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New Infrared Luminescence from Bi-doped Glasses

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1. Introduction

Infrared luminescent materials are widely used as laser media, for example, Nd:YAG and Er-doped silica fibers. In the infrared region, luminescence offers many advantages as laser media due to the variety of optics and excitation sources, such as semiconductor lasers and flash lamps. A frequency-doubled laser-beam with a nonlinear crystal can also obtained in the visible region. Therefore, infrared light source materials are very useful for scientific research and industrial uses.

Fujimoto et al. discovered a new infrared luminescent from bismuth-doped silica glass (BiSG)(Fujimoto & Nakatsuka 2001). Wide luminescent exists between 1000 and 1600 nm, and absorption is in the visible region. The lifetime is 630 µs at room temperature. Since this silica glass-based material has luminescence around 1.3 µm of zero-dispersion wavelength, it can be used as a core material of an optical fiber amplifier for telecommunication in the 1.3- μ m range. Fujimoto et al. also demonstrated an optical amplification at 1.3 μ m with 0.8µm excitation(Fujimoto & Nakatsuka 2003). Therefore, BiSG is expected to be a promising substitution for Pr³⁺:ZBLAN fiber or Raman amplifiers in the 1.3-µm region.

The spectroscopic properties of BiSG are different from previously reported Bi luminescent material(Fujimoto & Nakatsuka 2001), because the previous Bi-doped glasses or crystals with low Bi_2O_3 concentration showed that luminescences exist in the visible region (400 ~ 600 nm) and that absorptions are in the ultraviolet region. The lifetimes ($\sim 4 \mu s$) are much shorter than those of BiSG at room temperature. Such luminescences have been reported to originate from the Bi³⁺ ion in the media(M. J. Weber & Monchamp 1973; Parke & Webb 1973; van der Steen, van Hesteren et al. 1981). BiSG is also different from the Bi₂O₃-rich glass whose Bi₂O₃ concentration ranges from 18.7 to 42.5 mol%(Sugimoto, Kanbara et al. 1996) because the glass has no absorption in the visible region. Then the investigation of new glass compositions with Bi luminescence is also progressing vigorously in a past decade. Although the first discovered glass was a silica-based material with a SiO_2 of 97.5 mol%(Fujimoto & Nakatsuka 2001), many glass compositions were tested and shown to effectively generate Bi luminescence, such as silicate(Peng, Qiu et al. 2005; Ren, Yang et al. 2006; Suzuki & Ohishi 2006; Arai, Suzuki et al. 2007; Murata & Mouri 2007; Peng, Chen et al. 2007; Ren, Qiu et al. 2007; Zhou, Feng et al. 2007; Peng, Wu et al. 2008), germanate(Peng, Qiu et al. 2004; Meng, Qiu et al. 2005; Peng, Meng et al. 2005; Peng, Qiu et al. 2005; Peng, Wang et al. 2005; Wang & Xia 2006; Xia & Wang 2006; Murata & Mouri 2007; Ren, Wu et al. 2007; Source: Advances in Solid-State Lasers: Development and Applications, Book edited by: Mikhail Grishin, ISBN 978-953-7619-80-0, pp. 630, February 2010, INTECH, Croatia, downloaded from SCIYO.COM

Ren, Dong et al. 2007; Ren, Qiao et al. 2007; Ren, Qiu et al. 2007; Ren, Qiu et al. 2007; Peng, Wu et al. 2008; Qiu, Peng et al. 2008), borate(Meng, Qiu et al. 2005; Murata & Mouri 2007; Qiu, Peng et al. 2008), germanosilicate(Ren, Dong et al. 2007; Ren, Dong et al. 2008), and phosphate(Meng, Qiu et al. 2005; Qiu, Peng et al. 2008) glasses.

In this chapter, the author will introduce the basic properties of Bi doped silica glass (BiSG), such as a phase diagram and spectroscopic properties, and then mainly talk about the origine of luminescent center.

2. Basic properties of BiSG

2.1 Phase diagram

A phase diagram of BiSG was obtained for the Bi₂O₃-Al₂O₃-SiO₂ glass system. Silica powder (Aerosil 50; SiO₂, 99.8%), bismuth-oxide (Kojundo Chemical Lab. Co., Ltd.; α -Bi₂O₃, 99.99%), and aluminum oxide (Kojundo Chemical Lab. Co., Ltd.; α -Al₂O₃, 99.99%) were used as reagents. A properly mixed powder was inserted into a silica tube with an inner/outer diameter of 1/3 mm ϕ , and then the tube was heated by a mixed gas burner of natural gas and oxygen. After heating, the powder was checked for a distinctive glassy wetting. If it had such wetting, the composition was determined to be the glassy phase in the Bi₂O₃-Al₂O₃-SiO₂ glass system.

The measured phase diagram of the Bi₂O₃-Al₂O₃-SiO₂ glass system is shown in Fig. 1. The circle, triangle, and cross points show the glassy wetting states (pink to reddish-brown), unclear glassy states (pink to reddish-brown), and no glassy state (reddish-brown), respectively. This phase diagram shows a tendency where the glass whose Al₂O₃ content is larger than Bi₂O₃ is well glassified above a 90% SiO₂ concentration. On the other hand, all the samples are glassified at a 80% SiO₂ concentration without dependence on the Bi₂O₃ or Al₂O₃ content. The infrared luminescent spectrum is observed until 10 mol% of Bi₂O₃ concentration.



Fig. 1. Measured phase diagram of Bi₂O₃-Al₂O₃-SiO₂ system.

2.2 Influence of additives on luminescent intensitys

The Bi₂O₃-Al₂O₃-SiO₂ glass system is affected by alkaline metal additive. Figure 2 shows an effect of Li₂O additive in BiSG. The glass composition is Bi₂O₃(1.0 mol%)-Al₂O₃(7.0 mol%)-SiO₂(92-x mol%)-Li₂O(x mol%) glass system. The luminescent intensity decreases with increasing Li₂O additive drastically (Seo, Fujimoto et al. 2006). Therefore, in order to get luminescent center, we should carefully choose the host composition.



Fig. 2. Luminescent intensity dependence on Li_2O concentration for $Bi_2O_3(1.0 \text{ mol}\%)$ - $Al_2O_3(7.0 \text{ mol}\%)$ - $SiO_2(92-x \text{ mol}\%)$ - $Li_2O(x \text{ mol}\%)$ glass system.



Fig. 3. Dependence of lumienscent intensity on GeO₂ additive. The filled circle denotes BiSG with 0.1 mol% Bi₂O₃ concentration, while the filled square represents that of 1.0 mol% Bi₂O₃ concentration. The nomenclature of the samples is as follows: for example, the sample name "BiSG_1.0_5.0" means that the composition is Bi₂O₃ = 1.0 mol%, GeO₂ = 5.0 mol%.

On the other hand, GeO_2 is the one of attractive additive to increase luminescent intensity of BiSG (Fujimoto, Hirata et al. 2007). Figure 3 shows the dependence of luminescent intensity on GeO₂ concentration. All the samples contain 7.0 mol% of Al2O3. The nomenclature of the samples is as follows: for example, the sample name "BiSG_1.0_5.0" means that the composition is Bi₂O₃ = 1.0 mol%, GeO₂ = 5.0 mol%. The luminescent intensity of BiSG grows with increasing GeO₂ concentration almost linearly for BiSG_0.1_xx samples, and the luminescent intensity of BiSG_0.1_5.0 is 26.3 times larger than that of BiSG_0.1_0.0 only because of the GeO₂ additive. The increment effect on BiSG with GeO₂ additive is also observed for a Bi₂O₃ concentration of 1.0 mol% (BiSG_1.0_5.0), but it is less remarkable than for the lower Bi₂O₃ concentration of 0.1 mol%. Furthermore, the increment is only about 1.5 times larger than for BiSG_0.1_5.0 is almost same as that of BiSG_1.0_0.0. According to these data, the GeO₂ additive effectively increases the luminescent intensity of BiSG, and this effect is especially remarkable at low Bi₂O₃ concentrations.

2.3 Spectroscopic properties

The typical spectroscopic properties of BiSG are shown in Fig. 4. Five main absorption bands exist between 250 and 2000 nm: 300 (A), 500 (B), 700 (C), 800 (D), and 1000 nm (E) (Fig. 5(a)). It is considered that absorption band A is mainly derived from O^{2-} in the base glass(Scholze 1991). The other four absorptions (B~E) are connected to the infrared emissions shown in Fig. 5(b). Each luminescent peak wavelength and full width at half maximum (FWHM) were recognized as 1182 and 300 nm at 500-nm excitation, 1144 and 192 nm at 700 nm, 1265 and 336 nm at 800 nm, and 1130 and 188 nm at 1000 nm. A similarity of the luminescent spectra can be seen between 700- and 1000-nm excitations. These spectroscopic properties are different from the Bi³⁺ ones(M. J. Weber & Monchamp 1973; Parke & Webb 1973; van der Steen, van Hesteren et al. 1981)(Table 1).

	Bi ³⁺ :Glass*1 (Na ₂ O•P ₂ O ₅)	Bi ³⁺ :Crystal*2 (Bi ₃ Ge ₄ O ₁₂)	BiSG *3
Excitation wavelength [nm]	250	290	500~800
Luminescent wavelength [nm]	390	480	$1000 \sim 1400$
Lifetime [µs] (room temp.)	3.9	0.1	630

*1 (Parke & Webb 1973), *2 (M. J. Weber & Monchamp 1973), *3 (Fujimoto & Nakatsuka 2001) Table 1. Differences of spectroscopic properties between BiSG and previously reported Bi³⁺ luminescent material.

3. The origin of luminescent center

After the discovery of a new infrared luminescent bismuth center, several research groups try to understand what is the origine of the luminescence. Although there have been many previous discussions on this distinctive luminescent center, it has remained unclear. Different authors tentatively assigned the different origins of this near infrared luminescence to the electronic transition derived from Bi⁵⁺(Fujimoto & Nakatsuka 2001; Dvoyrin, Mashinsky et al. 2006; Fujimoto & Nakatsuka 2006; Wang & Xia 2006; Xia & Wang 2006; Ohkura, Fujimoto et al. 2007), Bi²⁺(Ren, Qiu et al. 2007; Ren, Qiu et al. 2007; Ren, Qiu et al. 2005; Meng, Qiu et al. 2005;

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Fig. 4. Typical spectroscopic properties of BiSG (sample B1 in Table 2): (a) transmittance spectrum shows five main absorption bands between 250 and 2000 nm, including 300 nm (A), 500 nm (B), 700 nm (C), 800 nm (D), and 1000 nm (E), and (b) luminescent spectra excited by 500, 700, 800, and 1000 nm.

Dvoyrin, Mashinsky et al. 2006; Arai, Suzuki et al. 2007; Ren, Qiu et al. 2007; Ren, Qiu et al. 2007; Ren, Qiu et al. 2007; Yang, Chen et al. 2007; Zhou, Feng et al. 2007; Qiu, Peng et al. 2008; Ren, Dong et al. 2008; Truong, Bigot et al. 2008) or to that of BiO molecules(Ren, Yang et al. 2006; Murata & Mouri 2007; Peng, Chen et al. 2007; Peng, Wu et al. 2008) dispersed in the glass host. Since most of these proposals were however analogized from only spectroscopic mesurement analysis, uncertainty remains in the present discussions. In addition, even the same researchers offer several explanations, suggesting that spectroscopic analogy is insufficient to correctly understand the luminescent center or the luminescent mechanism.

Based on this viewpoint, our group discarded the spectroscopic analogical approach to understand the structure of the luminescent center or its mechanism and instead chose such instrumental analytical approaches as nuclear magnetic resonance (NMR), x-ray diffraction (XRD), electron spin resonance (ESR), x-ray photoelectron spectroscopy (XPS), and extended x-ray absorption fine structure (EXAFS). NMR data revealed the coordination structure of the specific ions in a material, ESR and XPS data provided the valence state of ions, and EXAFS offered the bonding distance between the specific and neighboring ions. Once these analytic instrumental data are integrated to build one possible figure of a luminescent center, then the luminescent mechanism will be discussed, because knowledge of the luminescent center is very useful for understanding the luminescent mechanism, which is obviously important for such optical applications as lasers or amplifiers.

In the following paragraphs, let's try to build up an image of the local structure of the Bi luminescent center based on the analytical data, such as, spectroscopic properties (SPCT), luminesent intensity (LMI), NMR, XRD, ESR, XPS, and the EXAFS data of BiSG in a simple Bi₂O₃-Al₂O₃-SiO₂ glass system without discrepancies. From this approach, we will reach one strongly possible local structure of a distinctive Bi luminescent center that consists of Bi and Al that exist closely to each other.

3.1 Sample preparation

In these analyses, composition of BiSG is very important, because Bi infrared luminescent center, as shown in section 2.2, is disappeared with increasing some additives in the host. This means that majority part of Bi ions in the host shifts to non-emissive site, thus even if we choose the multi-component glass system and analyze the Bi centers, we could not judge the results form an inherent luminescent center or the other non-emissive site, then we would go into the complicated maze. Therefore, the analyses should be carried out to concentrate on the simple three components system, Bi_2O_3 -Al_2O_3-SiO_2 glass system. The samples for the instrumental analyses were prepared as follows.

- 1. A-series: The ratio of Bi₂O₃ to Al₂O₃ was kept at 3/7, and the Bi₂O₃ concentration varied from 0.1, 0.3, 0.5, 1.0, and 3.0 mol%. These samples were called A1-A5, respectively. Their compositions are inside the glassy phase area in Fig. 1.
- 2. B-series: B1 was composed of Bi₂O₃ (1.0 mol%), Al₂O₃ (7.0 mol%), and SiO₂ (92.0 mol%). This was used for the measurement of the spectroscopic properties because Al₂O₃-rich BiSG includes few bubbles. B2 was composed of Bi₂O₃ (0.3 mol%), Al₂O₃ (2.3 mol%), and SiO₂ (97.4 mol%) and B3 of Bi₂O₃ (3.0 mol%), Al₂O₃ (7.0 mol%), and SiO₂ (90.0 mol%). They were used for the ESR measurements.
- 3. C-series: C1 was composed of Bi₂O₃ (1.0 mol%) and SiO₂ (99.0 mol%) without Al₂O₃, and C2 was composed of Al₂O₃ (2.3 mol%) and SiO₂ (97.7 mol%) without Bi₂O₃. C1 and C2 were dark brown and colorless, respectively. C1 had no glassy wetting, but C2 did.

The chemical composition of the investigated BiSG samples and the several standards are listed in Table 2. Silica powder (Aerosil 50; SiO₂, 99.8%), bismuth-oxide (Kojundo Chemical Lab. Co., Ltd.; α -Bi₂O₃, 99.99%), and aluminum oxide (Kojundo Chemical Lab. Co., Ltd.; α -Al₂O₃, 99.99%) were used as reagents to make the glass samples of the Bi₂O₃-Al₂O₃-SiO₂ glass system. The mixtures of these powders were melted in an aluminum crucible in an electric furnace at 1750°C in air, and the melts were annealed by natural cooling. All of these glasses were reddish-brown. NaBiO₃ (Kanto Chemical Co., Inc.; >80%), which was identified as NaBiO₃•2H₂O by XRD(PDF#30-1161 2000), α -Bi₂O₃, and Bi-metal (Kojundo Chemical Lab. Co., Ltd.; 99.9%) were used as standard materials for the analyses.

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Sample	Bi ₂ O ₃ [mol%]	$Al_2O_3[mol\%]$	SiO ₂ [mol%]	Measurements		
<bisg></bisg>						
A1	0.1	0.23	99.67	LMI, NMR, XRD		
A2	0.3	0.7	99.0	LMI, NMR, XRD, EXAFS		
_A3	0.5	1.15	98.35	LMI, NMR, XRD, EXAFS		
A4	1.0	2.3	96.7	LMI, NMR, XRD, EXAFS, XPS		
A5, B3	3.0	7.0	90.0	LMI, NMR, ESR, XRD, EXAFS, XPS		
B1	1.0	7.0	92.0	SPCT		
B2	0.3	2.3	97.4	ESR		
C1	1.0	0.0	99.0	LMI		
C2	0.0	2.3	97.7	NMR, XRD		
<aluminium standard=""></aluminium>						
α - Al ₂ O ₃	0.0	100	0.0	NMR, XRD		
<bismuth standards=""></bismuth>						
NaBiO ₃				EXAFS, XPS		
α - Bi ₂ O ₃	100	0.0	0.0	EXAFS, XPS		
Bi metal				EXAFS, XPS		

LMI: Luminescent Intensity, NMR: ²⁷Al-NMR(Nuclear Magnetic Resonance), XRD: X-ray Diffraction, ESR: Electron Spin Resonance, XPS: X-ray Photoelectron Spectroscopy, EXAFS: Extended X-ray Absorption Fine Structure, SPCT: Spectroscopic Properties

Table 2. List of chemical composition and performed measurements for investigated Bi doped silica glasses and Al, Bi standards.

The prepared samples were evaluated by instrumental analyses, including spectroscopic measurements, NMR, XRD, ESR, XPS, and EXAFS. Luminescent intensity and spectra were measured by a spectrophotometer (JASCO Corporation; SS-25). Transmission spectra were obtained by a spectrophotometer (HITACHI; U-4100). The aluminum coordination state (ACS) in BiSG was determined by the ²⁷Al-NMR spectrum, and the ²⁷Al-NMR spectra were obtained on a nuclear magnetic resonance spectrometer (JEOL Ltd.; JNM-GSX-400). The XRD patterns were measured by an X-ray diffractometer (RIGAKU; RINT2500). The ESR signals were measured by an electron spin resonance spectrometer (JEOL Ltd.; JES-ME-2X) at both room and liquid nitrogen temperatures.

The XPS spectra were recorded on a SHIMADZU (KRATOS) AXIS 165 using MgK radiation (1253.6 eV) with a beam size of $0.7 \times 0.3 \text{ mm}^2$ and 50 eV analyzer pass energy. The operating vacuum pressure was maintained from 10^{-8} to 10^{-9} Torr. An electron beam was used to reduce the electrical charging during the measurements. Any binding energy shifts due to electrical charging were corrected by the C1s (284.6 eV) peak as an internal reference. 3-minute Ar-sputtering was used to remove the oxidized surface on the Bi-metal powder.

The Bismuth L_{III} XAFS spectra were measured in the transmission mode using a BL-7C beam line at the Photon Factory (PF). The ring energy was 2.5 GeV. The x-ray was monochromized by a Si(111) double crystal monochrometer. Two ionization chambers were used as detectors. A 170-mm length chamber was filled with Ar/N₂ (15%:85%) gas for

incident x-ray energy (I_0), and the other was 310-mm long with Ar-100% gas for transmitted x-ray energy (I). The EXAFS data of the Bi L_{III} edge (13426.5 eV) were collected between 12926 and 14526 eV with 481 energy points. Data analysis was carried out on *UWXAFS*. The back-scattering amplitude and the phase shift were theoretically calculated using *FEFF 8.2* code. The Debye-Waller factor was estimated by the Debye code implemented in *FEFF 8.2* based on Raman spectroscopy results(Narang, Patel et al. 1994).



Fig. 5. Dependence of luminescent intensity (LMI) on Bi_2O_3 concentration detected at 1120nm luminescence with 500-nm excitation.

The dependence of luminescent intensity (LMI) on Bi_2O_3 concentration is illustrated in Fig. 5. The measured samples were A-series (A1~A5) and C1. The excitation and detection wavelengths of the luminescence were at 500 and 1120 nm, respectively. The luminescent intensity nonlinearly increased with increased Bi_2O_3 concentration. At a 1.0 mol% of Bi_2O_3 concentration, the luminescent intensity from A4, which includes 2.3 mol% of Al_2O_3 , is three orders of magnitude larger than that of C1 without Al_2O_3 . Based on these results, we conclude the following:

- 1. Al₂O₃ additive can remarkably increase to generate a Bi luminescent center.
- 2. The generation of a Bi luminescent center has a nonlinear relation for Bi_2O_3 concentration.

3.4 ²⁷AI-NMR spectra

²⁷The Al-NMR spectra in BiSG are shown in Fig. 6(Fujimoto & Nakatsuka 2006). ²⁷Al chemical shifts were measured relative to $Al(H_2O)_6^{3+}$. The measured samples were A-series,

C2, and α -Al₂O₃. α -Al₂O₃ with a 6-fold coordinated state of corundum structure was used as a standard sample, and a peak exists at 15 ppm. The peaks at 70 and -40 ppm (marked by asterisks) were derived from spinning sidebands. The peaks of ²⁷Al-NMR from A1 to A3 only exist at 15 ppm and are the same as α -Al₂O₃. The A4 peak is still dominated by the 15 ppm peak, but a peak around 50 ppm begins to emerge, and then a peak of 56.4 ppm becomes dominant in A5 (Fig. 6(b)).



Fig. 6. ²⁷Al-NMR spectra of A1–A5, C2, and α -Al₂O₃: (a) whole view of spectra, and (b) expanded view of spectra in intensity scale

Sample A4, which includes Al_2O_3 of 2.3 mol%, has a weak 50 ppm peak in the ²⁷Al-NMR spectrum, while sample C2, which has the same amount of Al_2O_3 without Bi_2O_3 , shows no signal around 50 ppm (Fig. 6(b)), suggesting that the Bi ion affects ACS over a 1.0 mol% of Bi_2O_3 concentration. On the other hand, since the C2 spectrum is dominated by a peak at 15 ppm, the aluminum ions in the silica glass naturally configure the 6-fold coordinated state of the corundum structure up to 2.3 mol% of Al_2O_3 without Bi_2O_3 . This is also supported by the work of Mysen et al., who concluded the aluminum ions in silica glass work as a network modifier rather than a network former up to a 6.1 mol% Al_2O_3 concentration in the measurement of Raman spectra(Mysen, Virgo et al. 1980). The Al corundum structure clearly has certain important roles for the generation of the Bi luminescent center in BiSG.

3.5 XRD measurements

The XRD data of the BiSG samples were measured to check for existing crystallizations, including undissolved alumina, mullite, or crystbalite because crystallization influences the ²⁷Al-NMR spectra. The measured samples are A-series. Fig. 7 shows the XRD data on

samples A4, α -Al₂O₃ (alumina), and pure silica with a range between 10 and 80° in 20. Sample A4 is substituted for the other BiSG ones because these XRD patterns are almost the same. The peaks due to any kind of crystallization are not recognized in Fig. 7, especially undissolved alumina, where there is only a halo pattern. We previously confirmed an XRD pattern on a Nd₂O₃(3 wt%; 0.55 mol%)-SiO₂(97 wt%; 99.45 mol%) system that included undissolved 0.55 mol% of Nd₂O₃ at best(Fujimoto & Nakatsuka 1997). 7.0 mol% of alumina (at maximum in this experiment) is probably an adequate quantity for XRD detection if the quantity is changed to undissolved alumina or other crystals in the sample. Therefore, it is concluded that all our samples are in the amorphous phase.



Fig. 7. XRD patterns on α -Al₂O₃ (alumina), sample A4 and pure silica with range between 10° and 80° in 2 θ

3.6 ESR measurements

The presence of unpaired electrons in BiSG was verified by ESR signal. The measured sample was B2 and B3. There was no ESR signal due to the unpaired electrons for both B2 and B3, even at liquid N₂ temperature. The same phenomena without signals were also reported on Bi-doped multi-component glasses(Peng, Wang et al. 2005; Peng, Wu et al. 2008). According to Hund's rule, the valence states of bismuth ions without unpaired electrons should be $Bi^{3+}(\sim 5d^{10}6s^2)$ or $Bi^{5+}(\sim 5d^{10})$ (Ohkura, Fujimoto et al. 2007).

3.7 XPS measurements

3.7.1 Analysis on chemical shift

The results of the XPS measurements are shown in Fig. 8. The measured samples were A4, A5, and three standards, NaBiO₃, Bi₂O₃, and Bi-metal. The main Bi($4f_{5/2}$, $4f_{7/2}$) peaks of Bi₂O₃ exist at 163.7 and 158.4 eV, respectively. Bi-metal was treated with 3-minute etching by Arbeam in a vacuum chamber (1.0×10^{-7} Torr) to eliminate the oxidized Bi-metal surface before the measurement. Even after the treatment, weak residual peaks were found due to Bi₂O₃. The main Bi($4f_{5/2}$, $4f_{7/2}$) peaks of the Bi-metal exist at 162.4 and 157.1 eV, respectively. These Bi₂O₃ and Bi-metal peaks well agree with those previously reported(Wagner 1990; Saffarini

& Saiter 2000), and the chemical shifts of Bi_2O_3 and Bi-metal are very stable in the XPS measurement. NaBiO₃ is often used as a standard of the penta-valent state of Bi, but in our experiment, the main Bi(4f_{5/2}, 4f_{7/2}) peaks of NaBiO₃ were obtained at 163.8 and 158.5 eV corresponding to the Bi₂O₃ ones, and the second Bi(4f_{5/2}, 4f_{7/2}) peaks exist at higher bonding energy at 165.9 and 160.6 eV, respectively. The peaks of both BiSGs, that is, A4 and A5, are located at almost the same position at the second NaBiO₃ peaks. After arranging the chemical shifts for the measured samples, the order of the bonding energy is as follows:

[lower bonding energy] Bi metal (Bi⁰) -> Bi₂O₃ (Bi³⁺), first peaks of NaBiO₃ (Bi³⁺) -> second peaks of NaBiO₃ (Bi⁵⁺) = BiSG (A4, A5) [higher bonding energy]

In general, the valence state of the target ion becomes higher with increased bonding energy(Wagner 1990), and the same tendency is observed in my measurement. Therefore, Bi ions of the penta-valent state exist in BiSG.



Fig. 8. XPS peaks of Bi $(4f_{5/2}, 4f_{7/2})$ on A4, A5, NaBiO₃, Bi₂O₃, and Bi-metal. In observation order of binding energy, [lower bonding energy] Bi metal (Bi⁰) -> Bi₂O₃ (Bi³⁺), first peaks of NaBiO₃ (Bi³⁺) -> second peaks of NaBiO₃ (Bi⁵⁺) = BiSG (A4, A5) [higher bonding energy].

The peak positions of NaBiO₃, however, seem unstable. The previously reported Bi($4f_{5/2}$, $4f_{7/2}$) peaks(Kulkarni, Vijayakrishnan et al. 1990) were 164.1 and 158.7 eV, respectively, and they showed single peaks with almost the same binding energy of the Bi₂O₃ ones. In fact, although we measured the XPS data on the NaBiO₃ several times, the ratio of the main peaks corresponding to Bi₂O₃ to the second peaks was unstable. Therefore, we put the most probable XPS data of NaBiO₃ in Fig. 8.

Kumada et al. (Kumada, Takahashi et al. 1996; Kumada, Kinomura et al. 1999) reported that NaBiO₃ and LiBiO₃ are synthesized at 120-200°C and that the Bi⁵⁺ state is changed to a Bi³⁺ state over 400°C; similar unstability may occur for the Bi⁵⁺ state in NaBiO₃. The standard

material of NaBiO₃ was identified as NaBiO₃•2H₂O by XRD in Kumada's experiment. They also reported that Na ions in the A-site (A+B⁵⁺O₃) tend to be exchanged for Sr²⁺ or Ba²⁺ ions in NaBiO₃•nH₂O, and then the redistributed Bi ions in the A-site take the Bi³⁺ state(Kumada, Kinomura et al. 1999). Although they neglected to mention the redistribution of Bi ions in NaBiO₃ themselves, a similar phenomenon may occur for NaBiO₃ in their experiment.

3.7.2 Analysis on peak separation

By precisely observing the peak positions and the line widths, we recognized that the A4 peaks were slightly shifted to higher bonding energy than A5 and that the line widths of A4 and A5 were wider than the standards. Since the A4 and A5 peaks are composed of two or more peaks, we separated them with a Gaussian fitting curve to examine the origin of the peak shift. In this procedure, we make the following assumptions:

- 1. If such Bi ionic states as Bi^0 or Bi^{3+} are considered identical, the line widths of $Bi(4f_{5/2}, 4f_{7/2})$ are also identical.
- 2. The line widths of Bi $4f_{5/2}$ and Bi $4f_{7/2}$ are the same.
- 3. The ratio of Bi $4f_{5/2}$ to Bi $4f_{7/2}$ is constant for different ionic states in a sample. This ratio is theoretically calculated as Bi $4f_{5/2}$ /Bi $4f_{7/2}$ =l/(l+1)=3/4(Seah 1983).

The results are shown in Table 3. The peak separation results show five peak positions for all Bi $4f_{5/2}$ and Bi $4f_{7/2}$ peaks that are normalized at 100. No. 5 corresponds to Bi⁰, No. 4 to Bi³⁺, and No. 1 to Bi⁵⁺. Nos. 2 and 3 are the intermediate states between numbers 1 and 4, and these states have intermediate coordination states rather than intermediate valence states such as Bi⁴⁺ due to the ESR measurements. These results show that all the Bi ions in BiSG are not the penta-valent state, and therefore the Bi valence states were mixed states of Bi³⁺ with Bi⁵⁺. This mixed valence state of Bi³⁺ and Bi⁵⁺ is also supported by EXAFS analysis

Sample		Bi 4f _{5/2} (Bi 4f _{7/2})						
		1	2	3	4	5		
	Peak [eV]	166.0(160.6)	164.9(159.5)					
A4	FWHM [eV]	1.7	1.7					
	Height	100.0	64.6					
	Peak [eV]	166.2(160.8)	165.1(159.8)					
A5	FWHM [eV]	1.9	1.9					
	Height	50.6	100.0					
	Peak [eV]	165.9(160.6)		164.6(159.3)	163.8(158.5)			
NaBiO ₃	FWHM [eV]	1.8		1.4	1.4			
	Height	38.6		35.0	100.0			
	Peak [eV]			164.7(159.4)	163.7(158.4)			
α-Bi ₂ O ₃	FWHM [eV]			1.4	1.4			
	Height			19.9	100.0			
Bi metal	Peak [eV]			164.6(159.3)	163.4(158.1)	162.4(157.1)		
	FWHM [eV]			1.4	1.4	1.1		
	Height			22.4	22.4	100.0		

Table 3. Peak separation results on $Bi(4f_{5/2}, 4f_{7/2})$ of A4, A5, $NaBiO_3$, α - Bi_2O_3 , and Bi-metal. Peak position, FWHM, and normalized peak height are listed. Peak-heights are normalized at 100.

in the next section. The peak height ratio of Nos. 1 and 2 is counterchanged for A4 and A5 due to the existence ratio of Bi³⁺ and Bi⁵⁺. Thus, the peaks of A4 are slightly shifted due to higher bonding energy than A5.

3.8 Bi-O distance from EXAFS

Figure 9 shows the radial structure functions (RSF) to which the EXAFS oscillations were Fourier-transformed. The measured samples were A-series (A2~A5) and the two standards of α -Bi₂O₃ and NaBiO₃. The peak shown in about 1.0 Å is derived from the XANES region because it is too short for any Bi-O distance. Therefore, it is a ghost peak, and we conclude that the largest peak around 1.6-1.7 Å (the first relevant peak) corresponds to the first neighboring Bi-O bond. α -Bi₂O₃ and NaBiO₃ have the second peak at 3.5 and 3.2 Å, respectively. Since the BiSG ones have no secondary peak, the local environment of the Bi ion does not have any periodical structure; that is, BiSG should be an amorphous phase. These results are supported by the XRD data in Section 3.5. The RSF shows that all the BiSG peaks are about 0.1 Å shorter than α -Bi₂O₃. The NaBiO₃ peak is also shifted to a shorter position, but the line width is wider than that of any of the BiSG ones. Since RSF |F(r)|includes a phase shift, the radial distance in RSF shifted a shorter range than the actual Bi-O distance. To determine the length of the first neighboring Bi-O, the RSF of α -Bi₂O₃ and the BiSG samples were analyzed by the curve-fitting method in r-space with FEFF 8.2. In this curve-fitting calculation, we only took two coordination spheres due to the parameter number limitation in FEFF 8.2.



Fig. 9. Radial structure functions (RSF) of A-series (A2~A5), α -Bi₂O₃ and NaBiO₃ The fitting results of the BiSG samples, α -Bi₂O₃ and NaBiO₃, are listed in Table 4. We assumed amplitude reduction factor S₀² = 0.9(Manzini, Lottici et al. 1998) and absorption

edge energy $E_0 = 13426.5$ eV. The fitting range was selected from 1.2 to 2.1 Å in RSF (Fig. 9). The Bi-O distances of the first and second coordination spheres for BiSG were calculated as about 2.1 and 2.3 Å, respectively; on the other hand, the Bi-O distances for α-Bi₂O₃ were 2.2 and 2.4 Å, respectively. The Bi-O distance of 2.1 Å in BiSG is in good agreement with the previously reported Bi⁵⁺-O distance in LiBi(5+)O₃(Kumada, Takahashi et al. 1996) and Bi₂(3+,5+)O₄(Kumada, Kinomura et al. 1995). Therefore, the existence of the Bi⁵⁺ state is also indicated from the Bi-O distance in BiSG, and the first coordination sphere corresponds to the Bi⁵⁺-O distance. The second coordination sphere of 2.3Å corresponds to the Bi³⁺-O distance(Ohkura, Fujimoto et al. 2007). Therefore, the EXAFS curve-fitting results also show that the mixed valence state of the Bi ions exists in BiSG as Bi³⁺ and Bi⁵⁺.

The Bi-O distance of the first coordination sphere in the A-series is slightly shifted to a longer range with increased Bi₂O₃ concentration in Table 4. The ratio of Bi³⁺ to Bi⁵⁺ increases with increased Bi₂O₃ concentration, and the change of the Bi³⁺ to Bi⁵⁺ ratio can also explain the non-linear increment of LMI. NaBiO₃ is a well-known material as a standard for pentavalent state Bi ions. The first coordination state distance of NaBiO₃ is longer than the expected value of the Bi⁵⁺-O distance. This valence state of Bi ions in NaBiO₃ is also the mixed state of Bi³⁺ and Bi⁵⁺. These phenomena are also supported by the peak separation data of XPS.

Samples	First coordination sphere			Second coordination sphere			R-
Sumples	N ₁	R_1	$\sigma_1^2(\text{\AA}^2)$	N ₂	R ₂	$\sigma_2^2(\text{\AA}^2)$	factor(%)
α -Bi ₂ O ₃	2.01	2.18	3.91E-03	0.74	2.40	4.07E-03	8.82
NaBiO ₃	7.15	2.13	3.89E-03	3.12	2.38	4.06E-03	0.89
A2	2.19	2.08	3.89E-03	1.11	2.32	4.06E-03	1.65
A3	2.05	2.08	3.89E-03	1.33	2.31	4.06E-03	2.50
A4	1.81	2.11	3.89E-03	1.05	2.31	4.06E-03	2.84
A5	1.86	2.13	3.89E-03	0.99	2.33	4.06E-03	3.04

Table 4. FEFF fitting results providing two coordination spheres. Fitting results of A-series (A2 \sim A5), α -Bi₂O₃, and NaBiO₃ are listed.

3.9 Discussion (local structure of luminescent center)

In the previous section, several physical phenomena were observed in BiSG, especially regarding the local structure of the distinctive luminescent center. Now we consider the structural configuration on the Bi luminescent center.

First, the roles of the Al_2O_3 additive can be understood by luminescent intensity measurement. Based on Fig. 5, the luminescent intensity of A4, which includes 2.3 mol% of Al_2O_3 , is three orders of magnitude larger than that of C1 without Al_2O_3 ; clearly, the Al_2O_3 additive remarkably increases the generation of the Bi luminescent center. Second, Al_2O_3 assists the Bi ions to enter the silica glass network because C1 has no glassy wetting. This tendency is also supported by the phase diagram of the Bi_2O_3 - Al_2O_3 -SiO₂ glass system, because the glassy phase is likely achieved at the Al_2O_3 -rich composition. Therefore, aluminum ions have two roles in BiSG:

- 1. They assist the configuration of the distinctive luminescent center of Bi ions with a coupling effect that denotes that an aluminum ion behaves like a "generator" of the luminescent center.
- 2. They increase compatibility with the silica network.

These aluminum ion roles imply that both the Bi and Al atoms should be close together in BiSG. Based on the above discussion, the image view between Bi and Al ions in BiSG is illustrated in Fig. 10(a). Peng et al.(Peng, Qiu et al. 2005) reported that Ta ions also work as a "generator." Although aluminum is not the only element that behaves as a generator, the aluminum ion accepts its important role in the Bi₂O₃-Al₂O₃-SiO₂ glass system.



Fig. 10. Image view on local structure of infrared Bi luminescent center in BiSG: (a) image view determined by PDG (phase diagram) and LMI, (b) by ²⁷Al-NMR, (c) by ESR, (d) by XPS, EXAFS, (e) local structure of infrared Bi luminescent center.

Next, the aluminum cordination state (ACS) should be close to the Bi ion in BiSG, as seen from the ²⁷Al-NMR results. Since the relation between ACS and the chemical shift in the ²⁷Al-NMR measurement has been well studied(Laussac, Enjalbert et al. 1983), ACS is determined by a chemical shift in comparison between the standard materials and the target samples. In the case of BiSG, ACS is dominated by the α -Al₂O₃ corundum structure at lower Bi₂O₃ concentration up to 0.5 mol%. Al ions with corundum structure are crucial to generate a distinctive luminescent center, and the Al coordination state located near Bi should be a 6-fold corundum structure. Based on the above discussion, the image view between Bi and Al is illustrated in Fig. 10(b).

Next, information on the valence states of the Bi ions in BiSG is given by ESR measurements. Of course, it's not only for Bi ions, but since no signal exists for the unpaired electrons in the whole BiSG, the valence states of Bi ions without unpaired electrons are Bi³⁺ or Bi⁵⁺. These results show Bi³⁺ or Bi⁵⁺ ions close to the 6-fold coordination state of the Al ions. Based on the above discussion, the image view between Bi and Al is illustrated in Fig. 10(c).

Three types of coordination states of Bi3+ exist, including 5-, 6-, and 8-fold; on the other hand, only 6-fold coordination exists for Bi⁵⁺(Shannon 1976). Since BiSG is an oxide material, it is estimated that the neighboring ions of the Bi ion are oxygen. Although the ionic radius of O²⁻ has few differences with the coordination number, variation exists between 1.35 and 1.42 Å. If the coordination number of O²⁻ is 4, Bi³⁺(5)-O²⁻(4), Bi³⁺(6)-O²⁻(4), and Bi⁵⁺(6)-O²⁻(4) are calculated to be 2.34, 2.41, and 2.12 Å, respectively(Shannon 1976). The Bi-O distances in typical crystals including Bi³⁺ or Bi⁵⁺, such as LiBiO₃(Kumada, Takahashi et al. 1996) or Bi₂O₄(Kumada, Kinomura et al. 1995), show that the Bi⁵⁺-O distance is 2.1 Å. This value agrees well with the 2.1 Å of the first coordination sphere for A4 and A5. But the Bi³⁺-O distance varies from 2.15 to 3.26 Å in α-Bi₂O₃(Harwig 1978) or Bi₂O₄(Kumada, Kinomura et al. 1995). Therefore, the EXAFS data show that the Bi⁵⁺ ionic state exists in BiSG, and this is also supported by the XPS data. The previously reported Bi3+ spectroscopic properties are quite different in luminescent and absorption spectra and lifetime. It is concluded that the Bi valence state of the Bi luminescent center is Bi⁵⁺, not Bi³⁺. Therefore, the luminescent center model of Bi5+ with 6-fold coordination is expected to be close to Al3+ with 6-fold coordination of the corundum structure (Fig. 10(d)). Since the neighboring atom is oxygen, the local structure of the distinctive bismuth luminescent center is expected (Fig. 10(e)).

4. Applications

After the discovery of a new infrared luminescent bismuth center, several research groups started to study its applications, such as optical amplification(Fujimoto & Nakatsuka 2003; Seo, Fujimoto et al. 2006; Seo, Fujimoto et al. 2006; Ren, Wu et al. 2007; Ren, Dong et al. 2007; Ren, Qiao et al. 2007; Seo, Lim et al. 2007), waveguide inscription(Psaila, Thomson et al. 2006), or laser oscillation(Dianov, Dvoyrin et al. 2005; Dianov, Shubin et al. 2007; Razdobreev, Bigot et al. 2007; Rulkov, Ferin et al. 2007; Truong, Bigot et al. 2008) using Bi luminescent materials.

With respects to device applications, optical fibers with Bi luminescent center in the core material are very curious. Optical amplification around 1.3 μ m with Bi-doped multi-component glass fiber was achieved by Seo et al.(Seo, Fujimoto et al. 2006), and this is useful for metro area network optical amplifiers. Laser oscillation with Bi-doped optical fiber was firstly demonstrated by Dianov's group in 2005(Dianov, Dvoyrin et al. 2005), then the possibility of Bi-doped fiber is actively developed, now the oscillation power has achieved

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at 15 W at 1160 nm(Bufetov & Dianov 2009). It is known that the 570 – 590 nm band is very promising for ophthalmology and dermatology applications, thus the second harmonic of Bi fiber laser can be used in medical use. And the broad luminescence in near infrared region is also useful for a light source of optical coherence tomography.

5. Conclusions

In this chapter, we introduce the basic properties of BiSG and the analyzed local structure of Bi luminescent center. Several instrumental analyses, such as spectroscopic properties (SPCT), LMI, NMR, XRD, ESR, XPS, and EXAFS were advanced on a Bi₂O₃-Al₂O₃-SiO₂ glass system. The roles and the structure of the Al ions and the valence state of the luminescent Bi ions were examined.

The following are the roles and the structure of the Al ions: 1) to assist the configuration of the distinctive luminescent center of Bi ions with a coupling effect, which means that the aluminum ion behaves like a "generator" of the luminescent center; 2) to increase compatibility with the silica network; 3) to be a 6-fold corundum structure. The valence state exmination of Bi ions in BiSG reveals the following: 1) Bi³⁺ or Bi⁵⁺; 2) a mixed state of Bi³⁺ and Bi⁵⁺; 3) Bi⁵⁺ for the distinctive Bi luminescent center. Therefore, the distinctive bismuth luminescent center model was investigated with a 6-fold coordination state of Bi⁵⁺ that is combined with the 6-fold corundum structure of Al³⁺ through an oxygen ion (Fig. 10(e)). These results will bridge to verify the energy diagram and the mechanism of Bi luminescent center.

In the last place, this new infrared luminescent material, Bi-doped silica glass, which attains sensational progress in the past decade will continue to give us curious possibilities in the field of the optical science.

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Invention of the solid-state laser has initiated the beginning of the laser era. Performance of solid-state lasers improved amazingly during five decades. Nowadays, solid-state lasers remain one of the most rapidly developing branches of laser science and become an increasingly important tool for modern technology. This book represents a selection of chapters exhibiting various investigation directions in the field of solid-state lasers and the cutting edge of related applications. The materials are contributed by leading researchers and each chapter represents a comprehensive study reflecting advances in modern laser physics. Considered topics are intended to meet the needs of both specialists in laser system design and those who use laser techniques in fundamental science and applied research. This book is the result of efforts of experts from different countries. I would like to acknowledge the authors for their contribution to the book. I also wish to acknowledge Vedran Kordic for indispensable technical assistance in the book preparation and publishing.

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