Fiber Evanescent Wave Spectroscopy based on IR fluorescent chalcogenide fibers

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Highlights  
• Optical chloroform detection  
• Pr³⁺ doped chalcogenide fibers  
• Tapered optical fibers  
• Pr: GaGeSbS spectroscopy  
• Mid-IR incoherent emission sources
Abstract

Chalcogenide glasses, owing to their transparency in the infrared window and the appropriate solubility of rare earth, allows the generation of middle infrared (mid-IR) radiation from a near infrared or visible pumping source. These emitted mid-IR broad bands can probe the vibrational modes of several molecules, e.g. C-H, C=O or C-Cl. Relying on this principle, a mid-IR optical sensor using the mid-IR fluorescence of Pr$^{3+}$: Ga-Ge-Sb-S fibers has been developed. The detection principle is based on Fiber Evanescent Wave Spectroscopy (FEWS). The spectroscopic characterization of praseodymium ions (Pr$^{3+}$) was performed in the near and mid-IR and is discussed on the basis of comparison with Judd-Ofelt calculations. The broad emission spectrum of the Pr$^{3+}$: Ga-Ge-Sb-S fiber from 4 to 5μm could enable the monitoring of multiple pollutants. In this study, chloroform detection is carried out via a novel technique derived from FEWS. In this way, an infrared sensor was developed, composed of a pumping source in near-IR, a mid-IR detector and a tapered Pr$^{3+}$: chalcogenide fiber to enhance the detection sensitivity. These results demonstrate for the first time the feasibility of detecting molecules by FEWS using the mid-IR fluorescence emitted by rare earth ions doping chalcogenide fibers. This method is an effective alternative to the classical FEWS system, as RE doped chalcogenide fibers have the advantage of being a compact mid-IR source.

Keywords:

chalcogenide glasses, fibers, FEWS; praseodymium, mid-IR optical sensor, chloroform
1. Introduction

A wide range of (bio)-chemical molecules have vibrational modes in the fingerprint region localized in the middle infrared (mid-IR) spectral range. Thus, the detection of such species using their strong absorption bands in mid-IR could be achieved using a dedicated optical sensor. Chalcogenide fibers based on sulfur, selenium, or tellurium enable the propagation of an infrared optical signal and enable the absorption bands of (bio)-chemical species to be collected. Indeed, these materials present a large optical window extending in the mid-IR and covering the two atmospheric windows ranging from 3 to 5 and 8 to 12 \( \mu m \). Fiber Evanescent Wave Spectroscopy (FEWS) simply requires dipping the fiber into the liquid under analysis or putting it in direct contact with the analyte. FEWS using chalcogenide glass fibers is an efficient tool and the chalcogenide fibers can be specifically shaped for head sensing thanks to the fibers’ thermo-forming capability [1]. It also allows investigation in different fields; e.g. the detection of pollutants in waste water [2, 3], the monitoring of chemical or industrial processes [4, 5], the detection of bacterial contamination in food [6, 7], and applications in the medical field [8]. Consequently, these materials can contribute to the development of pioneering glass technologies in IR spectral windows to design innovative optical sensors.

In addition to these specific characteristics in the passive optical mode, chalcogenide glasses can provide valuable emissions in mid-IR [9-13]. Their low phonon energy enables them to be transparent up to 20 \( \mu m \) in the case of telluride based materials, and rare earth ions can be efficiently incorporated into the chalcogenide host matrix. In the case of fluoride materials, the phonon energy is not low enough to generate wavelength lasing or even luminescence further than 3\( \mu m \). The rare earth transitions are easily quenched between two relatively close energy levels involved in the IR emission processes. For the 3-5 \( \mu m \) domain,
sulfide and selenide materials are quite suitable materials for use in optical fiber detectors, as the fiber drawing processes from these RE doped preforms is well mastered [10-15].

An optical sensor based on mid-IR fluorescence to probe gas molecules has already been developed using a transmission configuration instead of FEWS for the detection of carbon dioxide by means of the 4.3 μm emission of a Dy$^{3+}$ doped chalcogenide fiber [16]. Due to their high degree of mechanical integration and reliability (connectors, fiber coating techniques), optical fiber-based sensors are designed for commercial purposes. Compared to FEWS using undoped chalcogenide fibers as the sensitive media, this sensor has been developed in order to produce its own mid-IR fluorescence source. The resolution of real time detection could be enhanced and the overall compactness and complexity of the optical system can be efficiently reduced. FEWS relies on the optical absorption of the surrounding environment of the optical fiber, with a typical penetration depth of a few micrometers for the evanescent waves. The degree of integration of such devices is potentially high thanks to the rare earth doped chalcogenide fiber converting the pump signal from a visible or near infrared laser diode to mid-IR. By means of frequency conversion, the sensor could allow remote monitoring using silica fibers considering that the inverse photon conversion from mid-IR to visible or near-IR can be used based on a rare earth up-conversion phenomenon [17].

2. Materials and methods

2.1. Pr$^{3+}$:Ga$_5$Ge$_{25}$Sb$_{10}$S$_{65}$ bulk glass, fiber and taper fabrication

The composition of sulfide glass is Ga$_5$Ge$_{25}$Sb$_{10}$S$_{65}$ doped with 500, 1000 and 3000 ppm Pr$^{3+}$ ion concentrations. These glasses were prepared using conventional melting and quenching methods. Ga, Ge, Sb and S with high purity (≥ 5N) and praseodymium sulfide with 3N purity were used. Commercial sulfur was purified by successive distillations to remove
carbon (CO₂, CS₂, COS), hydrate, or sulfide hydride (H₂O, OH, SH) impurities [18]. Then, chemical reagents were put in silica tubes and pumped under vacuum (10⁻⁴ mbar). After sealing, the chemical reagents were melted under slow heating to 850°C for 10 hours in a rocking furnace to ensure homogenization. The ampoule was quenched in water, then annealed at 280°C for 3 hours. Single index fibers with a 350 μm diameter were obtained by drawing 7 mm diameter and 100 mm length Pr³⁺-doped Ga₅Ge₂₀Sb₁₀S₆₅ preforms. Tapers were obtained by heating the fiber locally and slowly stretching it. To perform the optical and spectroscopic characterizations, bulk glass slices were cut and polished.

2.2. Pr³⁺: GaGeSbS spectroscopic properties

Pr³⁺: Ga₅Ge₂₀Sb₁₀S₆₅ absorption spectra were recorded using a Perkin-Elmer Lambda 1050 and a Bruker FTIR Tensor 37 spectrophotometer for the near-IR and the mid-IR spectral domains, respectively. Fluorescence spectra were recorded using two pumping sources: a Lumics LU1470-T015 laser diode tuned at 1490 nm and a homemade Tm: YAG laser emitting at 2.0 μm. These two sources were used to investigate emission spectrum shape differences from one pumping wavelength to another [19]. The luminescence signal was collected using a nitrogen cooled InSb detector, a lock-in amplifier, and a monochromator with a resolution of 25 nm. The lock-in amplifier integration time was 1 s with a 35 Hz chopping frequency. The spectral response of the whole acquisition device was wavelength-dependent; emission spectra were calibrated using a tungsten-halogen lamp source and a heat source with emission spectra similar to a black-body source. Therefore, to record fluorescence signals without any parasitic absorption, such as carbon dioxide at 4.25 μm, the monochromator was flushed with dry nitrogen.

The incident power on bulk samples and fiber input was 800 mW. For lifetime measurements, a Continuum Horizon OPO laser delivering 5 ns pulses at a repetition rate of
10 Hz was used for the optical excitation. The signal was imaged on the InSb detector using a lens and passed through 2.36 μm, 4.00 μm and 4.64 μm centered band-pass filters respectively.

2.3. Fluorescent FEWS optical sensor

The Pr$^{3+}$: Ga$_5$Ge$_{20}$Sb$_{10}$S$_{65}$ fiber was pumped using a homemade Tm: YAG laser emitting at 2.0 μm pumped by a Ti: Sapphire laser tuned at 780 nm. The beam was focalized on the fiber using a first lens, and the output was imaged on the monochromator entrance slit using a second CaF$_2$ optics (Figure 8). To deduce the chloroform absorption from the emission spectra of the doped fiber, the emission signal obtained through an air flushed cell is used as the reference line. Two spectra with and without chloroform were then recorded, which gave the $I_{\text{diff}}(\lambda)$ curve as a function of the wavelength (eq.1).

$$I_{\text{diff}}(\lambda) = \frac{I_{\text{(empty cell)}} - I_{\text{(CHC$_3$)}}}{I_{\text{(empty cell)}}} \quad (\text{eq.} \; 1)$$

These $I_{\text{diff}}(\lambda)$ curves were then investigated as a function of the geometrical parameters of the taper. The signal was collected in a nitrogen cooled InSb detector.

3. Results and discussion

3.1. Pr$^{3+}$: GaGeSbS spectroscopic properties

3.1.1. Absorption and Judd-Ofelt calculations

The room temperature absorption cross-section spectrum of a 1000 ppm doped Pr$^{3+}$: Ga$_5$Ge$_{20}$Sb$_{10}$S$_{65}$ glass is presented in Figure 1-A. The Pr$^{3+}$ GaGeSbS glass shows overlapped of the $^3F_4$ (1580 nm), $^3F_3$ (1475 nm), $^3F_2$ (2020 nm) and $^3H_6$ (2250 nm) absorption bands. At a wavelength of 1020 nm, the weak $^1G_4$ absorption band could be identified. The infrared absorption cross-section between the fundamental $^3H_4$ and the first excited $^3H_5$ manifolds is presented in Figure 1-B. These absorption cross-sections feature peak values comparable to
other sulfide praseodymium doped materials [20]. In Figure 1-B, the emission band was calculated using the cross section reciprocity properties, as formulated by McCumber [21, 22]. The corresponding zero-line phonon energy was 2178 cm⁻¹.

Before performing the Judd-Ofelt analysis, the overlapped contributions had to be isolated by means of a pseudo-Voigt fit [23]. The mid-IR absorption band corresponding to the 3H₄ → 3H₅ transition extended from 3.8 to over 5.2 μm (Figure 1-B), indicating that the 3H₅ manifold Stark splitting was leading to a broad absorption level. From these integrated absorption bands, a Judd-Ofelt analysis was performed to estimate the emitting manifold lifetimes and branching ratios [24, 25]. Using a standard method of least squares adjustment, the calculated Ω₂,₄,₆ parameters were 10.3 ; 2.6 ; 8.0 x 10⁻²₀ cm², which is in good agreement with some other works [26]. The calculated manifold lifetimes were compared to the experimental values, as shown in Table 1.

In the following section, this comparison will be discussed in relation to some glass matrix effects, e.g. multiphonon relaxation rates and cross-relaxation processes. Even if the Judd-Ofelt calculation requires some approximation and its results should usually be considered with care, a huge discrepancy between the measured and the calculated values clearly indicates that another relaxation process occurs, invalidating the isolated ion case.

### 3.1.2. Emission properties

The near-IR emission spectrum of a 1000 ppm Pr³⁺ doped fiber with 1490 nm optical pumping is shown in Figure 2. The full width at half maximum (FWHM) of the emission band is about 250 nm, and is mainly composed of two major contributions from the 3F₃ → 3H₅ and 3H₆ → 3H₄ transitions (Figure 2 inset and Table 1). The observation of simultaneous emissions from the (3F₂, 3H₆) and (3F₄, 3F₃) manifolds is possible due to multiphonon relaxation rate being in the same order of magnitude as the radiative process rate. Considering the phonon energy of the germanium sulfide glass matrix (~ 340-430 cm⁻¹), approximately
three to four phonons are required to bridge the energy gap between the $^{3}F_{3} \rightarrow ^{3}F_{2}$ levels ($\approx 1300 \text{ cm}^{-1}$).

For the ($^{3}F_{2}, ^{3}H_{6}$) manifold, the luminescence is mainly due to the $^{3}H_{6} \rightarrow ^{3}H_{4}$ transition. Some reciprocity calculations separately performed on the $^{3}H_{6}$ and $^{3}F_{2}$ absorption bands showed that the calculated $^{3}H_{6} \rightarrow ^{3}H_{4}$ emission band matches the experimental data. The focus is on the mid-IR emission band, which is of interest for the above mentioned applicative purposes. Some investigations were performed regarding the influence of pumping wavelength, fiber drawing, and fiber length on the shape of the emission bands (Figure 3).

According to the absorption data and reciprocity calculations, the expected emission bands are in a good agreement with Park et al. for selenide materials [16(Par08)]. This broad mid-IR emission band is composed of the ($^{3}F_{2}, ^{3}H_{6}$) $\rightarrow ^{3}H_{5}$ ($3.5 - 4.2 \mu\text{m}$) and $^{3}H_{5} \rightarrow ^{3}H_{4}$ ($4.0 - 5.5 \mu\text{m}$) transitions, the structures of which are similar to those found in some other studies [14, 19]. In Figure 1-B, the emission line was calculated using the cross-section reciprocity properties, as formulated by McCumber [21, 22]. The calculated emission cross-section shows a maximum at a wavelength of 4750 nm, which is also the case in the experimental bulk glass emission spectrum (Figure 3), indicating that the fluorescence contributions above a wavelength of 4.5 $\mu\text{m}$ are due to the $^{3}H_{5} \rightarrow ^{3}H_{4}$ contribution.

The differences between the band shapes of the spectra arising from the pumping of the $^{3}F_{2}$ and ($^{3}F_{3}, ^{3}F_{4}$) manifolds were first studied (Figure 3-A). Few intensity variations could be noted at 4.5 $\mu\text{m}$ on 500 ppm doped bulk glass for the two pumping wavelengths of 1.49 $\mu\text{m}$ and 1.5 $\mu\text{m}$; however, the spectra remain quite similar. In the sulfide glasses, no modification in the emission band shape comparable to that reported by Park et al. in Pr$^{3+}$: GaGeSbSe glass [14] was observed. In Pr$^{3+}$: GaGeSbS glass, a strong contribution arises at 5.25 $\mu\text{m}$ with 2.0 $\mu\text{m}$ pumping, which is not the case with a 1.5 $\mu\text{m}$ pumping scheme.
Figure 3-B shows the comparison between the spectrum produced by a 92 mm long fiber and the corresponding bulk glass emission spectrum, and some remarkable differences are noticeable. Due to the length of the fiber relative to the optical path length available in bulk glass, the S-H and C-S absorption bands are more noticeable at 4.0 and 4.9 μm, respectively, on the fiber emission spectrum than on the bulk emission spectrum. Some amplified spontaneous emissions and reabsorption occurs along the propagation in the fiber, locally enhancing some contributions (4.8 and 5.0 μm for instance). In the fiber emission spectrum, the third order of the pump is not visible due to efficient pump absorption. The fiber length also has an impact on the emission band shape, allowing a level of reabsorption, amplified spontaneous emission (ASE), and impurity quenching processes. Some non-normalized emission spectra from fluorescence cut-back experiments for 500 ppm Pr\(^{3+}\) doped fibers are shown in Figure 4.

In Figure 4, for fiber lengths of about 10 cm (81 and 137 mm presented here), the emission intensities are quite comparable, indicating that the optimum length for this 500 ppm doped fiber mid-IR emission band is about 10 cm. The 179 mm long fiber 4.25-5.0 μm luminescence is affected by strong \(^3\)H\(_4\) → \(^3\)H\(_5\) reabsorption, which is a consequence of the previous pump absorption.

S-H and C-S impurity absorption bands are logically slightly more visible on the 179 mm long fiber emission spectrum. These trends are also noted for 1000 ppm doped fibers, with an optimum length of about 7 cm.

### 3.1.3. Lifetimes

The experimental lifetimes were measured at room temperature. Figure 5 shows the lifetimes of the \(^3\)F\(_4\), \(^3\)F\(_3\), \(^3\)F\(_2\), \(^3\)H\(_6\) and \(^3\)H\(_5\) manifolds with pumping at 1650 nm \(^3\)F\(_4\). 2.36 ± 0.40 μm, 4.00 ± 0.40 μm and 4.64 ± 0.40 μm filters were used to isolate the \(^3\)F\(_4\) \(^3\)F\(_3\) → \(^3\)H\(_5\),
$(^3F_2 \, ^3H_6) \rightarrow ^3H_5$, and the $^3H_5 \rightarrow ^3H_4$ transitions, respectively. The transmission width of these filters ensures the selective lifetime measurement of the two different contributions [14].

For the experimental fluorescence lifetime, the usual trend of faster decay as the Pr$^{3+}$ concentration increases was observed. The 500 ppm doped glass experimental lifetime was then compared to the Judd-Ofelt calculations. For the $^3F_3 \rightarrow ^3H_5$ transition, the calculated lifetime is about 400 $\mu$s, and the experimental lifetime was 109 $\mu$s. Considering that the Judd-Ofelt theory does not take into account any energy transfers, including luminescence quenching due to impurities, reasonable agreement remains between the calculated and measured values. The fluorescence lifetimes of the levels involved in transitions with lower energy differences witness an intense quenching process. The experimental lifetime for the $(^3F_2, ^3H_6) \rightarrow ^3H_5$ transition was shorter by more than an order of magnitude than the calculated value, and the same was true for the $^3H_5 \rightarrow ^3H_4$ transition, where the calculated lifetimes were respectively 8.2 and 15.9 ms and the experimental values were some hundreds of microseconds.

In other studies using selenide glasses, the measured lifetimes were much closer to the calculated values [14], such as 11.5 ms for the $^3H_5 \rightarrow ^3H_4$ transition for Pr: GeAsGaSe glass [11]. This indicates that a strong quenching effect seems to occur in sulfide glasses. This could be explained by the S-H bonds (4.0 $\mu$m), whose resonance belongs to the same spectral domain as the Pr$^{3+}$ emission, and also the S-C bond (at 4.9 $\mu$m). Absorption analysis of the glass preforms showed a low concentration at around 30 ppm for [SH] species. As also observed for other rare earth ions, this matrix could lead to some rare earth clustering, which could promote the quenching of luminescence and also contribute to a drop in fluorescence lifetime. This spectroscopic analysis of lifetimes allowed us to identify the necessary means to
develop a glass preform synthesis to improve the efficiency of the Pr\textsuperscript{3+}: sulfide mid-IR sources.

3.2. FEWS experimentations

The sensor is essentially composed of a tapered doped fiber hermetically mounted into a glass cell containing chloroform (Figure 6).

The detection experiments were performed by pouring chloroform into the glass cell and comparing the fiber emission spectra recorded with those obtained without chloroform. As already mentioned in the introduction, for FEWS detection, chalcogenide fibers have already proved to be suitable materials. This technique is based upon the evanescent field attenuation caused by the absorption of the surrounding media and related to the Goos-Hänchen shift in the case of total internal reflections. The evanescent field penetration depth is a function of the refractive index contrast between the analyte and the fiber [27]. For a Ga\textsubscript{5}Ge\textsubscript{20}Sb\textsubscript{10}S\textsubscript{65} fiber, the evanescent field penetration depth is about 1 μm for a wavelength in the 2-12 μm range. Previous experiments show that this value is sufficient to achieve a useful level of sensitivity for a passive optical fiber [3, 28, 29]. In this study, the fluorescence of the rare earth incorporated in the fiber generated the probing signal, instead of a more expensive and less compact mid-IR laser source.

To enhance the sensitivity of a FEWS sensor, the number of total internal reflections per unit length should be maximized, which is the aim of the tapered fiber [29, 30]. The thinner the taper is, the larger the number of total internal reflections that occur per length unit, therefore making the detection more sensitive. The slopes between Φ\textsubscript{1} and Φ\textsubscript{2} should not be too steep in order to minimize the optical losses of the infrared beam during the change in diameter (Figure 7).
For these tapers, the conical shape spreads over a length of about 1 cm for a 350 μm single index fiber tapered to diameters ranging from 80 to 40 μm. In our study, a 350 μm diameter was maintained at the input and output of the taper to ensure an efficient pumping beam injection for pumping powers at about 1 W. The experimental setup is shown in Figure 8. The CHCl₃ infrared absorption spectrum was also recorded using an FTIR spectrophotometer with a sealed cell with CaF₂ windows (Figure 9).

Chloroform possesses four main absorption bands corresponding to the stretching and bending modes of C-H and C-Cl bonds. Alongside these bands are some harmonic frequencies of lower intensity, like the band observed at 4.16 μm. Figure 9-B shows the liquid chloroform absorption spectrum versus the Pr: GaGeSbS mid-IR fluorescence signal. The CHCl₃ absorption band with a moderate absorption cross-section at 4.1 μm is overlapped by the Pr³⁺ emission generated by both ³H₆ → ³H₅ and ³H₅ → ³H₄ transitions, and suggest that chloroform detection could be implemented by the fluorescent FEWS technique. Apart from the feasibility of CHCl₃ detection using a Pr³⁺ doped chalcogenide fiber, the taper diameter effect on the absorption of the evanescent field and the effect of the rare earth concentration were studied.

The absorbed signal was higher with a tapered fiber, and could hardly be measured with 500 ppm Pr³⁺: GaGeSbS with a 350 μm core diameter. This absorption could not even be observed with a 1000 ppm doped fiber.

In a simplified geometrical approach, the reduction of taper diameter involves more reflections of light at the interface between the fiber and the chloroform, so that the evanescent wave is in greater contact with the absorbing solution, leading to more efficient sensing (Figure 7). For the 1000 ppm doped fiber, this enhancement was clearly visible, but this gain was more obvious on the 500 ppm doped fiber. The highest absorption signal was measured for a 40 μm tapered 500 ppm doped sulfide fiber.
4. Conclusion

During this work, fibers were successfully drawn from Pr: GaGeSbS preforms, and the mid-IR spectroscopic properties (absorption, emission, and lifetimes) of both bulk glass and the fibers were investigated as a function of the doping rate. Judd-Ofelt calculations were also performed.

For 350 μm initial diameter rare earth doped fibers, 80 to 40 μm tapers were fabricated. This tapered fiber was then optically pumped, and this embedded mid-IR source was used to perform FEWS experiments. Praseodymium doped fibers were used to detect the chloroform (CHCl₃) 4160 nm absorption band. For the first time, the detection of a pollutant was performed by the FEWS method using rare earth luminescence; more generally, this paper shows the possibility of using rare earth doped chalcogenide fibers in sensors. The use of the broad emission band of Pr³⁺ to detect CHCl₃ is witness to the fact that this particular fiber is able to detect several species having fingerprints in the 3-5 μm spectral range. On this basis, a dedicated sensor could be fabricated using a 4.1μm centered band-pass filter, combined with a monitoring application controlling both the laser diode source and a reference arm to enable differential detection.

Acknowledgements:

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References


Biographies

**R. Chahal** is a PhD student at the Chemical Sciences Institute (ISCR) at the University of Rennes, France. The subject of his thesis, supervised by Prof. Bruno Bureau and Dr. Virginie Nazabal, is devoted to the development of mid-IR optical sensors based on chalcogenide glasses applied for detection and monitoring of CO₂.

**Dr. F. Starecki** received his Ph.D. degree from the University of Caen, France, in 2013. This work was focused on visible and near-infrared laser emitting waveguides, grown by the liquid phase epitaxy technique (LPE). Now working as a post-doctoral associate at the Chemical Sciences Institute (ISCR) on supercritical CO₂ optical detection embedded in microreactors.

**Dr. Catherine Boussard-Pledel**, defended her PhD, focused on glasses in the system B–O–F, at the University of Rennes (France) in 1997. Since 1999, she is engineer of research at the CNRS, in the Glasses and Ceramics group at ISCR in Rennes. She is in charge of the development of fibers (multimode, single mode) and she works on the development of infrared sensors having applications in biomedicine, environment and spatial areas. She is involved in several international research programs. She is the author and co-author of about 100 publications and several patents.

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Dr. K. Michel is a scientific researcher at BRGM. She holds a PhD in Chemistry from University of Rennes. Her thesis is entitled “Infrared fiber optical sensor dedicated to detection of water pollution”. She has been involved on CO₂ storage since the Weyburn, Geocarbone monitoring, Sentinelle, Optique CO₂ projects. She participates to various European and national CO₂ storage projects, including CO₂GeoNet. Her key qualifications are: chemistry, soil gas, monitoring, sensors and mineral characterization.

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**Pr. B. Bureau**, PhD, “Local order investigations in fluoride glasses by multinuclear solid state NMR”, defended on 1998 at the University of Le Mans (France). From 1999 to 2006, he was Associated Professor in the “Glasses and Ceramics Laboratory” of the University of Rennes. After his accreditation to supervise research devoted to “optical and structural properties of non-conventional glasses”, he is Full Professor in charge of the passive optic group working on infrared optical fibers for sensors. He is also a solid-state NMR expert and is responsible for several international research programs. He is author and co-author of about 100 publications and more than 20 invited lectures.

**Dr. V. Nazabal** got her PhD degree in Solid State Chemistry from the University of Bordeaux (France) in 1999 after a master at Ecole Normale Superieure, at University of Paris VI. She joined the NIMS (Tsukuba, Japan) as post-doctoral fellowship in 2000. She belongs to the CNRS since 2001 and got accreditation in 2009. Her research activity is performed at the Glasses & Ceramics team of the Chemical Sciences Institute at the University of Rennes (France) with more than 100 publications as author or co-author and several invited contributions in international conferences. She received in 2010 a bronze medal of CNRS.
Figure 1. (A): Pr\(^{3+}\): GaGeSbS near IR absorption cross section, calculated from the 1000 ppm doped glass spectrum. (B): mid-IR absorption cross-section and emission cross-section calculated using the reciprocity method.

Figure 2. 1000 ppm Pr\(^{3+}\): GaGeSbS near (A) and mid-IR (B) fluorescence signal from a 1490 nm pumping. Inset: scheme of the major contributions.

Figure 3. mid-IR emission lines comparison: 1.49 and 2.0 \(\mu\)m optical pumping (A); fiber and bulk materials (B).

Figure 4. raw mid-IR emission spectrum of a 500 ppm doped Pr\(^{3+}\): Ga\(_5\)Ge\(_{20}\)Sb\(_{10}\)S\(_{65}\) fiber for several lengths. The 1490 nm pump power is of 800 mW. The 1490 nm pump power is fixed at about 800 mW.

Figure 5. Pr\(^{3+}\) GaGeSbS bulk glasses experimental lifetimes versus the Pr\(^{3+}\) concentration, ranging from 500 ppm to 3000 ppm (pumping at 1650 nm).

Figure 6. Glass tank containing chloroform with fiber mounted inside, side and top view.

Figure 7. Propagation of the light in a raw and a tapered fiber.

Figure 8. Experimental setup for detection of chloroform.

Figure 9. (A) Liquid chloroform infrared absorption spectra recorded by FTIR spectrophotometer (Red box: detected absorption band). (B) Pr: GaGeSbS mid-IR emission band and chloroform absorption at 4.1 \(\mu\)m.

Figure 10. FEWS experimentations results: 5000 ppm doped fiber (A) and 1000 ppm doped fiber (B).
Fig. 1

A

Pr$^{3+}$ GaGeSbS (1000 ppm)

Absorption cross section [$10^{-20}$ cm$^2$]

wavelength [nm]

B

Absorption
Emission

Cross section [$10^{-20}$ cm$^2$]

wavelength [nm]

Fig. 2

Fluorescence signal [A.U.]

Pr$^{3+}$ GaGeSbS

wavelength [nm]
Fig. 7

Fig. 8

2 μm Tm$^{3+}$: YAG laser

monochromator entrance slit

imaging optics

CHCl$_3$ cell

focusing optics
Fig. 9

Fig. 10
Table 1. Pr$^{3+}$: Ga$_5$Ge$_{20}$Sb$_{10}$S$_{65}$ Judd-Ofelt analysis report for the near and mid-IR involved transitions. Transitions with a branching ratio less than 5% are not listed ($\lambda_{em}$ is the mean wavelength, $\tau_{rad}$ the radiative lifetime, and $\beta$ the branching ratio).

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