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Regulating the Bi NIR luminescence behaviours in fluorine and nitrogen co-doped germanate glasses

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Bi-Doped nitridated germanate glasses covering the whole NIR region from 800–1600 nm have attracted considerable attention due to their potential application in tunable fiber lasers and optical amplifiers. However, it remains challenging to regulate the luminescence behaviour of Bi because of the coexistence of multi-centers in the glasses. Here, we successfully regulated multi Bi NIR centers to obtain a flatter and ultrabroadband emission in nitridated germanate glasses. By varying the alkaline earth metal fluorides from MgF₂ to BaF₂, the glass structures were gradually depolymerized. A looser glass network, on one hand, promoted Bi in lower valences to be oxidized into Bi²⁺ and Bi³⁺, thus reducing the NIR emission intensity at ~1150 and ~1260 nm; on the other hand, it also enhanced the energy transfer from Bi⁺ to Bi⁰, which resulted in a relatively flatter emission band. In addition, the depolymerised glass structure could facilitate more nitride bonds, which are favorable for dispersing and stabilizing the new Bi NIR active centers related to germanate for ~930 and ~1490 nm emission. This investigation offers an efficient way to manipulate the multi-center luminescence behaviour of Bi in nitridated germanate glasses, and is beneficial to understand the mechanism of Bi NIR emission centers in glasses.

1 Introduction

The arrival of the 5th generation of communication technology (5G) is often accompanied by the explosive growth of information and data, which require bigger information transmission capacity for the current optical communication system. One possible way to improve the performance of the optical communication system is to broaden the data transmission channels, thus increasing the information transmission capacity. Laser materials operating in the broadband NIR spectral region (1200–1600 nm) are the major components for developing modern communication systems with unprecedented performance.¹⁻⁴ Due to the excellent fluorescence characteristics, rare-earth-doped materials, as a type of efficient gain medium, particularly near-infrared (NIR) optical amplifiers and tunable lasers, have attracted widespread attention. Unfortunately, the f-f electronic transition, which is shielded by the

In the past decades, Bi-doped glasses and fibers have been confirmed to exhibit intriguing broad NIR photoemission from 1000 to 1600 nm, which made them a promising medium for optical amplifiers and NIR lasers. $^{10-21}$ Although a significant progress has now been made in improving the performance of Bi-doped glasses and fibers, the emission intensity of the widely used C and L bands (1530–1625 nm) is relatively weak. In recent years, Cao $et\ al.$ realized a super-broad emission band shape, which extends over the whole NIR spectral region (850–1600 nm) via the topochemical reduction of bismuth in germanate glasses co-doped with AlN, Si $_3$ N $_4$ or SiC. $^{20,22-24}$ The topochemical reduction in germanate glasses not only enhances the near-infrared luminescence of bismuth but also generates new NIR emission centers located at \sim 930 and \sim 1500 nm, which are related to germanium. Due to the coexistence of

outer-layer electrons, is almost unaffected by glass composition, making it difficult to modulate the emission bandwidth and peak position. As a result, the gain bandwidth of traditional rare-earth-doped materials has always been quite narrow (not exceeding 100 nm), which cannot make full use of the ultra-low-attenuation telecommunication windows of silica glass fibers (1260–1675 nm).^{5–9} Therefore, it is urgent to explore new photonic materials with an ultra-broadband NIR emission and long lifetime as a gain medium for potential applications in broadband optical amplifiers and tunable lasers.

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multiple Bi NIR activate centers (such as Bi^+ , Bi^0 and Bi related to germanium) and their super sensitivity to the changes in the crystal field, it remains challenging to regulate the Bi NIR luminescence behaviour. $^{21,25-28}$

At present, Bi-doped fibers are mainly based on oxide glasses, which show superior chemical and optical stability; however, simultaneously, they suffer from the large phonon energy (\sim 1200 cm⁻¹ for silica glass). Compared to oxide glasses, the phonon energy of a fluoride glass ($\sim 550 \text{ cm}^{-1}$ for ZBLAN glass) is relatively low, which helps to reduce the non-radiative transition and to enhance the luminescence, but its chemical stability is poor. Hence, oxyfluoride glasses combining the advantages of both oxide and fluoride glasses, may be one of the best candidates for optical materials. Owing to this reason, oxyfluoride glasses have been confirmed to be promising rare-earth-doped optical materials in recent years.^{29–31} In 2012, Xu et al. reported that the substitution of magnesium fluoride for magnesium oxide in Bi-doped silicate glasses could significantly increase the intensity and lifetime of the NIR luminescence.32 In 2020, Chen et al. manifested that the partial substitution of barium fluoride for barium oxide in Bi-doped nitridated germanate glasses could facilitate the formation of nitride bonds to induce NIR luminescence in the glasses.³³ However, to the best of our knowledge, there are a few reports related to a systematic study on different fluorides of alkali earth metals on the regulation of multi-center luminescence behaviour of Bi in nitridated germanate glasses.

In this study, taking topochemically reduced multicomponent germanate glasses as models, a series of nitridated germanate glasses co-doped with a low Bi content and different fluorides of alkaline earth metals were investigated. The results here indicated that the Bi NIR luminescence behaviour can be manipulated by varying the alkaline earth metal ions, and flat and tunable NIR emission spectra spanning from 850 nm to 1600 nm were obtained by adjusting the glass structures. We believe that the present work is beneficial to promoting the design of Bi-doped germanate glass fibres with a tunable and ultra-broadband NIR luminescence.

2 Experimental section

2.1 Synthesis

Glass composition with molar compositions of $76\text{GeO}_2-9\text{Al}_2\text{O}_3-x\text{CaF}_2-(12-x)\text{CaO}-3\text{AlN}-0.005\text{Bi}_2\text{O}_3$ (x=0,4,6,8, and 10) and $76\text{GeO}_2-9\text{Al}_2\text{O}_3-6\text{MF}_2-6\text{BaO}-3\text{AlN}-0.005\text{Bi}_2\text{O}_3$ (M=Mg,Ca,Sr, and Ba) were prepared by traditional melting and quenching methods. For convenience, the glass samples were denoted as xCF (x=0,4,6,8, and 10) and MF, CF, SF, BF, respectively. Analytical grade Al_2O_3 , MgF_2 , CaF_2 , SrF_2 , BaF_2 , CaCO_3 , BaCO_3 and high-purity GeO_2 (99.999%), AlN and Bi_2O_3 (99.99%) were used as the starting materials. Batches of raw materials (15 g) were mixed homogeneously in an agate mortar. Each batch of the mixture was melted in an alumina crucible at a temperature of 1520 °C in air for 30 min. Afterwards, the melt was cast onto a stainless-steel plate, and finally pressed by another stainless-steel plate to speed up the cooling process and avoid

crystallization. Finally, the glasses were cut and polished for optical analysis.

2.2 Characterization

Optical absorption spectra of the samples were recorded using a PerkinElmer Lambda 900 UV/Vis spectrophotometer over the spectral range of 300–1000 nm. Static excitation and emission spectra, dynamic emission spectra excited at 460 nm, and luminescence lifetime were obtained using an Edinburgh FLS920 spectrofluorometer equipped with a continuous wave 450 W Xe lamp and a microsecond flashlamp (μ F900) as excitation sources. Fourier transform infrared (FT-IR) spectra were measured in the transmission mode using a Bruker Vector 33 spectrometer by dispersing glass powders into KBr pellets. Electron paramagnetic resonance (EPR) spectra were collected using an X-band spectrometer (Bruker E500). All measurements were performed at room temperature.

3 Results and discussion

In order to verify that the introduction of an appropriate amount of fluorine in the glasses is beneficial to improve the NIR luminescence property of Bi, a series of Bi-doped germanate glasses containing different contents of fluorine were investigated. Fig. 1 illustrates the emission spectra of 76GeO₂-9Al₂O₃-xCaF₂-(12 - x)CaO-3AlN-0.005Bi₂O₃ (x = 0, 4, 6, 8, and 10). The intensities of the emission bands at ~920 nm and ~1484 nm first increase, and then, decease with the further increase in the fluorine content. However, the intensity of the emission band at ~1150 nm kept decreasing continuously. As shown in the inset of Fig. 1, the intensities of the emission bands at \sim 920 and \sim 1484 nm reached the appropriate state at 4 mol%. The intensity of the emission band at \sim 1500 nm decreases continuously, which indicates that fluorination can effectively manipulate the Bi NIR emission bands, and enhance the emission intensity, particularly for the emission bands at \sim 920 and \sim 1484 nm.

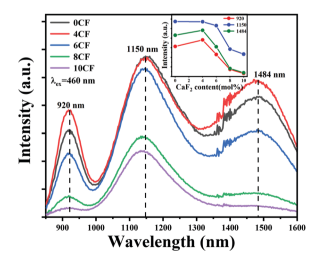


Fig. 1 The emission spectra ($\lambda_{\rm ex}$ = 460 nm) of the xCF (x = 0, 4, 6, 8, and 10) glass samples. The inset is the integral intensity of the ~920, ~1150, and ~1484 nm emission bands as a function of the content of fluoride.

When we introduced different fluorides of alkaline earth metals to Bi-doped nitridated germanate glasses, the luminescence spectral configuration of the glasses could be tuned. As presented in Fig. 2(a), under an excitation at 460 nm, an ultrabroadband NIR luminescence covering 850-1600 nm with three predominant peaks at ~ 924 , ~ 1150 and ~ 1496 nm related to the Bi NIR centers is observed in all samples. According to previous reports, the emission bands at \sim 924 and 1496 nm can be attributed to the new Bi-activated centers generated by the addition of nitride in germanate glassses, 22,23,34 while the emission band at ~ 1150 nm originated from the typical transition ${}^{3}P_{1} \rightarrow {}^{3}P_{0}$ of Bi⁺. ^{20,35} Moreover, as shown in Fig. 2(a), the integral intensity of the ~1150 nm emission band showed a dramatical decreasing trend as the radius of alkaline earth metal ion increased, while the integral intensity of the \sim 924 and \sim 1496 nm emission bands got slightly enhanced. Moreover, different from the \sim 1150 nm emission band, the emission bands located at \sim 924 and \sim 1496 nm demonstrate an increasing trend, and the shapes of their emission bands are similar, as indicated by the corresponding normalized emission spectra (to the intensity at ~ 1150 nm) in Fig. 2(b). The above phenomenon indicates that the Bi-activated NIR centers responsible for ~924, ~1150, and ~1496 nm responded differently to the change of alkaline earth metal fluorides.

More importantly, the FWHM of the ~1150 nm emission band increased from 187 to 443 nm by varying the alkaline earth metals from Mg to Ba. Although the overall emission intensity at ~1150 nm shows a weakening trend, a new emission band appeared at ~1260 nm, which is due to the typical transition $^{2}D_{3/2} \rightarrow ^{4}S_{3/2}$ of Bi⁰, and emerged gradually in the SF and BF glass samples (Fig. 2(b)).18 Owing to the strong overlap between the \sim 1150 and \sim 1260 nm emission bands, it was hard to detect the \sim 1260 nm band in the MF and CF glass samples. A Gauss fitting of the emission spectra of the four samples was conducted towards the emission spectra of the glasses with different fluorides of alkaline earth metals. As shown in Fig. 2(c), there are four emission bands in each sample. Although the overall emission intensity at \sim 1150 nm decreased, the ratio between the emission intensity of \sim 1260 nm to that of \sim 1150 nm increased with the increase in the radius of alkaline earth metal ions (Fig. 2(d)). This result was slightly different from the phenomenon reported by a previous investigation, in which only a redshift and an intensity decrease occurred as the radius of alkaline earth metal ions increased.36-38 Here, a tentative reason to explain this phenomenon is that the energy transfer between Bi⁺ and Bi⁰ occurred.

The absorption spectra of the glasses are shown in Fig. 3. These samples all exhibited three absorption peaks centered at

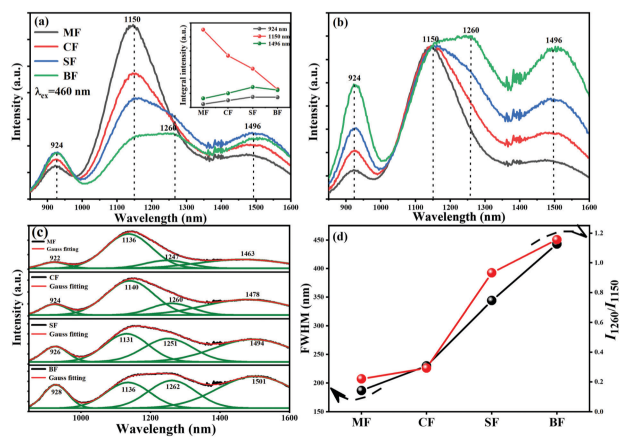


Fig. 2 (a) Emission spectra (λ_{ex} = 460 nm) of the MF, CF, SF, BF glass samples. The inset is the integral intensity of the ~924, ~1150, ~1260 nm and ~1496 nm emission bands as a function of alkaline earth metal fluorides. (b) Corresponding emission spectra normalized to the emission peak at ~1150 nm. (c) Comparison between the emission spectra of sample MF, CF, SF, BF by Gauss fitting. (d) Dependence of the FWHM of the ~1150 nm emission band and the ratio of I_{1260}/I_{1150} on the alkaline earth metal fluorides.

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Fig. 3 The optical absorption spectra of MF, CF, SF, BF glass samples.

 $\sim\!460,\,\sim\!574$ and $\sim\!710$ nm, respectively. It is widely accepted that the $\sim\!460$ nm absorption band is attributed to the typical absorption of Bi⁰, while the $\sim\!710$ nm band is from Bi^{+,26,27,37} However, as compared to the $\sim\!460$ and 710 nm bands, the $\sim\!574$ nm band presented a flatter and wider shape, which may be caused by the new Bi-activated NIR centers. By varying alkaline earth metal fluorides from MgF₂ to BaF₂, all the Bi-related absorption intensities gradually decreased.

To obtain more information on the Bi-related centers, the excitation spectrum of different emission bands of the BF glass is collected, as shown in Fig. 4. It can be seen that the four NIR emission bands could be well excited by blue light, which shows a great potential as a blue LED-excited broadband NIR light source. The four excitation spectra, however, are different in some details. When monitored at 930 nm, five excitation bands at \sim 338, \sim 394, \sim 456, \sim 574, \sim 672 and \sim 800 nm are observed. These complicated excitation features are different

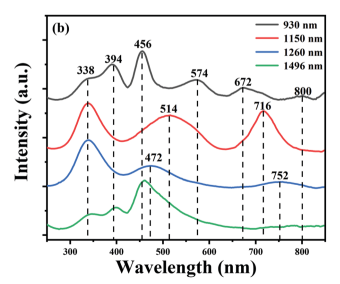


Fig. 4 Excitation spectra at 930, 1150, 1260 and 1490 nm for the BF sample.

from the excitation bands located at \sim 338, \sim 394, and \sim 456 nm for the 1490 nm emission band. Therefore, we inferred that the \sim 924 and \sim 1496 nm emission bands may originate from different Bi-activated NIR centers. While monitoring at \sim 1150 nm, the excitation spectra consist of three strong bands locating at \sim 338, \sim 514 and \sim 716 nm, respectively. The \sim 1260 nm emission band presents a strong absorption band at \sim 338 nm and other broad excitation bands at \sim 472 and \sim 752 nm. The variation tendency observed in the excitation spectra can well explain the different behaviour of the multiple Bi-activated centers in some degree.

The fluorescence lifetime is another important parameter to characterize the fluorescence performance of Bi-doped glasses. The decay curves of multi-center NIR emission at 930, 1150, 1260 and 1490 nm under the excitation of 460 nm are presented in Fig. 5(a)–(d). All the decay curves followed a double exponential decaying law, as described by eqn (1):

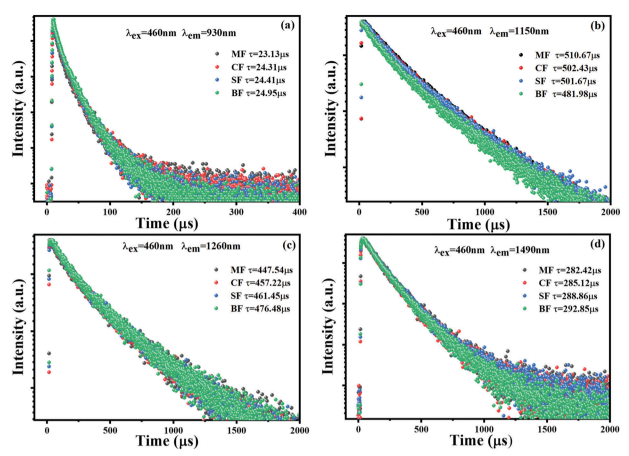
$$I(t) = A_1 \exp(-t/\tau_1) + A_2 \exp(-t/\tau_2)$$
 (1)

where I(t) is the intensity at time t; τ_1 and τ_2 are short and long decaying components, respectively; parameters A_1 and A_2 are fitting constants. Fitting each curve to the equation produced τ_1 and τ_2 , and the correlation coefficients were always between 99.50% and 99.94%. As shown in Fig. 5, the obtained mean lifetimes for 930, 1150, 1260, and 1490 nm shared different responses with varying alkaline earth metals from Mg to Ba. As listed in Table 1, the mean lifetime for ~ 930 and ~ 1490 nm was extended slightly, and that for ~1260 nm significantly increased. However, the opposite trend was observed in the lifetimes of the emission at 1150 nm. The opposite variation trends between ~1150 and ~1260 nm gave further evidence that the energy has transferred from Bi⁺ to Bi⁰. Moreover, compared to oxide glasses, Bi-doped oxyfluorogermanate glass with a low phono energy could reduce the probability of the nonradiative relaxation of Bi-activated NIR centers, thus obtaining a relatively long lifetime. 22,39

In consideration of the above-mentioned results, here, proper energy level diagrams for Bi $^{+}$ and Bi 0 were depicted on the basis of previous studies to explain the energy transfer mechanism. 18,26 As shown in Fig. 6, both Bi $^{+}$ and Bi 0 can be excited by 460 nm light from the ground state to the excited levels of $^{1}D_{2}$ and $^{2}P_{1/2}$, respectively, and the active Bi $^{+}$ then relaxes non-radiatively to the first excited level $^{3}P_{1}$. The energy transfer process can be described as $^{3}P_{1}(\text{Bi}^{+}) + ^{4}S_{3/2}(\text{Bi}^{0}) \rightarrow \, ^{3}P_{0}(\text{Bi}^{+}) + \, ^{2}D_{3/2}(\text{Bi}^{0})$, which helps to emit a 1260 nm photon. In order to quantify the energy transfer, we calculated the efficiency in CF, SF, and BF glasses compared to the MF sample by the following eqn (2):

$$\mu_{\text{Bi}^+ \to \text{Bi}^0} = 1 - \frac{\tau_{1150}}{\tau_{1150_{\text{MF}}}} \tag{2}$$

where $\mu_{\rm Bi^+ \to Bi^0}$ is the energy transfer efficiency from Bi⁺ to Bi⁰; τ_{1150} is the mean lifetime of Bi⁺ at the 1150 nm emission in CF, SF and BF glasses; $\tau_{1150_{\rm MF}}$ is to the mean lifetime of Bi⁺ at the 1150 nm emission in the MF glass. As listed in Table 1, the energy transfer efficiency increased from 1.61% to 5.62% with varying CF to BF glasses compared to MF glass.



Decaying curves of (a) 930, (b) 1150, (c) 1260, and (d) 1490 nm emission of MF, CF, SF, and BF glass samples (λ_{ex} = 460 nm).

Table 1 The mean lifetime of Bi at 930, 1150, 1260, and 1490 nm, the energy transfer efficiency from Bi⁺ to Bi⁰ in CF, SF, and BF glasses as compared to MF glass, and the values of the optical basicity in the glass

Glass samples	τ ₉₃₀ (μs)	$\begin{array}{l} \tau_{1150} \left(\mu s \right) \\ Bi^{^{+}} \end{array}$	$\begin{array}{l} \tau_{1260} \left(\mu s \right) \\ Bi^0 \end{array}$	τ ₁₄₉₀ (μs)	$\begin{array}{c} \mu_{\mathrm{Bi}^{+}\rightarrow\mathrm{Bi}^{0}} \\ \left(\%\right) \end{array}$	Optical basicity $(\Lambda_{\rm th})$
MF	23.13	510.67	447.54	282.42	_	0.601
CF	24.31	502.43	457.22	285.12	1.61	0.606
SF	24.41	501.67	461.45	288.86	1.76	0.609
BF	24.95	481.98	476.48	292.85	5.62	0.611

Owing to the co-existence of multiple Bi-activated NIR centers in the glasses, the ultrabroadband NIR emission in Bi-doped germanate glasses prepared through topochemical reduction could be tuned widely to span the NIR region by adjusting the excitation wavelength from 420 to 700 nm, as presented in Fig. 7(a). Compared to RE-doped or RE-Bi co-doped glasses, the NIR emission performance of Bi-doped glasses was more flexible and efficient in terms of the emission bandwidth and peaks. 40,41 To further reveal the relationship between the emission and excitation behaviour in detail, the emission-excitation map from the BF glass sample is denoted in Fig. 7(b). Clearly, there are multiple NIR centers co-existing in the glass and Bi-activated centers responded differently to the excitation wavelength. For NIR emission centers at \sim 1100 nm, two wide absorption bands

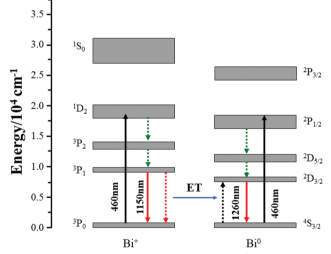


Fig. 6 Simplified energy level diagram of Bi⁺ and Bi⁰, and the possible energy transfer between Bi⁺ and Bi⁰.

ranging from 420 to 600 nm and 660 to 720 nm are observed, which indicates that the emission band around ~1100 nm could be excited by visible light. For the NIR emissions centered at \sim 930 and \sim 1500 nm, there was only a narrow absorption band between 430 and 470 nm. By increasing the excitation wavelength

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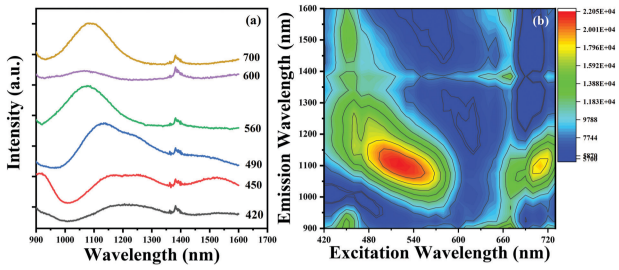


Fig. 7 (a) The NIR spectra evolution of the BF glass sample. (b) The contour plot of the emission-excitation map from the BF glass sample.

from 420 to 720 nm, the emission band can cover the 900-1600 nm region at the beginning, but set to ~ 1100 nm at the end. These results provided a flexible pumping scheme of Bi-doped glasses for the potential application as NIR materials.

In order to understand why the above changes occurred, the concept of optical basicity, which has an important influence on the valence state and luminescence behaviour of Bi, was employed to provide more clues.^{37,38} The values of the Duffy's optical basicity (Λ_{th}) of glasses can be calculated by the eqn (3):

$$\Lambda_{\text{th}} = (X_1 \Lambda_1 + X_2 \Lambda_2 + \dots + X_n \Lambda_n) \tag{3}$$

where Λ_n is the optical basicity values of each composition of glass, and X_n is the equivalent fraction of the oxides and the fluorides that composition n contributes. Values of the optical basicity Λ_n can be decided by referring to ref. 42. As presented in Table 1, the optical basicity values of glasses increased from 0.600 to 0.611 for the samples varying from MF to BF. It is widely accepted that the low optical basicity favors Bi ions with a low valence state, while higher ones exist easily in glasses with a higher optical basicity. 32,37 Increasing the optical basicity values of the glass will unavoidably change the local environment around Bi-activated centers, thus triggering the chain reaction of Bi clusters or metal particles: Bi⁰ \rightarrow Bi⁺ \rightarrow Bi²⁺ \rightarrow Bi³⁺, resulting in a diversity of luminescence behaviours.

To the best of our knowledge, the Bi-activated centers are highly sensitive to the variation of the local crystal field and glass structures around them. 43-45 To reveal the deeper mechanism behind what we have discussed above, the structural analysis by FTIR spectroscopy and EPR was performed to offer more detailed information concerning the relationship between the microstructure of the glasses and the optical properties of Bi-activated NIR centers. FTIR spectra are illustrated in Fig. 8. Two absorption bands around 578 cm⁻¹ and 862 cm⁻¹ are observed in the spectra, which are assigned to the symmetric stretching vibrations of Ge-O-Ge and the asymmetric stretching vibrations of Ge-O-Ge in the GeO₄ tetrahedra.^{24,37,46} As the radius of the alkaline earth-

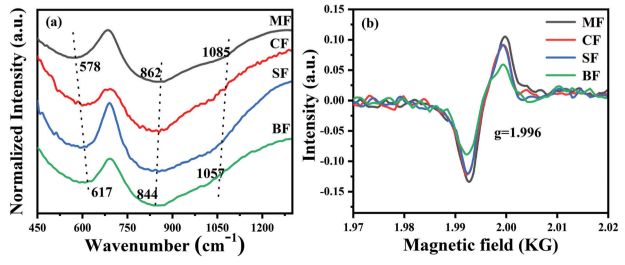
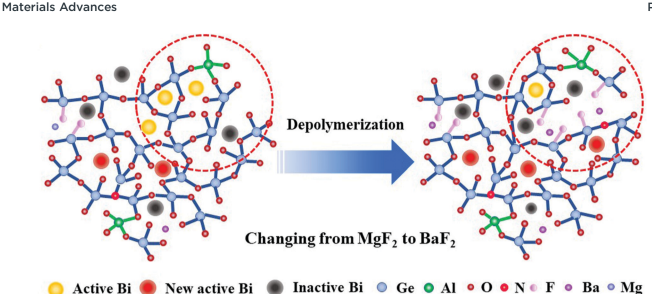


Fig. 8 (a) Normalized FTIR spectra and (b) EPR spectra of MF, CF, SF, BF glass samples.



Evolution of the microscopic structure of the glasses MF to BF.

metal ions increases, two remarkable changes are observed in the FTIR spectra. First, the former absorption peak located at 578 cm⁻¹ ultimately shifted to 617 cm⁻¹, indicating that the sixunit GeO₄ rings are broken up into smaller ones, and eventually, three-unit GeO₄ rings. Second, the latter absorption peaks blueshift from 862 cm⁻¹ to 844 cm⁻¹ and from 1085 cm⁻¹ to 1057 cm⁻¹, which are related to the conversion of the GeO₄ tetrahedra from Q3 (tetrahedral with one non-bridged oxygen) to Q² (tetrahedral with two non-bridge oxygens), Ge-O⁻ bands, and finally forming GeO₆ octahedron. ⁴⁷⁻⁴⁹ To sum up, the phenomenon observed above implies that more and more non-bridging oxygens are generated in the glasses, which therefore resulted in the depolymerisation of the glass network. The EPR technique is adopted to offer more information about the glass structure. As presented in Fig. 8(b), an obvious signal at g = 1.996 appeared in the four samples, which is attributed to the oxygen vacancy. Moreover, the signal intensity decreases continuously with the increase in the radius of the alkaline earth metal ions, indicating that more and more oxygen were involved into the glass structures, thus causing the Bi-activated NIR centers to be oxidized into non-NIR emission centers. This fact shows that the NIR luminescence behaviour of Bi could be regulated by modulating the overall glass structure.

Based on the above analysis, the structural evolution of the glasses from MF to BF is proposed in the schematic diagram in Fig. 9. As the ionic field strength decreases from Mg²⁺ to Ba²⁺, more and more F⁻ replace O²⁻ with varying alkaline earth metal fluorides from MgF₂ to BaF₂, which results in an increase in the content of non-bridging oxygen. As a result, the glass networks are gradually depolymerized. On one hand, it creates a negatively charged environment, which requires more positive charges to neutralize the excess negative charges, and the looser glass structures can easily expose the Bi NIR emission centers within the glasses to the oxygen atmosphere.21 The above two reasons drive the oxidation of Bi in low valence into higher valence, which are not active NIR centers. 50-52 Simultaneously, the depolymerized glass network promotes interactions between Bi⁺ and Bi⁰. As a result, the energy transfer between Bi⁺ and Bi⁰ is enhanced, which is why we could observe the increased energy transfer efficiency from Bi⁺ to Bi⁰ in Table 1. On the other hand, the depolymerized glass network may facilitate the formation of more nitride bonds connecting with Ge to the glass network around Bi, thus generating more new Bi NIR emission centers related to germanate by nitridation.³³ As a consequence, the emission intensity at \sim 924 and \sim 1496 nm improved. This result indicates that the multi-center Bi NIR luminescence behaviour can be regulated differently by adjusting the glass structures, and provides a better understanding about the mechanism of the Bi NIR emission.

4 Conclusions

In summary, through adjusting the glass structure, we successfully manipulated the multi-center Bi NIR luminescence behaviours in fluorine- and nitrogen-doped germanate glasses. A flatter emission spectrum spanning from 850 to 1600 nm with predominant emission bands located at ~924, ~1150, ~1260 and ~1496 nm was realized. By increasing the radius of alkaline earth metal ions, the glass structures were gradually depolymerized. The looser glass structures, on one hand, promoted the oxidization of Bi⁰ and Bi⁺ into inactive centers, thus reducing the emission intensity at \sim 1150 nm. On the other hand, it enhanced the energy transfer from Bi⁺ to Bi⁰, which was conducive to enhance the emission intensity at \sim 1260 nm. Moreover, the depolymerized glass network may facilitate the formation of more nitride bonds connecting to Ge to the glass network around Bi, thus generating new Bi NIR emission centers related to germanate by nitridation. As a result, the emission intensity and lifetime of ~924 and ~1496 nm emission bands got a certain degree of improvement. The study presented here is expected to promote our understanding to the mechanism of the Bi NIR emission behaviours in glass.

Conflicts of interest

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There are no conflicts to declare.

Acknowledgements

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