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ORIGINAL ARTICLE

Mesoscale engineering of photonic glass for tunable luminescence

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Control of the optical behavior of active materials through manipulation of their microstructure has led to the development of high-performance photonic devices with enhanced integration density, improved quantum efficiencies and controllable color output. However, the achievement of robust light-harvesting materials with tunable, broadband and flattened emission remains a long-standing goal owing to the limited inhomogeneous broadening in ordinary hosts. Here, we describe an effective strategy for the management of photon emission by manipulating the mesoscale heterogeneities in optically active materials. Importantly, this unique approach enables control of dopant–dopant and dopant–host interactions on the extended mesoscale. This allows the generation of intriguing optical phenomena such as a high activation ratio of the dopant (close to 100%), dramatically inhomogeneous broadening (up to 480 nm), notable emission enhancement and, moreover, simultaneously extension of the emission bandwidth and flattening of the spectral shape in glass and fiber. Our results highlight that the findings connect the understanding of and manipulation in the mesoscale realm to functional behavior on the macroscale, and the approach to manage the dopants based on mesoscale engineering may provide new opportunities for the construction of a robust fiber light source.

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INTRODUCTION

Multicomponent glass fibers are considered the building blocks of the next-generation of fiber photonics because they hold promise for the realization of a compact light source in a wide spectral range, which can be used in advanced applications, such as high-capacity tele-communication and remote sensing and in the defense industry.^{1–3} The ultimate challenge in achieving these operational modes is to develop a glass fiber material capable of emitting in a wide wavelength range. A conventional approach, tuning the glass composition, can achieve emission bandwidths up to 100 nm, but extending the bandwidth remains a challenging task.^{3,4} Another effective strategy for extending the luminescence spectral range is based on a multicorefiber design, in which cores, with a variety of rare-earth dopants, are physically arranged into a single fiber.^{5,6} Unfortunately, generating broadband flatline-shape emission has been achieved with limited success because of the discrete energy levels in rare-earth ions.⁷

From the traditional point of view, the structural origin of the spectral broadening is the site-to-site variation of dopants in a specific

host matrix.^{8,9} The corresponding structural ordering in homogeneous glasses is strictly limited to a short-range order of ~ 5 Å.^{10,11} In contrast, the dopant-host matrix and the dopant-dopant interactions are known to operate on a length scale of at least one order of magnitude larger than the short-range order in glass.^{12,13} It seems to be accepted now that supercooled liquids develop structural heterogeneities on cooling, which are preserved in the glass and can, for example, be the origin of an inhomogeneous response to thermal strain.14 Motivated by these significant clues, we recognize that functionality of doped photonic glasses manifests itself not only on the atomic or short-range order scales but also in the extended mesoscale. Here, we validate this hypothesis by providing a link between fundamental electron transition processes in a dopant and the mesoscale behavior of glass. We show that manipulating the mesoscale heterogeneities in a multiphase glass-ceramic fiber can result in significant inhomogeneous emission broadening beyond the 100 nm limit and, for the first time, in successful simultaneous extension of the emission bandwidth with a flattened spectral shape.

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EXPERIMENTAL PROCEDURES Materials

The transparent multicomponent glass was designed and prepared based on a standard phase diagram and the glass-forming ability of an LiTaO3-SiO2-Al2O3 system. The optimized composition of the host glass was Li₂O/Ta₂O₅/SiO₂/ Al₂O₃ = 25.53/21.53/35.29/17.65 (in mol.%). NiO powder was used for Ni²⁺-doping at the level of 0.15 mol.%. For a typical synthesis, ~40 g of the raw oxide materials (99.99% in purity) was homogeneously mixed and melted in a corundum crucible at 1600 °C for 90 min in air using an electric muffle furnace. The investigated glass system has a high propensity for devitrification. To avoid the crystallization of glass, we employed the melting-quenching method. The melt was poured onto a preheated stainless brass mold and pressed with another brass plate. This method can provide a rapid cooling rate. The thickness of the as-produced glass was 2 mm, and the size of the glass after cutting was 10 mm × 10 mm. The thermal properties of the prepared glass were measured by calorimetry and were used in the heat-treatment procedure to obtain the glass-ceramic composite. The glass-ceramic composites, with high optical transparency, were obtained by isothermal annealing at 750 °C, typically for 3 h. All samples were polished to optical quality.

Characterizations

The glass-transition and crystallization temperatures were measured by differential thermal analysis using an STA449C Jupiter (Netzsch, Bavaria, Germany) instrument in an argon atmosphere at a heating rate of 10 K min⁻¹. The X-ray diffraction was done using a D8 advance X-ray diffractometer (Bruker, Faellanden, Switzerland) equipped with Cu/K α radiation. Raman spectroscopy, using a Renishaw InVia spectrometer with a 532-nm laser source, was performed for structural studies. The transmission and absorption spectra were recorded using a Perkin-Elmer Lambda 900 UV/VIS/NIR spectrophotometer (Waltham, MA, USA). The excitation and photoluminescence spectra were measured using an FLS920 fluorescence spectrometer (Edinburgh Instrument Ltd., Edinburgh, UK) equipped with a xenon arc lamp (Xe900). The luminescence decay curves were recorded with a microsecond flash-lamp (µF900). The microstructure in both the glass and the glass-ceramic composite was characterized by transmission electron microscopy (TEM) and scanning

transmission electron microscopy (STEM). The images were acquired by a JEOL ARM200F electron microscope (JEOL Ltd., Tokyo, Japan) with a prober spherical aberration corrector. The TEM/STEM specimens were prepared by crushing the samples in an agate mortar and pestle with ethanol, followed by instilling a drop of suspension onto a copper grid covered with a lacy carbon film. The microstructure of the fiber was studied on a scanning electron microscope (ZEISS Merlin, Jena, Germany) equipped with an energy-dispersive X-ray spectrometer (EDX, Oxford Instruments, Oxford, UK). The X-ray absorption near-edge structure (XANES) spectra of Ni²⁺-doped samples at the Ni *K*-edge were collected at room temperature in the transmission mode using the 1W1B beam line (XAFS station) at Beijing Synchrotron Radiation Facility. The ring energy was operated at 1.85 GeV and 205 mA. The X-rays were monochromatized using a Si (111) double-crystal spectrometer, and the energy was calibrated by a Ni-foil reference. The Athena software, based on the IFEFFIT program, was used to analyze XANES data.

Theoretical simulations

Classical molecular dynamics simulations were performed to establish a structural model of the glass. To avoid atoms overlapping, the initial configurations were randomly generated with constraints between atom pairs. The time step for this simulation was set at 1 fs. The system was first equilibrated at 4000 K for 2000 time steps and then quenched to 293 K over 10 000 time steps. A final relaxation, for 2000 time steps, was carried out at room temperature (293 K). The atomic configurations of the final step were used for preparation of the multiphase glass. More details about the simulations are shown in the Supplementary Materials.

Fiber preparation

A 'molten-core' fiber-drawing method was used for fabrication of a high optical quality fiber.¹⁵ The Ni²⁺-doped bulk glass with high homogeneity was shaped into a cylindrical rod with a diameter of 3 mm and a length of 55 mm. To avoid the introduction of impurities into the fiber, the surface of the rod was carefully polished and washed by an acid (10% hydrofluoric acid). The glass rod was inserted into a high-purity silica tube (99.999%) with an inner



Figure 1 Mesoscale structure in the host $Li_2O-Ta_2O_5-Al_2O_3-SiO_2$ glass. (a) HAADF-STEM glass micrograph shows the presence of mesoscale heterogeneities. (b) Line profile of the contrast from **a**. (**c** and **d**) MD simulations of the chemical bond distribution. The red and blue sticks represent Ta–O and Si/Al–O bonds, respectively. (**e–h**) The average coordination numbers and the partial distributions calculated from the structure model. The average coordination numbers of Li and Ta cations are 5.6 and 5.7, and their bonds distances were calculated to be 2.04 and 2.25 Å, which match well those in the LiTaO_3 crystal (Ta–O: 2.07 Å; Li–O: 2.29 Å). The average coordination numbers of Si and Al are 4.02 and 4.62, and their bonds distances were calculated to be 1.62 and 1.8 Å. HAADF-STEM, high-angle annular dark-field scanning-transmission electron microscopy; MD, molecular dynamics.

The fiber-drawing process was performed at a temperature of 1950 °C, at which the core precursor is melted and the silica cladding is sufficiently soft. To effectively reduce the inter-diffusion between the core and the cladding, a fast drawing speed is required. In a typical fiber-drawing process, the preform was fed into a furnace from the top with a speed of 2 mm min⁻¹, and the fiber was pulled out from the bottom of the furnace with a speed of 1500 mm min⁻¹.

The fiber was optically activated by annealing at 800 °C for 20 h, which is higher than that used for the glass sample. This is because the degrees of structural heterogeneity in the glass and the fiber samples are different due to the different cooling rate during the fabrication process. In addition, in fiber sample, the core material was covered with SiO_2 cladding. In this case, the heat-transfer process during the heat treatment is also different from the case of the glass sample.

RESULTS AND DISCUSSION

Mesoscale structure in multicomponent host glass

Origin of heterogeneity. A prototypical multicomponent Li₂O-Ta₂O₅-Al₂O₃-SiO₂ (LTAS) glass was designed and prepared based on the standard phase diagram and glass-forming properties of the LiTaO₃-SiO₂-Al₂O₃ system (see Experimental procedures on details about the synthesis). This glass system was chosen because of its unique bonding features. Ta⁵⁺ ions with a high valence state, an intermediate field strength (F=1.2) and a high Ta–O bond energy (300 – 400 kJ mol⁻¹) show a high tendency to attract surrounding anions, resulting in local

compositional/structural fluctuations in the glassy matrix.¹⁶ Light alkali Li⁺ ions in the low valence state are highly mobile, have a low activation energy of diffusion (~67 kJ mol⁻¹) and facilitate structural rearrangements in the glass.¹⁶ The unique bonding features of Ta⁵⁺ and Li⁺ ions in this study favor the so-called amorphous phase separation, potentially leading to the formation of a Ta- and Si/Al-rich glass phase. The initial localized heterogeneities are expected to be quenched from the melt, starting from a liquid–liquid phase separation.

To validate the hypothesis of the local heterogeneities present in the glassy Li_2O - Ta_2O_5 - Al_2O_3 - SiO_2 system, the microstructure of LTAS was studied by high-angle annular dark-field STEM (Supplementary Figure S1). As shown in Figure 1a and b and Supplementary Figure S2, the STEM images show a typical pattern of regular heterogeneities on the mesoscale. The line profile, taken from Figure 1a, indicates periodic density fluctuations with an average characteristic length of ~ 1.8 nm (Figure 1b).

In the high-angle annular dark-field STEM mode, the image intensity (*I*) depends on the sample properties as follows: $I \propto \rho \times d \times Z^{1.5}$, where ρ is the material density, *d* is the TEM sample thickness and *Z* is the atomic number. The relative atomic masses of Ta, Al and Si are 180.9, 27.0 and 28.1, respectively. It is assumed that the brighter regions may correspond to areas containing more of the heavier elements, such as Ta, while the darker regions are richer in the lighter elements, such as Si and Al. The size of the heterogeneities is in



Figure 2 Structural characterization of the glass-ceramic composite. (a) XRD patterns of the glass and glass-ceramic composite. (b) Raman scattering spectra of the glass and glass-ceramic composite. (c) Bright-field TEM of the glass-ceramic composite. (d and e) Atomic-resolution HAADF-STEM images of, respectively, LiTaO₃ and LiAlSi₂O₆ nanoparticles present in the glass-ceramic composite. (f) Transmission spectra of the glass and glass-ceramic composite. The inset photographs show transparency of the glass (left) and glass-ceramic composite (right). HAADF-STEM, high-angle annular dark-field scanning-transmission electron microscopy; TEM, transmission electron microscopy; XRD, X-ray diffraction.

the range of nanometers. The STEM images can illustrate the idea of $[TaO_x]$ and $[Si/AlO_x]$ structural units demixing.

To shed more light on the origin of the observed heterogeneities, we performed molecular dynamics simulations to reveal the local coordination configuration and mesoscale ordering in the prototypical glass. Figure 1c and d show the distribution of Ta–O and Si/Al–O bonds, represented by the red and blue sticks, respectively. Local separation between $[TaO_x]$ and $[Si/AlO_x]$ units can be observed, as highlighted by the red circles (Figure 1c and d). Statistical analysis of the cation abundance ratio is presented in Supplementary Figure S3. In all these cases, the cation abundance ratio deviates from the average

value in the regions with the reduced scale, representing local chemical heterogeneities. The study of the coordination features of the glass network former shows that the local heterogeneities can mainly be assigned to the existence of two coordination units, $[TaO_6]$ and $[Si/AlO_4]$ (Figure 1e–h). The presence of heterogeneities in the glass was further tested for varying cooling rates, 3.7×10^{12} , 7.4×10^{12} and 3.7×10^{13} K s⁻¹, in molecular dynamics simulations (Supplementary Figure S4). In the limits of molecular dynamics cooling rates, a negligible influence of the cooling rate on the coordination configurations and the mesoscale architecture of the prototypical glass was observed.



Figure 3 Optical properties of the Ni²⁺-doped glass-ceramic composite. (a) Absorption spectra and a glass-ceramic composite photograph, in the inset, are shown. (b) NIR luminescence excited at $\lambda = 288$, 390 and 620 nm. (c) Decay curves of the NIR emission bands with the central emission wavelength at $\lambda = 1250$ and 1480 nm. (d) Photoluminescence excitation spectra of the NIR emission bands with the central wavelength at $\lambda = 1250$ and 1480 nm. (d) Photoluminescence excitation spectra of the NIR emission bands with the central wavelength at $\lambda = 1250$ and 1480. (e) XANES spectra of the Ni *K*-edge in the glass-ceramic composite, NiO and NiCr₂O₄. (f) Ni-Ni partial distribution functions of the glass-ceramic composite, NiO and NiCr₂O₄. NIR, near-infrared; XANES, X-ray absorption near-edge structure.

Multiphase glass-ceramic composite. The existence of mesoscale heterogeneities inside the glass can further be exploited in the development of multiphase glass-ceramic composite materials because the nucleation and crystal growth can be greatly affected by structural heterogeneities. On the one hand, the nucleation barriers are largely overcome by the structural heterogeneities where Ta- or Si/Al-rich regions form critical supersaturation domains.¹⁷ On the other hand, the local heterogeneities bear some similarities to the structure of the nucleating crystals, which can decrease the energy required to achieve the crystal growth.¹⁸ The LTAS glass was annealed for 3 h at 750 °C, which is slightly above its glass-transition temperature of $T_g = 700$ °C, but well below the crystallization temperature of 800 °C (Supplementary Figure S5). At this temperature, the crystallization of glass occurs at a moderate rate. As a result, the size of nanocrystals can be well controlled by tuning the heat-treatment parameters, and a glass-ceramic composite with a high transparency can be achieved. The X-ray powder diffraction pattern of the annealed glass (Figure 2a) shows fingerprint peaks that can be indexed to the LiTaO3 and LiAlSi2O6 crystalline phases. The Raman scattering spectrum (Figure 2b) shows intensive peaks that are consistent with the vibrational modes of crystalline LiTaO₃ and LiAlSi₂O₆.^{19,20} The band at \approx 464 cm⁻¹ (E mode) can be assigned to Ta–O bending, and the bands at ≈ 593 cm⁻¹ (A₁ mode) and ≈ 655 cm⁻¹ (E mode) originate from Ta–O stretching.¹⁹ The band at \approx 486 cm⁻¹ can be assigned to a stretching mode of T–O–T, where T represent Si or Al in the network. $^{\rm 20}$

In contrast, the as-prepared LTAS glass exhibits broad 'amorphous' halo peaks in both X-ray diffraction and Raman scattering data, indicating its glassy nature. Bright-field TEM micrographs (Figure 2c and Supplementary Figure S6) show the presence of dense crystalline nanoparticles, with average diameters of ~ 5 nm, embedded in the glassy LTAS matrix. Atomic-resolution high-angle annular dark-field STEM images, shown in Figure 2c–e, reveal a single-crystalline nature of the two typical particles, and the measured lattice fringes with *d*-spacings of 3.69 and 3.98 Å are in good agreement with the lattice spacing of the (012) plane of LiTaO₃ (3.69 Å, JCPDS #29-0836) and (102) of LiAlSi₂O₆ (3.98 Å, JCPDS #35-0797), respectively.

Importantly, the prepared glass-ceramic composite shows a high transparency in both the visible and the near-infrared (NIR) spectral regions (Figure 2f). The high transparency of the composite is primarily attributed to the uniformly dispersed nanocrystals within the glassy matrix.

The newly prepared nanocrystalline glass-ceramic composite, when optically activated by a proper dopant, Ni²⁺ ions in this study (see Materials in Experimental procedures), can become a novel multiphase photonic material, which is the main focus of the following sections.



Figure 4 Mesoscale structure-mediated tunable NIR emission of the Ni²⁺-doped glass-ceramic composite. (a) Schematic representation indicating the configurations of the (NiO₆)¹⁰⁻ octahedron and the crystal-field splitting of the 3*d* orbitals of Ni²⁺ in octahedral coordination. (b) Tanabe–Sugano diagram; the full rectangles indicate the crystal field of Ni²⁺ corresponding to the luminescence centers E₁ and E₂. (**c**-**e**) Fitted NIR luminescence spectra (the blue curves show the measured data, and the red curves represent the corresponding best fits). The bottom to top panels show the Ni²⁺-doped glass-ceramic composite activated by increasing annealing time from 5 to 10 and 20 h at 750 °C. The deconvoluted peaks show the emission centers E₁ (orange) and E₂ (green). NIR, near-infrared.

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 $N\dot{r}^{2+}$ doping. The presence of a rich mesoscale structure in the glassceramic composite may increase the chance of obtaining optically active dopants, and site-to-site variations offer unique opportunities for inhomogeneous spectral emission bandwidth broadening. To verify this suggestion, Ni²⁺ ions were chosen as the probing dopant, taking into account its fundamental role in the active photonics.^{21,22} The Ni²⁺ doping level was 0.15 mol.%, and the doped glass-ceramic composite was produced by the same thermal treatment as the host LTAS glass, that is, annealing at 750 °C and for 5 h.

Figure 3a shows the representative absorption spectrum of the Ni²⁺-doped glass-ceramic composite. Characteristic absorption peaks ${}^{3}A_{2}(F) \rightarrow {}^{3}T_{2}(F) \ (\lambda = 1270 \text{ nm}) \text{ and } {}^{3}A_{2}(F) \rightarrow {}^{3}T_{1}(F) \ (\lambda = 716 \text{ nm}),$ corresponding to electronic transitions in (NiO₆)10⁻ clusters, are observed.^{23,24} Importantly, we observed an ultra-broadband NIR luminescence with the highest full-width at half-maximum of ~400 nm in the glass-ceramic composite (Figure 3b). Interestingly, the shape of the steady-state emission is highly sensitive to the excitation wavelength (Figure 3b and Supplementary Figure S7). The examination of the decay process shows that the decay times cannot be simply fit to the exponential decay equation (Figure 3c). Moreover, the steady-state excitation measurements show a spectral shift when changing the detection wavelength (Figure 3d). All these results provide solid proof of the remarkable NIR emission broadening originating from the inhomogeneous glass-ceramic structure, in contrast to the reference glass-ceramic composite Ni2+-doped Ga2O3, for which only negligible inhomogeneous broadening is observed (Supplementary Figures S8 and S9).

The structure of the Ni^{2+} doped glass-ceramic composite was studied by XANES measurements of the active centers. As shown in Figure 3e, the XANES spectrum of the Ni dopant in the glass-ceramic composite resembles that of NiO, suggesting that the average first-shell coordination number of Ni²⁺ is approximately eight, which is consistent with the optical analysis presented above. The XANES spectra of the Ni–O shell in *R*-space show notable extension, strongly indicating the existence of site-to-site variation of Ni²⁺ in the glass-ceramic composite (Figure 3f and Supplementary Figure S10). Origin of luminescence. Another distinct advantage of the unique mesoscale structure in the multiphase Ni2+-doped glass-ceramic composite is that it can also facilitate effective management of the emission output. In a further set of experiments, we synthesized samples using variable annealing times of 0, 5, 10 and 20 h at a temperature T = 750 °C. The fully glassy sample shows negligible emission in the NIR spectral region (Supplementary Figure S11). By inducing growth of the nanocrystals in the glass, the emission not only shows remarkable intensity enhancement but also changes its spectral band shape (Figure 4c-e). The contribution of Ni²⁺ from the glassy phase to the radiative transitions is negligible. Distinctive bands of (NiO₆)10⁻ active units in LiTaO₃ and LiAlSi₂O₆ crystalline nanophases were obtained by deconvoluting the emission spectra, as indicated in Figure 4c-e. To further assign the observed emission bands, we performed a ligand-field theory analysis of the electronic transitions in (NiO₆)10⁻ clusters (Figure 4a and b). The electrostatic field produced by the six ligands, for an octahedrally coordinated unit, can be expressed by the potential V_{oct} according to equation (1):^{25,26}

$$V_{\text{oct}} = \frac{6Z_{Le}}{R} + \frac{35Z_{Le}}{4R^5} \left(x^4 + y^4 + z^4 - \frac{3}{4}r^4 \right) \tag{1}$$

where (Z_{Le}) is the charge of the ligand separated by the distance R from the cation, and x, y, z and r are the polar coordinates of the electrons. On the basis of this expression, the critical physical factor that determines the energy level configuration of $(NiO_6)^{10-}$ clusters, that is, the crystal-field splitting parameter Δ_0 , can be obtained according to equation (2):

$$\Lambda_{\rm o} = \frac{Q\langle r^4 \rangle}{R^5} \tag{2}$$

where Q is a constant. The theoretical analysis indicates that Δ_0 is inversely proportional to the fifth power of the distance between the central cation and the surrounding ligands. According to the crystalfield splitting illustration of the 3*d* orbitals of Ni²⁺ in the octahedral coordination state (Figure 4a), the large crystal-field splitting parameter Δ_0 directly leads to the enhancement of the energy gap between the ground state and excited states. Thus, it is believed that



Separation Distance

Figure 5 Mesoscale structure-mediated management of dopant-dopant interaction. Schematic illustration showing the dependence of the energy transfer process on the separation distance, r, between the two emission centers E_1 and E_2 . The line profile in the TEM image indicates the contrast from the dotted line.



Figure 6 Photonic glass fiber. (**a** and **b**) Photographs of the starting bulk glass $(1 \times 1 \times 5.5 \text{ cm}^3)$ and the drawn fibers. (**c**) SEM image of the splice joint between the commercial silica fiber (on the left of the joint) and the activated multiphase glass-ceramic composite gain fiber (on the right of the joint). The inset shows the corresponding optical micrograph. (**d**) Optical cross-section micrograph of the activated fiber. The core diameter is $10 \pm 1 \,\mu$ m, and the cladding diameter is $130 \pm 2 \,\mu$ m. (**e** and **f**) Raman mapping of the fingerprint peaks of LiTaO₃ (593 cm⁻¹) and LiAlSi₂O₆ (486 cm⁻¹) units in the core region of the activated fiber. (**b** and **i**) EDX mappings of Ta and AI elements in the core region of the activated fiber. EDX, energy-dispersive X-ray spectrometer; SEM, scanning electron microscope.

 $(NiO_6)10^-$ clusters will emit in the high-energy region of the NIR spectral range in the LiAlSi₂O₆ matrix with a tight framework (indicated as E₁ in Figure 4) and in the low-energy part of the NIR region in the LiTaO₃ host matrix, which has a relatively open structure (indicated as E₂ in Figure 4). A kinetic Raman study shows that LiTaO₃ nanocrystals are already formed within 5 h of the annealing, while it takes up to 20 h for the LiAlSi₂O₆ to fully grow (Supplementary Figure S12). Therefore, the intensity enhancement of the E₁ band originating from $(NiO_6)10^-$ in LiAlSi₂O₆, is rather weak with increasing annealing time, leading to the notable changes in the emission shape (Figure 4c–e).

In addition to the interesting dopant–host matrix interaction, the physical origin of the controllable emission output can also be attributed to the unique dopant–dopant interactions within the multiphase glass with the special mesoscale structure. As schematically shown in Figure 5, energy migration can occur between two different active sites of a material embedded in a host matrix.^{27–29} We assume that there is a contribution from an electric exchange to the energy migration between the emission centers E_1 and E_2 due to the overlapping orbitals. The energy transfer rate K_T and the efficiency E_T can be expressed according to equations (3) and (4):

$$KT = \frac{1}{\tau D} \cdot \left(\frac{R_0}{r}\right)^6 \tag{3}$$

$$ET = \frac{R_0^6}{R_0^6 + r^6} \tag{4}$$

where τ_D , R_0 and r are the donor lifetime, the critical distance and the separation distance, respectively. Dopants in a conventional homogeneous host matrix are in close physical proximity to each other, less than 3 nm at typical doping levels of 0.15 mol.%. In this case, dopant

interactions falls within the strong coupling region and the unexpected 'internal friction', such as the non-radiative energy transfers, prevails (Figure 5). It becomes extremely difficult to separately extract photons from the emission centers E1 and E2. Unlike in a homogeneous glass, the unique mesoscale heterogeneities present in the prepared glass-ceramic composite allow complete isolation of the emission centers E1 and E2 because of the notable increase in the separation distance, ≈ 10 nm. This shifts the correlation state of the E₁ – E₂ interaction to the weak coupling region (Figure 5). As a result, each dopant contributes to the emission output of the system, leading to a significant enhancement of the emission efficiency and a tunable spectrum shape. Importantly, the emission output with the highest full-width at half-maximum of approximately 480 nm and an extremely flat spectral shape covering the entire O, E, S, C and L telecommunication bands ($\lambda = 1260 - 1625$ nm) can be obtained (Figure 4c). To the best of our knowledge, this is the first time ever that simultaneous emission in the second and third telecommunication windows was achieved in a single-active material excited by a commercially available pumping source.

Construction of optical fibers

The newly developed Ni²⁺-doped glass-ceramic composite, with intriguing optical properties, was drawn into an optical fiber (see Experimental procedures). In a typical preform-to-fiber fabrication approach, a large size bulk glass with excellent optical quality is needed. As shown in Figure 6a, a macroscopic preform $(1 \times 1 \times 5.5 \text{ cm}^3)$ was successfully prepared by thorough mixing and continuous bubbling processes. This specially designed glass has a high propensity for devitrification, which may potentially degrade the optical performance of the drawn fibers. To circumvent this issue, a 'molten-core' strategy was used to ensure the core homogeneity

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| Table 1 | . Com | parison | of the | optical | properties | between | the | constructed | photonic | glass | -ceramic | fiber | and | the | typical | commercial | fibers |
|---------|-------|---------|--------|---------|------------|---------|-----|-------------|----------|----------|----------|-------|-----|-----|---------|------------|--------|
| | | | | | | | | | | <u> </u> | | | | | ~ . | | |

| Core material | Fiber preparation method | Emission center (nm) | Bandwidth (nm) | Ref. |
|--|---------------------------|----------------------|----------------|-----------|
| LiTaO ₃ and LiAISi ₂ O ₆ :Ni ²⁺ glass ceramics | Molten-core | 1250-1500 | 480 | This work |
| TeO ₂ :Er ³⁺ glass | Jacketing method | 1530 | 83 | 38 |
| ZBLAN:Pr ³⁺ | Jacketing method | 1310 | 100 | 39 |
| Υ-(Ga,Al) ₂ O ₃ :Ni ²⁺ glass ceramics | Rod-in-tube | 1250 | 250 | 40 |
| YAG:Cr ⁴⁺ crystal | Rod-in-tube | 1350 | 240 | 41 |
| SiO ₂ -GeO ₂ :Bi glass | Chemical vapor deposition | 1100 | 200 | 42 |



Figure 7 OA. (a) Schematic illustration of *in situ* optical signal amplification in a communication system. (b) Optical amplification properties at $\lambda = 1300$ nm. (c) Schematic representation of an optical amplifier used for ultra-long distance telecommunication. OA, optical amplifier.

(Supplementary Figure S13). The preform was inserted into a silica cladding tube, and fibers were drawn at 1950 °C. The drawing temperature was kept above the melting point of the core phase, $T\approx$ 1550 °C. We believe that fluent floppy-mode network glass cross-links with the cladding phase, resulting in the formation of strong interface bonding. Figure 6b shows the photograph of the drawn fibers. The obtained fibers show good uniformity of density and the standard core-cladding configuration and can also be well connected to the commercial silica fibers (Figure 6c and d). Structural, compositional and optical analysis showed that the key functional units, that is, LiTaO₃ and LiAlSi₂O₆ nanocrystals (Figure 6e–i and Supplementary Figure S13), could be obtained during fiber activation by annealing at T = 800 °C for 20 h. Importantly, the drawing process (Supplementary Figure S14) can prepare long fibers at commercially relevant speeds.

The optical properties of the typical active fibers are summarized in Table 1. The emission bandwidth of the Ni^{2+} -doped glass-ceramic fiber extends the range that can be achieved with commercially available fibers. Furthermore, the prepared fiber shows potentially strong nonlinear effects (Supplementary Figure S15), which originate from the spectral modulation function of the precipitated LiTaO₃ nanocrystals. Therefore, this novel fiber could also be applied as a multifunctional light source with a high level of integration.

It is believed that the excellent optical performance of the constructed fiber benefits from the proposed unique mesoscale engineering approach.

Optical amplifier

To examine the feasibility of the potential use of the glass-ceramic material in communication technology, a proof-of-principle device was constructed to measure the optical signal amplification performance. The test was conducted in a configuration resembling the conventional two-wave mixing geometry. Figure 7b shows the optical amplification property at $\lambda = 1300$ nm with an excitation power of ~1.1 W. The oscilloscope image with and without an optical pump clearly indicates that the intensity of the modulated probe signal can be amplified by ~40%. This equals an optical gain of 1.4 dB. The proposed principal optical amplifier device could potentially be embedded into a communication system and could support ultra-long distance telecommunication, for example, $\approx 14\,000$ km from Beijing to Washington, by regenerating the attenuated optical signal (Figure 7a and c). Another key advantage of our new composite material is the extremely flat spectral shape and notable emission bandwidth, which allows for in situ amplification of an optical signal in a wide wavelength region covering the entire telecommunication windows. As is well known, traditional Erbium-doped optical

amplifier can only provide limited bandwidth amplification (~11 THz).³⁰ In contrast, the Ni²⁺-doped glass-ceramic composite can potentially offer broad gain from 1140 to 1620 nm with a bandwidth of ~78 THz, thus being promising for use in the next-generation super-capacity information transmission systems. As an added benefit, the profound extended emission of the constructed material is also highly beneficial for minimizing signal cross-saturation and cross talk in these modern systems.⁸

CONCLUSIONS

We have successfully demonstrated an effective way for tuning and controlling photon emission by manipulating mesoscale heterogeneities in a multicomponent-glass-ceramic composite. Unlike the conventionally used approach, which has achieved only limited possibilities for tailoring the spectral broadening, the inhomogeneous mesoscale structure engineering protocol presented in this paper allows broadening of the emission bandwidth up to ~480 nm, that is, two times larger than that with the conventional approach, ~240 nm. For the first time, simultaneous extension of the emission bandwidth and a flat spectral shape in a multicomponent-glassceramic composite have been demonstrated. We believe not only that our findings will provide major steps toward precise control of the dopant-dopant and the dopant-host matrix interactions on the mesoscale but also that the results offer exciting opportunities for the construction of next-generation composite and fibers, which are urgently needed in high-capacity telecommunication, remote sensing and defense applications.^{31–33} Furthermore, the understanding of and manipulation on the mesoscale could lead to new material functionalities, which could potentially be applied in engineering a variety of materials, such as ceramics,34 cements,35 organic dyes36 and other composites,³⁷ where the mesoscale architecture becomes the typical microstructure of the applied system.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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