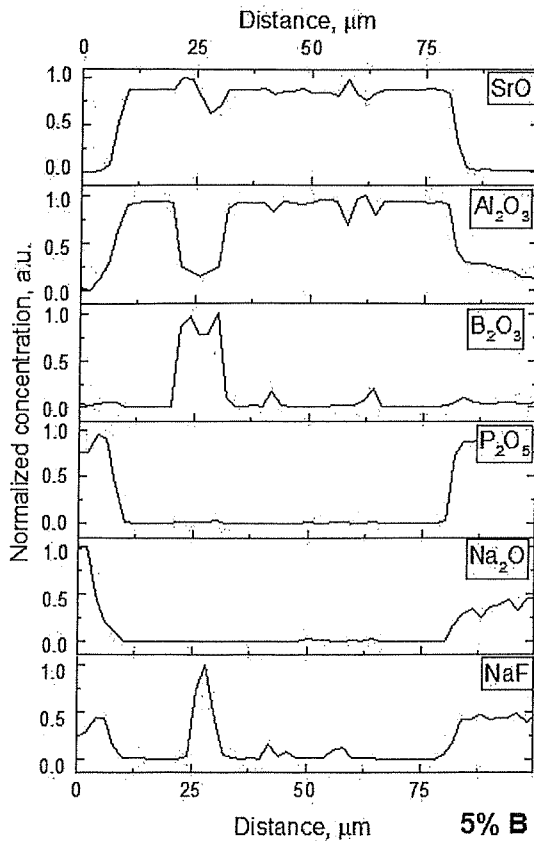


Figure 2. Photoluminescence spectra of the PL glass and of the PL particles alone with a) 5% B and b) 7% B. The spectra were measured at room temperature and at 10K.

The samples were cooled to 10K temperature and the spectra were recorded. Both particles exhibit a narrowing of the emission band due to the lack of phonon interactions and a second peak also emerges at 455 nm. This peak is attributed to Eu^{2+} emission from differently coordinated Sr sites and is thermally quenched at temperatures above 150 K as explained in [3]. We can note that the PL spectra measured at RT do not change significantly when incorporating the luminophores in glass matrix: the intensity of the band at 455 nm compared to the main band at 530 nm increases when performing the measurement at 10K indicating that the site of the RE ions in the particles is slightly changed after embedding the particles in the glass. As explained in [10], the changes in the site of the RE ions can be due to the partial decomposition of the particles occurring during the glass preparation. As depicted in Fig. 3, Al and Sr can be found in the glass at the glass-particles interface confirming the partial decomposition of the particles associated with the diffusion of the elements from the luminophore to the glass.



a)

b)

Figure 3. Composition analysis of the $\text{SrAl}_{2-x}\text{B}_x\text{O}_4$: Eu, Dy particle after embedding in glass with with 5% B (a) and 7% B (b)

The average size of the particles found at the surface of the glasses is around 100 μm indicating that the particles have agglomerated in the glass increasing their size and therefore decreasing the surface/volume ratio. The composition analysis reveals that the particles conserved their compositional integrity: they exhibit a quite homogeneous distribution of Al_2O_3 and SrO in their

center. It should be noted that there are no characteristic Eu^{3+} lines visible in the photoluminescence spectra of the glasses. Therefore we can conclude that although Eu^{2+} are also suspected to diffuse in the glass, there is no (or little) oxidation from Eu^{2+} to Eu^{3+} taken place during the melting process.

The PL decay of the particles prior to and after being embedded in the glass are shown in Fig. 4a. The microparticles alone show an afterglow of several hours and for several minutes in the low temperature region before they reach the level of 0.32 mcd. However, when incorporated in glass, the afterglow time is reduced due to the particle degradation during melting procedure (Table 1).

	5% B		7% B	
	RT	10K	RT	10K
Particles alone	>8 hours	> 50 min	>8 hours	> 50 min
Particles embedded in glass	150 min \pm 10s	50 s \pm 10s	20 min \pm 10s	300 s \pm 10s

Table 1: Afterglow time until reaches 1% of initial intensity of the investigated particles containing glass measured at room temperature (RT) and at 10K

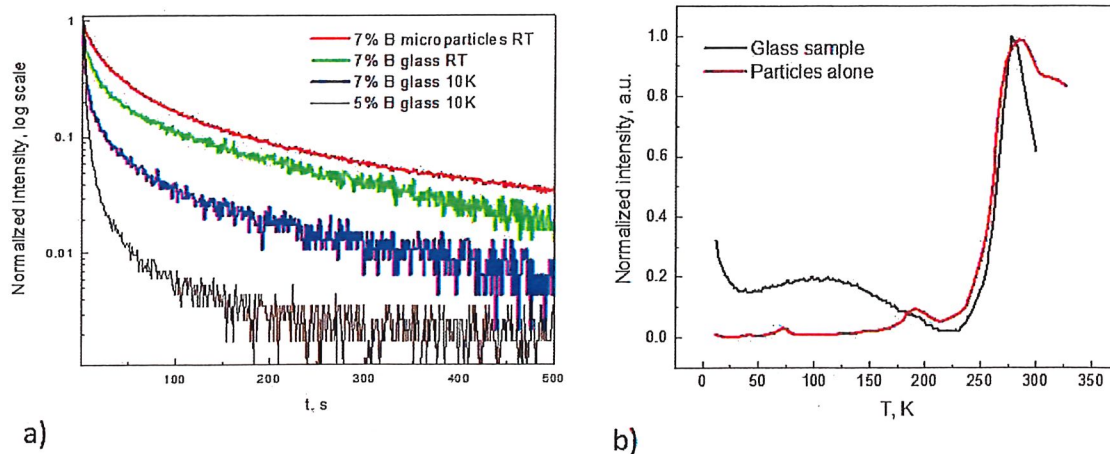


Figure 4. Photoluminescence decay kinetics comparison of the particles alone and of the glasses (a) and thermally stimulated luminescence (TSL) spectra of the particles with 7% B prior to and after being embedded in the glass (b)

The TSL analysis of the particles and particles containing glass were carried out in temperature range 10-320K (Fig. 4b). The particles alone exhibit quite distinct TSL peaks which can be related to well-defined trapping center (intrinsic SrAl_2O_4 defects as discussed in [16]) energies. However, the wide TSL peak with maximum at around 100K of the particles embedded in the glass is a clear

1 indication that the trapping center energies are spread out after embedding the particles in the glass.
2 The defect concentration in the particles increases when adding the particles in the glass and the
3 defects might migrate with respect to their position in particles due to the partial decomposition of
4 the particles during the glass melting, resulting in a different trapping center distribution.
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10 In summary, phosphate glasses containing $\text{SrAl}_2\text{O}_4:\text{Eu}^{2+}$, Dy^{3+} were successfully prepared with
11 persistent luminescence at low temperature by adding the B containing PL particles in the glass melt.
12 From the EPMA analysis of the particles alone, B tends to incorporate in already distorted crystal
13 lattice surroundings in the proximity of those Sr lattice sites, that have been replaced by Dy. The
14 addition of B presumably creates different lattice distortions, that can lead to electron trapping. The
15 afterglow decay of the glasses was considerably lower compared to that of the particles alone due to
16 the decomposition of the particles occurring during the glass preparation. However, due to B in the
17 particles, the glass-based materials exhibit afterglow at low temperature creating possible sensing
18 and biomedical applications.
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Figure 1
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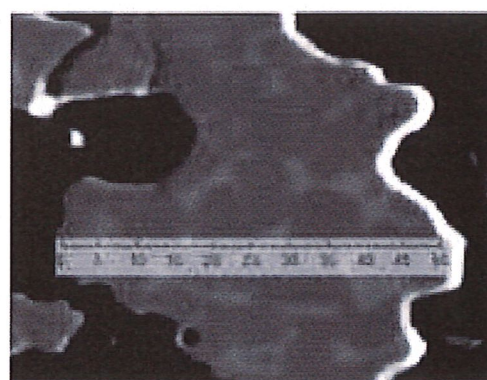
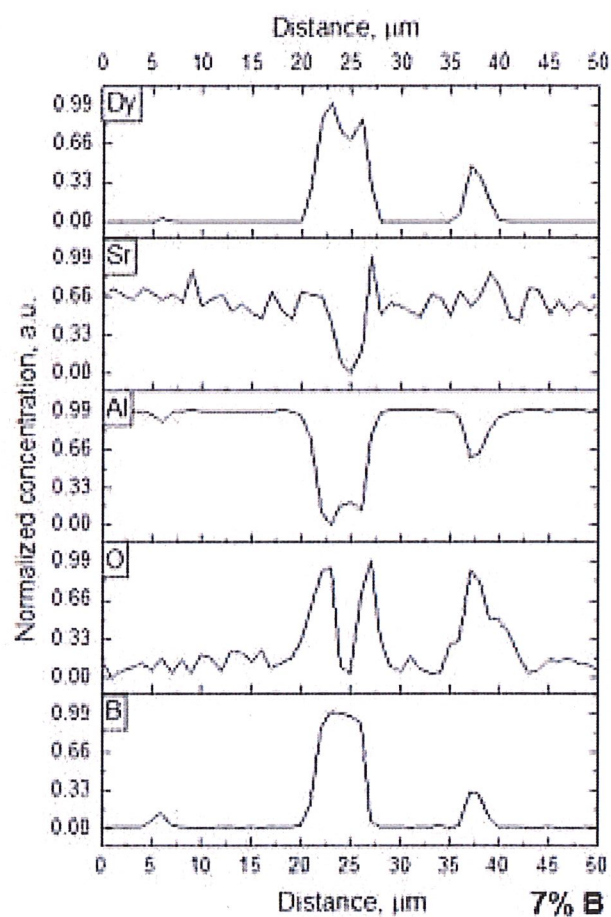
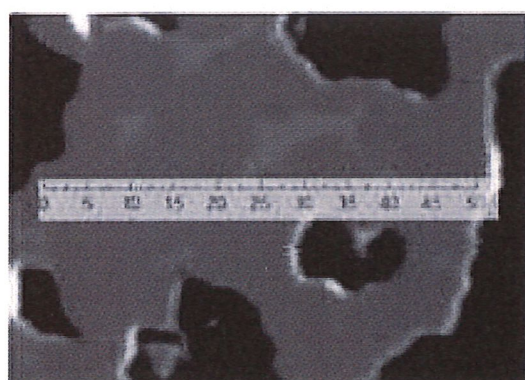
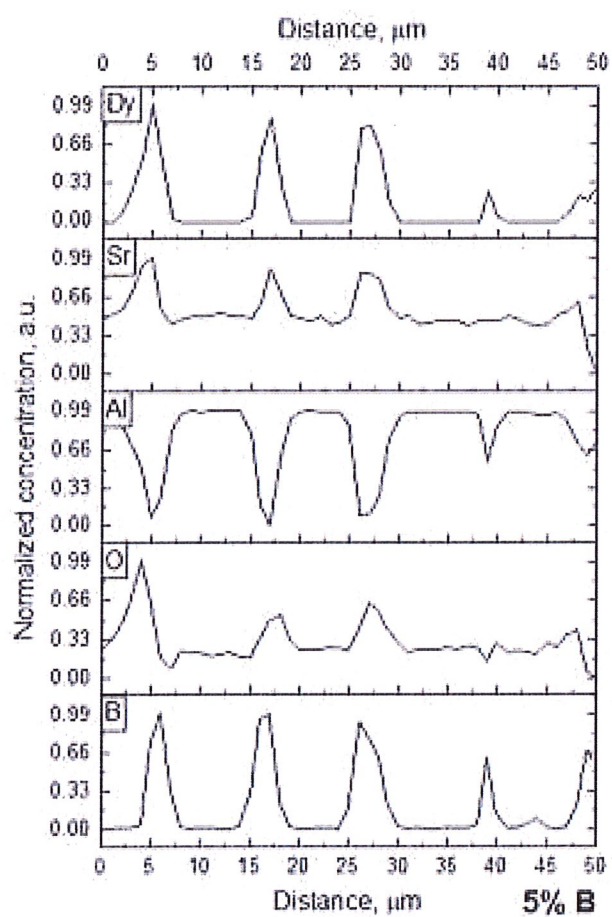


Figure 2
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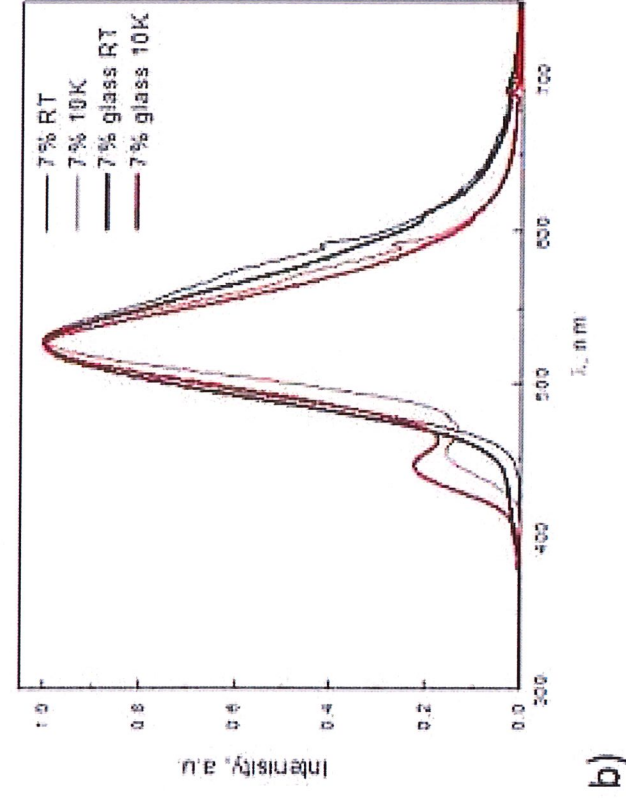
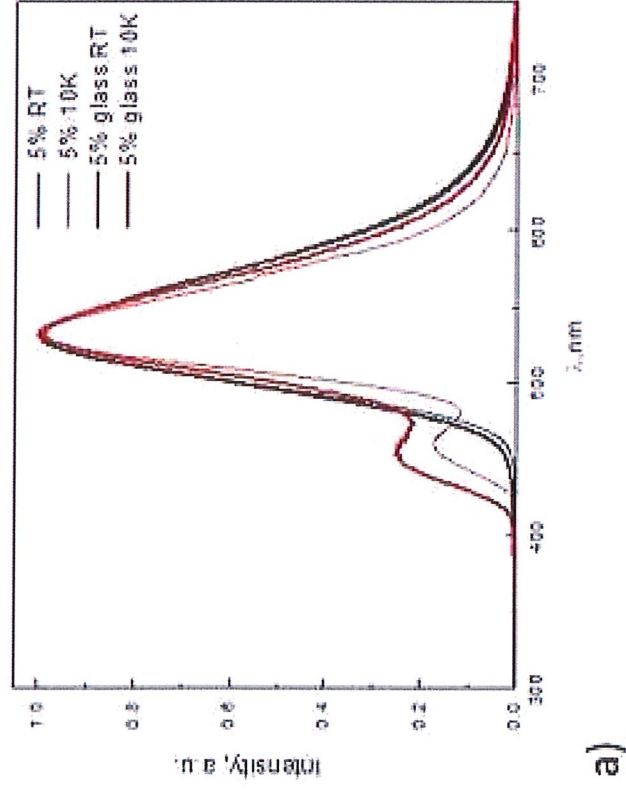


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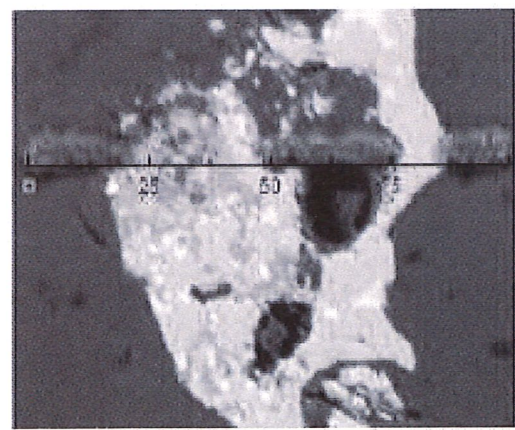
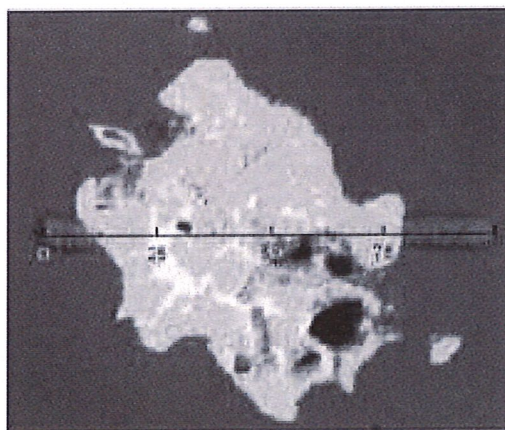
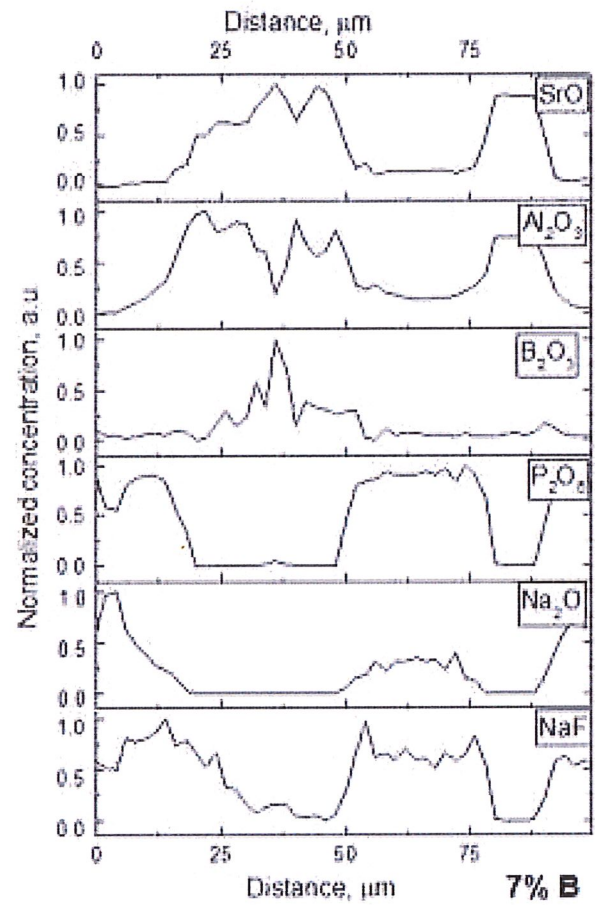
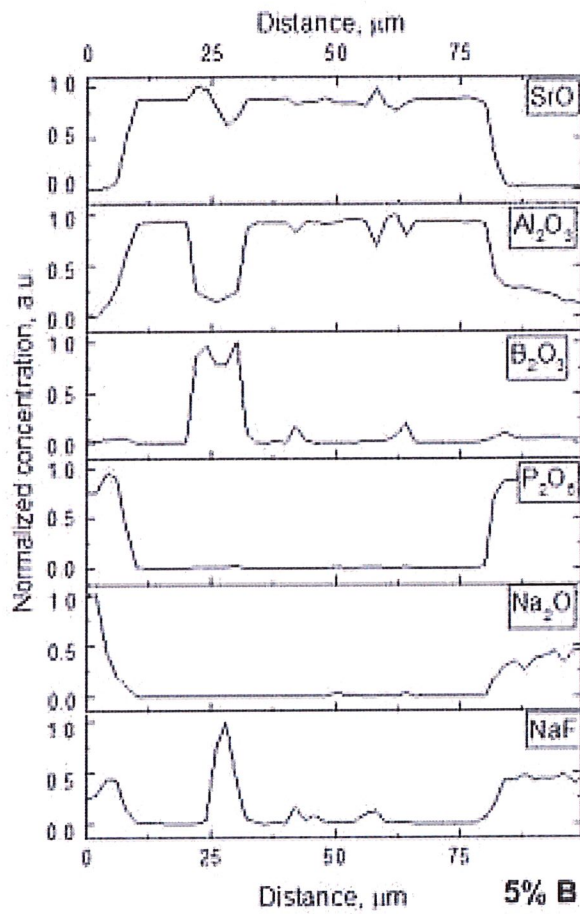
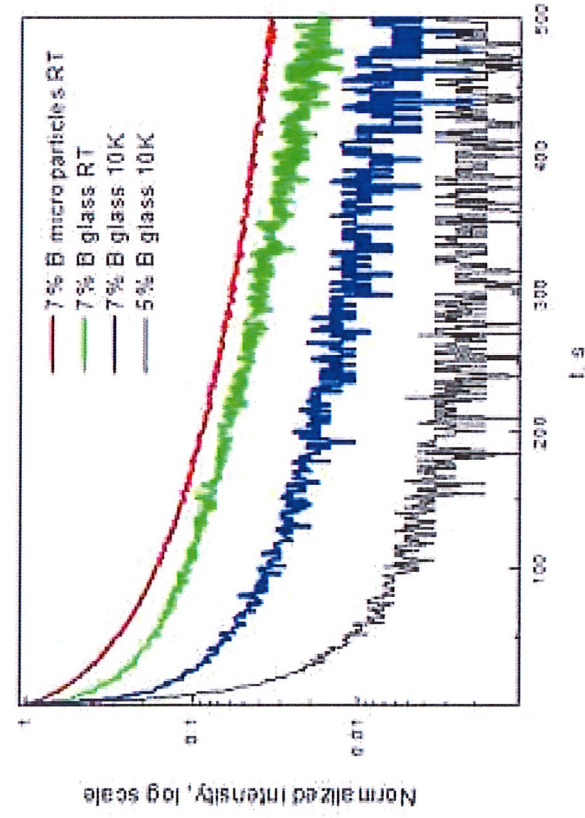
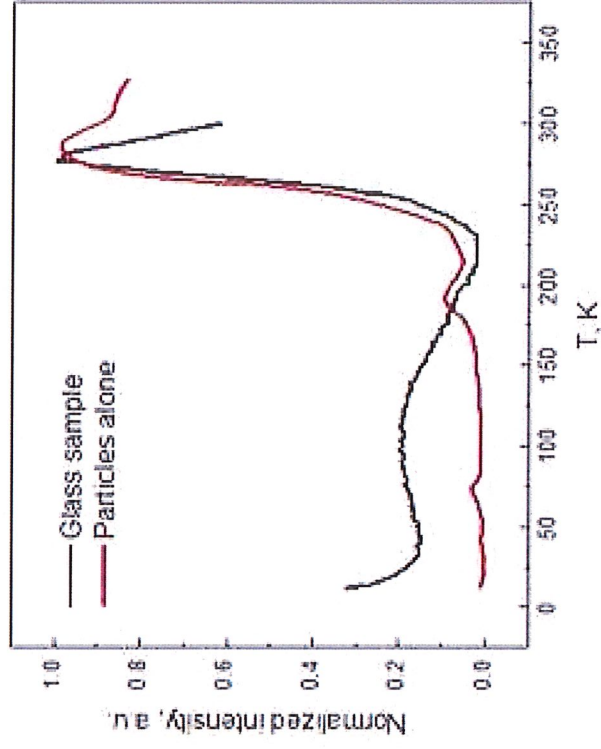


Figure 4
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