

Synthesis and Luminescence Properties of Doped Magnesium Boro-Tellurite Ceramics

Nur Zu Ira Boharia,* R. Hussina, Zuhairi Ibrahima, Hendrik O. Lintangb

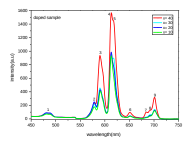
^aPhosphor Research Group, Department of Physics, Faculty of Science, *Universiti Teknologi Malaysia, 81310 UTM Johor Bahru, Johor Malaysia* ^bIbnu Sina Institute for Fundamental Science Studies, Universiti Teknologi Malaysia, 81310 UTM Johor Bahru, Johor Malaysia

*Corresponding author: nzuira@gmail.com

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Graphical abstract



Abstract

Glass has been widely utilized in the field of lighting, telecommunication and spectroscopy. Boro-tellurite is one of the suitable glasses used for solid state lighting and laser application. The investigation on the luminescence properties of rare earth doped ceramic is rarely used due to the opacity. In this paper boro-tellurite prepared in ceramic can show the better luminescence with the less advantage. The aim of this paper is to present the effect and advantages in luminescence results of boro-tellurite ceramics doped with the constant amount of rare earth. Doped magnesium boro-tellurite with Eu³+ and Dy³+ ceramic have been prepared using solid state reaction method with the compositions of xTeO₂-(70-x)B₂O₃-30MgO with 10≤x≤40, and have been doped with Eu₂O₃ (1mol%) and Dy₂O₃ (1mol%) . The characterizations of the samples have been investigated by means of X-Ray diffraction, Raman, Infrared and Photoluminescence spectroscopy. From the X-ray diffraction results, two phases are assigned to MgTe₂O₃ and Mg₂B₂O₃. Raman spectroscopy showed strong bands observed in the vicinity of 140, 175, 220, 266, 332, 403, 436, 646, 694, 723, 757 and 806 cm⁻¹. FTIR spectra showed bands located in the range between 400-800 cm⁻¹ are assigned to the bending mode of Te-O-Te, TeO₃ and TeO₄. In the range of 800-1400 cm⁻¹, the bands are associated with B-O, B-O-B, BO₃ and BO₄ bonds. The emission transition ⁵D₀⁻¬T₂ corresponded to the red emission (612 nm) was found to be the most intense in all the Eu³+-doped magnesium boro-tellurite ceramics.

Keywords: Magnesium boro-tellurite ceramic, luminescence, doped Eu3+ and Dy3+

Abstrak

Kaca telah digunakan secara meluas dalam bidang pencahayaan, telekomunikasi dan spektroskopi. Borotellurite adalah satu kaca yang sesuai digunakan dalam pencahayaan keadaan pepejal dan aplikasi laser. Penyelidikan luminesens bagi nadir bumi dop seramik adalah jarang digunakan disebabkan oleh kelegapan. Dalam kertas ini boro-tellurite seramik menunjukkan luminesens yang baik dengan kelebihan kurang. Matlamat kertas ini adalah untuk menunjukkan kesan dan kelebihan dalam keputusan luminesens bagi seramik boro-tellurite dop dengan amaun nadir bumi yang tetap. Dop magnesium boro-tellurite dengan Eu³+ and Dy³+ seramik disediakan dengan menggunakan kaedah tindakbalas keadaan pepejal yang berkomposisi xTeO₂-(70-x)B₂O₃-30MgO dengan 10≤x≤40, dan dop Eu₂O₃ (1mol%) and Dy₂O₃ (1mol%). Pencirian sampel disiasat oleh spektroskopi pembelauan sinar-X, Raman, infra merah and Fotoluminesens. Dari keputusan XRD, dua fasa dilihat sebagai MgTe₂O₅ and Mg₂B₂O₃. Spektroskopi Raman menunjukkan jalur kuat dilihat sekitar 140, 175, 220, 266, 332, 403, 436, 646, 694, 723, 757 dan 806 cm⁻¹. Spektra FTIR menunjukkan jalur terletak dalam lingkungan 400-800 cm⁻¹ adalah mod lentur bagi Te-O-Te, TeO₃ and TeO₄. Dalam lingkungan 800-1400 cm⁻¹, jalur yang berkaitan adalah B-O, B-O-B, BO₃ and BO₄. Peralihan pancaran ⁵D₀-⁷F₂ yang merujuk kepada pancaran merah (612 nm) didapati menjadi yang paling tinggi dalam semua seramik Eu³+-dop magnesium boro-tellurite.

Kata kunci: Seramik magnesium boro-tellurite, luminesens, dop Eu³⁺ and Dy³⁺

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■1.0 INTRODUCTION

Ceramic based on boro-tellurite host matrices doped with rare earth have tremendous applications for lasers, optical amplifier, photo sensitivity, optical storage, and bio-ceramics materials [1-8]. Currently, a great deal of research has been focused on rare earth

(RE) doped glasses owing to their extensive applications [9-10]. But, the investigation on the luminescence properties of rare earth doped ceramic is rarely used due to the opacity and hence, limited their applications. Eu³⁺ ion doped sodium-aluminum-tellurite (Eu³⁺: NAT) was prepared in opaque ceramics compound [11].

The host material with low phonon energy that can reduce the non-radiative loss, play important role for obtaining highly efficient luminescent properties via multiphonon relaxations and thus achieve strong luminescence [12]. In this study, tellurite oxide based has been utilized due to their desirable physical properties, such as high refractive index, excellent infrared transmittance and high dielectric constant, good chemical durability and low melting temperature. On the other hand, borate has important physical properties such as high transparency, low melting point, high thermal stability, good rare earth ions solubility, resistance to

■2.0 EXPERIMENTAL

High purity of H₃BO₃ (99.99%), TeO₂ (99.99%), MgO (99.99%), Eu₂O₃ (99.99%) and Dy₂O₃ (99.99%) were used as the raw materials. Two types of sample were prepared for the study; magnesium boro-tellurite ceramic and magnesium boro-tellurite with Eu³⁺ and Dy³⁺ ceramic. The samples with the compositions of *x*TeO₂-(70-*x*)B₂O₃-30MgO with 10≤*x*≤40 mol%, doped with Eu₂O₃(1mol%) and Dy₂O₃(1mol%) have been prepared using solid state reaction method. Analytical grade reagents of H₃BO₃, TeO₂, MgO, Eu₂O₃ and Dy₂O₃ powders in appropriate amounts (mol%) were thoroughly mixed in agate mortar. The mixtures were pressed into pellets using hydraulic press with 10 ton/cm² of pressure. Pelletized samples were then heat treated at 650 °C for 6 hours. The dried pellet was grounded into fine powder for further characterizations.

Several experiments have been setup to determine the physical nature and characteristics of the prepared samples. The examination of the structure was examined using X-ray diffraction (XRD) using Siemen Diffraction D500 diffractometer with CuK α radiation [13]. Fourier transform infrared spectroscopy (FTIR) was examined using Perkin-Elmer spectrometer (Spectrum 100) [13]. Raman spectroscopy was examined using Bruker model RFS 100/S FT-Raman spectrometer. The emission spectra were obtained using Jasco Fluorescence spectrophotometer, Model FP8000 series, at room temperature.

■3.0 RESULTS AND DISCUSSION

X-ray diffraction technique was used to identify the crystalline phase of the doped magnesium boro-tellurite with Eu³⁺ and Dy³⁺ ceramics. The XRD patterns, Figure 1, show the influence of varying composition of the xTeO₂-(70-x)B₂O₃-30MgO with 10≤x≤40 mol%. The x-ray diffractogram showed well-defined peaks which indicate the crystalline and phase formation of the synthesized compounds. There are two main phases; namely MgTe₂O₅ (ICDD: 01-073-3922) and Mg₂B₂O₅ (ICDD: 01-073-2107). The intensity increased with B₂O₃ and decrease with TeO₂, ranging from x = 10 to 40. MgTe₂O₅ became a dominant phase in the sample followed by Mg₂B₂O₅ phase. The majority phase of MgTe₂O₅ is an evident that the TeO₂ contribute to the formation of the phase. The percentage of the crystalline phase can be estimated by comparing the highest peak heights summarized in the Table 1. From the estimation of Table 1, it can be concluded that the MgTe₂O₅ was the dominant phase.

chemicals and can enhance mechanical durability [3]. Addition of alkali oxide (MgO) as network modifier has also been employed since the utilization of modifier doped with boro-tellurite based are rarely reported.

The aim of this work is to prepare doped magnesium borotellurite ceramics via solid state reaction route and to study the behavior through X-Ray Diffraction (XRD), Infrared (IR), Raman and Photoluminescence (PL) spectroscopic for both materials. The concentration of Eu³⁺ and Dy³⁺ will remain constant.

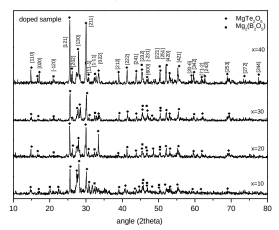


Figure 1 X-ray diffraction pattern of $x\text{TeO}_2$ -(70-x)B₂O₃-30MgO doped Eu³⁺ and Dy³⁺ ceramic (10 $\le x \le 40$ mol%)

Table 1 The estimation of percentage of the crystalline phase of xTeO2- $(70-x)B_2O_3$ -30MgO doped Eu³⁺ and Dy³⁺ ceramic $(10 \le x \le 40 \text{ mol}\%)$

x	MgTe ₂ O ₅ phase	Mg ₂ B ₂ O ₅ phase
	(%)	(%)
10	58.03	41.97
20	62.94	37.05
30	67.94	32.06
40	76.00	23.99

Raman spectrum of $x\text{TeO}_2$ -(70-x)B₂O₃-30MgO are presented in Figure 2. Raman spectroscopy show the strong band are observed at 140, 175, 220, 266, 332, 403, 436, 646, 694, 723, 757 and 806 cm⁻¹. Raman spectrum shows that besides the expected TeO₄, TeO-Te, and BO₃ (peak 7 to 12) existent, the vibrations corresponding to the metal ions are also present in ceramics (peak 1 to 6), respectively.

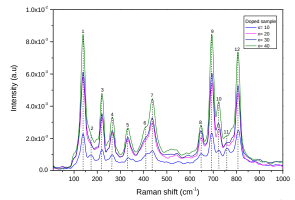


Figure 2 Raman spectra of xTeO₂-(70-x)B₂O₃-30MgO doped Eu³⁺ and Dy³⁺ ceramic (10 \le x \le 40 mol%)

The infrared spectrum of $x\text{TeO}_2$ - $(70-x)\text{B}_2\text{O}_3$ -30MgO with $10 \le x \le 40$ mol% in the range 1400- 4000cm^{-1} are presented in the Figure 3. There are four bands in this range when the peaks represent OH-and H- units are at around 3215 cm^{-1} , 2920 cm^{-1} and 2261 cm^{-1} . The peaks observed between 3375 cm^{-1} and 4000 cm^{-1} are the hydroxyl groups due to OH- stretching and the peaks observed between $(2800\text{-}2900)\text{ cm}^{-1}$ are the hydrogen bond [9]. Based on the Figure 3, the peaks around 1500 cm^{-1} are due to the B-O stretching of BO₃ units which are observed and the band became smaller (x=40), indicating a decrease in TeO₂ amounts and increasing B₂O₃.

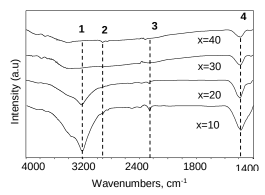


Figure 3 IR spectra of xTeO₂-(70-x)B₂O₃-30MgO doped Eu³⁺ and Dy³⁺ ceramic (10 \le x \le 40 mol%)

In Figure 4, the IR spectra of *x*TeO₂-(70-*x*)B₂O₃-30MgO with 10≤*x*≤40 mol% in the range (800-1400) cm⁻¹ is presented. The bands located in the range between (400-800) cm⁻¹ are assigned to the bending mode of Te-O-Te, TeO₃ and TeO₄. In the range of (800-1400) cm⁻¹, the B-O bond of B-O-B, BO₃ and BO₄ also appeared. The bands located around (1331-1362) cm⁻¹ are assigned to BO₃ units [15, 21] and in the range between (881-1200) cm⁻¹ which is the BO₄ units [10, 14, 16]. The bands positions around 431 cm⁻¹-564 cm⁻¹, 610-680 cm⁻¹ and 700- 780 cm⁻¹ showed the Te-O-Te, TeO₂ and TeO₃ respectively. All of these assignments are in good agreement with the reported literature [10, 17-18]. In Figure 4, the band gradually increased with TeO₂. The intensity of the band in the range of 431-780 cm⁻¹ increased gradually with TeO₂ and the intensity of the band slightly decreased with B₂O₃ in the range of 881-1362 cm⁻¹.

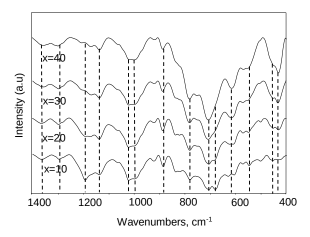


Figure 4 IR spectra of xTeO₂-(70-x)B₂O₃-30MgO doped Eu³⁺ and Dy³⁺ ceramic ($10 \le x \le 40$ mol%)

The emission spectrum of doped $x\text{TeO}_2$ - $(70-x)\text{B}_2\text{O}_3$ -30MgO with $10 \le x \le 40 \text{ mol}\%$ is shown in Figure 5. The emission transitions

 $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_0$, $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_0$, $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_1$, $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_2$, $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_4$, $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_4$, $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_4$, $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_4$ and $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_4$ corresponded to the band position at 484.58, 578.17, 589.89, 612.82, 618.38, 651.23, 683.48, 691.53 and 700.85 nm, respectively. It can be noted that a series of Eu³⁺ characteristic emission lines are between 540 nm and 780 nm, which includes the most prominent peak at 612 nm and corresponding to the $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_J(J=0, 1, 2, 3, 4)$ transitions [19]. The luminescence spectrum was found to be lowest for the sample x=10 followed by x=20, 30 and 40. The $^5\mathrm{D}_0 \rightarrow ^7\mathrm{F}_2$ transition is more intense than the other transitions as reported in the other literatures [20-22].

The emission peaks of Dy^{3+} were observed at 484.58 nm and 578.17 nm (peak 1 and 2) similar with the other reported values [23]. Figure 5 shows the emission peaks of Eu^{3+} are more dominant with high intensity compared the peaks of Dy^{3+} . The Dy^{3+} emission peaks are not present as dominant peak which indicates that Dy^{3+} acts as trap centers that cause long afterglow characteristics, rather than the luminescent centers in the host lattice [24].

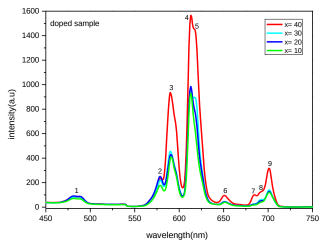


Figure 5 Emission spectrum of xTeO₂-(70-x)B₂O₃-30MgO doped Eu³⁺ and Dy³⁺ ceramic (10 \le x \le 40 mol%)

■4.0 CONCLUSION

The samples of xTeO₂-(70-x)B₂O₃-30MgO with $10 \le x \le 40 \text{ mol}\%$ doped with Eu₂O₃(1mol%) and Dy₂O₃(1mol%) have been prepared using solid state route. The structural studies of the samples have been investigated using X-ray diffraction (XRD), Infrared and Raman spectroscopy. The XRD profiles show that MgTe₂O₅ became a dominant phase in the sample followed by Mg₂B₂O₅ phase. The IR spectrum shows two series of band regions which obtained in 4000-1400cm⁻¹ and the second region of 1400-400cm⁻¹. The ceramics doped with Eu³⁺ shows a bright red emission at 612nm which belongs to the electric dipole (5 D₀ \rightarrow ⁷F₂) transition of Eu³⁺ ions. There are two groups of emission at 484 nm and 578 nm occurring in the emission spectrum of the ceramics, which probably can be applied for white LEDs application.

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