Development of regenerated fiber Bragg grating sensors with long-term stability

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Abstract: The effect of annealing cycle on regeneration efficiency was investigated through isothermal treatments between 700 and 1000°C. We determined an inverse relationship between the recovery rate of the peak reflectivity and temperature. A regeneration efficiency of 85.2% and long-term stability at 1000°C for 500 hours were achieved via a slow regeneration process. Thermal sensors developed by isothermal regeneration were determined to be reliable up to 1000°C (±2 °C). Experimental findings suggest the involvement of both diffusion related phenomena and stress variation through densification of the fiber core in type-I FBG during the thermal regeneration process.

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References and Links
1. Introduction

Optical fiber based devices such as Fiber Bragg Grating (FBG) sensors have become highly attractive for nuclear, submarine, aerospace and space applications due to their resistance to electromagnetic interference, low density, mechanical flexibility, as well as great configurability [1–10]. A standard (type-I) FBG consists of a protective silica fiber cladding surrounding a photosensitive germanosilicate glass fiber core where periodic gratings are obtained by UV-inscription through a micrometric mask [11]. The periodic modulation of the refractive index generates highly selective spectral filtering effects that can be used to measure temperature and pressure in real time [8]. However, the sensing capability of FBGs, which depends on peak reflectivity with precise wavelength, degrades with temperature, limiting their implementation in power generating stations, space exploration modules and engine turbines [6].

In the last decade, different types of FBGs with superior thermal stability (Type-II, Type-In, Chemical Composition Gratings (CCG) and Regenerated Fiber Bragg Gratings (RFBG))
were developed in attempts to increase the thermal resistance of standard type-I gratings that can function only up to 425°C [4, 12, 13]. While CCG and RFBG have both shown superior performances, withstanding temperatures above 1000°C with exploitable spectral quality for operation in harsh environments over a few hours [4, 13, 14] to date there is still a strong controversy about the underlying mechanism of the so-called ‘thermal regeneration’ [12]. The challenge in understanding the regeneration phenomenon relates to the complex processing steps involving numerous parameters which affect the regeneration efficiency, defined as the percentage of the seed-grating reflectivity that remains upon regeneration [2, 4, 12, 15]. Gas loading (H\textsubscript{2} or Helium), pre-annealing, UV-inscription power and laser wavelength, the strength of initial (seed) grating and annealing cycle, as well as material properties such as fiber composition were all found to influence thermal recovery [2, 4, 12, 15]. In addition, strain is another important factor which affects the central wavelength and the thermal regeneration efficiency [1, 16]. The reported recovery rates vary enormously from 4% up to 74% depending on the parameters mentioned earlier [4, 6, 7, 17]. Maximizing the recovery rate upon regeneration is crucial for the development of FBG sensors with long-term stability, as peak reflectivity will be available for temperature measurements for longer durations.

This work focuses on the effect of annealing temperature on the regeneration process. In previous studies, two annealing approaches were mainly used for the thermal regeneration of FBGs: i) the first approach involves a single step continuous rise in temperature using a constant heating rate from room temperature (RT) up to 1000°C and above, whereas ii) in the second approach, the FBG is heated from RT to an intermediate temperature (~850–900°C), held isothermally for 10 to 60 minutes and then ramped up to 1000°C and above [4, 6,16, 18, 19]. The ultimate regeneration efficiency was reported to be superior when thermal regeneration occurs during isothermal holding at an intermediate temperature below 1000°C, in comparison with an annealing cycle where regeneration takes place during continuous heating up to 1000°C and above [6]. However, the value of this intermediate temperature used for isothermal treatments is dependent on seed grating strength, composition, dopant concentration and hydrogen loading [4]. Hence, this intermediate temperature, which is specific to the FBG under study, needs to be determined empirically before each thermal regeneration experiment. For FBGs with same processing route, even using different holding temperatures and annealing cycles allowed thermal regeneration and caused significant variations in recovery rates [4, 6].

Here we primarily investigate the effect of isothermal annealing in the 700-1000°C temperature range for thermal regeneration and also study the process dynamics to maximize recovery efficiency and long-term stability. Our experimental approach is based on the simultaneous measurements of FBG peak reflectivity, its Bragg wavelength and peak width, whose time evolutions were reported for FBGs of same origin, composition and optical properties. RFBG developed in this work were also tested for their reliability to be used as thermal sensors via calibration experiments up to 1000°C. Photoluminescence (PL) measurements were conducted on FBGs before and after regeneration for supplementary information. We finally compared our experimental findings with existing theories on the origin of the thermal regeneration mechanism.

2. Theoretical background

The reflection by a uniform FBG can be described by using four parameters: a central wavelength $\lambda_B$, the peak reflectivity, denoted as $\rho_B$, the peak width $\omega_B$ and the surrounding noise level which is primarily determined by the sensitivity of the detector. The Bragg wavelength obeys the following relation [13, 20]:

$$\lambda_B = 2n_A\Lambda$$

(1)
where $n_{\text{eff}}$ is the effective modal index of the fiber around 1.5 $\mu$m ($n_{\text{cladding}} < n_{\text{eff}} < n_{\text{core}}$) and $\Lambda$ is the period of the refractive index modulation inside the fiber core. The peak reflectivity ($\rho_B$) of spectral width ($\omega_B$) for strong gratings ($\Delta n_m > \sim 1.10^{-4}$) is given by the relation [13, 20]:

$$\rho_B = \tan h \left( \frac{\pi \eta \Delta n N}{2n_{\text{eff}}} \right)$$

$$\omega_B \propto \Delta n \lambda_0$$

where $\eta$ is the overlap integral of the guided mode in the fiber core, $N$ is the number of periodic variation ($\sim 2 \cdot 10^4$ in our case) and $\Delta n_m$ is the average amplitude of the refractive index modulation.

3. Experimental

The UV-inscription of commercial photosensitive optical silica fibers (IXFiber; 8.2 $\mu$m core diameter, 125 $\mu$m outside cladding diameter) containing 3.5 mole % $\text{GeO}_2$ to produce type-I FBGs was conducted at MPB Technologies Inc. Laboratories. Prior to FBG writing, the fibers were loaded with deuterium (D) under high pressures ($\sim 2000$ psi) at room temperature (RT) for 120 hours. After conducting a low temperature heat treatment (120 °C, 1 day), type-I (seed) FBGs with a transmission loss of $- 33 (+/- 2)$ dB and grating length of 10 mm were obtained via pulsed UV inscription ($\lambda$: 193 nm, power: 4-5 mJ, frequency: 70 Hz).

An erbium broadband laser source (1530-1565 nm) was used throughout the annealing cycles and reflectivity measurements were recorded in situ with 10 seconds (1 to 10 minutes at $T = 700$ °C) acquisition time via a Burleigh WA-7000 Multi-Line wavemeter of 0.5 pm wavelength measurement resolution. Additionally, reflectivity and transmission measurements of all fibers before and after regeneration were conducted using an Optical Spectrum Analyzer (OSA). Peak reflection values are presented as relative optical power in dB scale. The recovery rate of the peak reflectivity is based on the transmitted signal [6]. The experimental setup for high temperature regeneration experiments is depicted in Fig. 1.

Type-I gratings were annealed in steps with a constant heating rate of 25 °C/min between each step. For short-term stabilization and to compensate overshooting, samples are held for 5 minutes at each step. Firstly, the temperature was ramped up from RT to 500 °C and then using 100 °C temperature increments (50 - 200 °C for the final temperature raise depending on $T_{\text{final}}$), we reached $T_{\text{final}}$ between 700°C and 1000°C. For $T_{\text{final}}$ values above 900 °C, a direct ramp from 800 °C was used to avoid the onset of regeneration prior to reaching $T_{\text{final}}$, thereby ensuring for all samples that regeneration occurred at (stabilized) $T_{\text{final}}$. $T_{\text{final}}$ is kept constant ($\pm 1$°C) until the regeneration process is complete and the variation in reflectivity is negligible ($< 0.2$ dB). Finally, the samples are cooled down to RT inside the furnace before OSA transmission measurements. The data were extracted with a dedicated Matlab code.

For calibration of $\lambda_0$ shifts and the quality assessment of thermally regenerated FBG sensors, temperature measurements from 26 °C to 1000 °C with four repetitions were done. The calibration curves were drawn for RFBG sensors upon long-term annealing cycles.

Confocal PL measurements were performed, before and after regeneration, using an Olympus BX71 microscope, coupled to a Horiba iHR320 spectrometer equipped with a thermoelectrically cooled CCD array and a 0.9 N.A objective mounted orthogonally for both excitation and collection of the emission. The FBG is excited perpendicular to the fiber axis using the TEM00 mode of a 473 nm solid state laser, corresponding to a sample volume associated within a prolate ellipsoid of about 3 microns in height, with a radius of the airy disc of $\sim 350$ nm. The fiber is fixed horizontally on the X-Y-Z scanner, whose vertical position (Z) is adjusted from interfacial reflections to position the optical focus inside the fiber core, with a precision of about +/- 1 $\mu$m.
4. Thermal regeneration experiments

Thermal regeneration experiments were conducted at different final temperatures ($T_{\text{final}}$) on FBGs with initial seed grating strength of about $-33$ dB, fabricated using similar processes. A representation of the complete heating cycle is shown by the wavelength vs. time graph for a $T_{\text{final}}$ of 900 °C [Fig. 2]. Except for the experiment at 1000 °C, where $T_{\text{final}}$ is reached just before the disappearance of the Bragg peak, ramping to the final temperatures are completed in all samples before a major decrease in peak reflectivity and regeneration occurs. We found that regeneration takes place at all $T_{\text{final}}$ used (700-1000 °C), with different kinetics. The variation of peak reflectivity as well as peak width with time is given in Fig. 3 for $T_{\text{final}}$ values ranging from 900 to 1000 °C. We define the recovery time during which the peak reflectivity is not measurable as $\tau_{\text{rec}}$ [Fig. 3]. As vertical arrows indicate the time ($t_0$) when ramping to $T_{\text{final}}$ is completed, the onset time of peak width decrease occurs at a much earlier stage than the decrease in peak reflectivity. Even though the change of total refractive index already occurs around 300 °C, the effect on the peak reflectivity only becomes detectable later, often after $T_{\text{final}}$ was reached. This feature stems from the relationships given by Eqs. (2) and (3), where the peak width varies linearly with the refractive index [Eq. (3)] as compared to a highly non-linear relation for the reflectivity [Eq. (2)]. The FBG remains an almost perfect mirror at $\lambda_B$ even though the refractive index modulation decreases. In other words, structural or stress variations occur, resulting in a change in refractive index modulation at an early stage (=300°C, after 6-7 minutes of heating) in the regeneration process, yet the effect of this change can only be observed by peak reflectivity variation at a later stage. We operate FBGs with roughly 20,000 periods so even though the penetration depth of the beam at $\lambda_B$ increases with decreasing refractive index modulation, the combination of this modulation with the length guarantees almost 100% reflectivity until the modulation becomes too low to be compensated by the length of the grating.

Upon heating from ambient temperature to the maximum temperature [Fig. 2], $\lambda_B$ shifts almost linearly with a temperature coefficient of approximately 13 pm/K towards longer wavelengths following Eq. (1) in which, both the FBG periodicity (thermal expansion) and the mean refractive index (thermo-optic effect) of the fiber core depend on temperature.
The thermo-optic effect $n(T)$ dominates over thermal expansion so that typically 90% of the observed wavelength shifts are attributed to changes of the refractive index of the fiber core [20]. While holding the sample at $T_{\text{final}}$, the peak reflectivity drops to the noise level thus becoming undetectable, and after the recovery time $\tau_{\text{rec}}$ it recovers slowly and non-linearly to a value smaller than the initial one. Simultaneously, the peak width drops to zero, then becomes undetectable during the absence of a detectable reflectivity and recovers to a value
which is also inferior to the initial peak width [Fig. 3]. As far as the peak reflectivity is concerned, the process is referred to as FBG regeneration. During isothermal treatment, the Bragg wavelength decreases slowly [Fig. 2] while the noise level remains unaltered.

In Figs. 4(a)-4(f), we compare the measured peak reflectivity ($\rho_B$) with that obtained from the amplitude of the index modulation determined from independent measurements of $\omega_B$ and $\lambda_B$, using Eqs. (2) and (3) which required the use of a parameter related to $N$, $n_{eff}$ and $\eta$. Considering this parameter constant over time, shows a good overlap of the measured and calculated data in Fig. 4. These data are obtained for regeneration experiments conducted at 700 to 1000 °C. The good agreement between these curves makes our simultaneous independent measurements consistent with theoretical predictions. It indicates that all peak reflectivity variations during the regeneration process can be almost entirely attributed to the variation of the refractive index modulation inside the fiber core. This suggests that both the fraction of light inside the fiber core and the number of Bragg gratings remain unchanged during and after thermal regeneration. While the latter is not a surprise, the fact that the fraction of light inside the fiber core remains constant allows us to consider this as a one-dimensional problem along the fiber axis, with negligible contributions from radial modifications. This corresponds to the fact that the fiber core diameter is one order of magnitude larger than the grating wavelength.

![Fig. 4](image)

**Fig. 4.** (a-f) show the comparison between the measured Bragg peaks intensity ($\rho_B$) with that obtained from the amplitude of the index modulation determined from independent measurements of $\omega_B$ and $\lambda_B$.

4. Effect of regeneration temperature

4.1. Process efficiency and long-term stability

In the present work, instead of using a two-step annealing process as described in previous studies [6, 21], thermal regeneration was conducted isothermally (upon reaching $T_{final}$) in the 700-1000 °C temperature range. As we observed that regeneration was possible at all investigated temperatures, we obtained a clear understanding of the effects related to the isothermal temperature ($T_{final}$) on the regeneration efficiency. In Fig. 5(a), we plot the recovery rate determined from OSA measurements upon regeneration as a function of $T_{final}$.
From 700 to 1000 °C, we found an inverse relationship between $T_{\text{final}}$ and the percentage of the peak reflectivity recovered after thermal regeneration. The latter reaches a remarkable maximum of 85.2% at 700 °C, and a minimum of 10.1% at 1000 °C. The time evolution of the peak reflectivity measured at 700 °C is presented in Fig. 5(b). However, as expected, the time necessary for a complete regeneration process increases substantially at low temperatures: it required 450 hours at 700 °C compared to 1.3 hours at 1000 °C.

![Fig. 5. (a) shows the plots of regeneration recovery (%) vs. $T_{\text{final}}$, and $\Delta \lambda$ vs. $T_{\text{final}}$ (b) thermal regeneration at $T_{\text{final}} = 700$ °C (ρ vs. time).](image)

In addition, a higher peak reflectivity generates longer operation times of RFBG-based thermal sensors at high temperatures, because a more intense reflectivity will have better wavelength precision for temperature sensitivity for longer durations. Also, at very high temperatures the decay time of the higher peak reflectivity will be longer. To determine the differences in long-term thermal stabilities of RFBGs developed at ($T_{\text{final}}$) 700 °C (I) and 775 °C (II), we conducted isothermal treatments at 1000 °C. RFBG-I is initially regenerated at 700 °C with 85.2% reflectivity, and then after the furnace is cooled to RT, long-term annealing at 1000 °C for 500 hours was conducted. Similarly, RFBG-II was regenerated at 775 °C with a reflectivity of 49.7%, and then annealed at 1000 °C for 140 hours. The variation in $\rho_B$ upon long-term thermal exposure given in Fig. 6(a) indicates the superior thermal stability at 1000 °C of the grating regenerated at 700 °C (I) compared to the one regenerated at 775 °C (II). The higher peak reflectivity of RFBG (I) upon isothermal treatment at 1000 °C even after 500 hours is shown in Figs. 6(b) and 6(c) in comparison to the RFBG-II reflectivity after only 140 hours at 1000°C. Such increase in the thermal stability via regeneration at a lower $T_{\text{final}}$ represents a significant added-value for FBG-based thermal sensors used in harsh environments because it significantly extends their operability.

The final reflectivities of RFBG-I and RFBG-II were determined to be 43.8% and 15.9%, respectively. Hence, upon long-term thermal exposure (1000 °C, 500 hrs) RFBG-I preserves a high peak reflectivity with low degradation (Figs. 6(b) and 6(c)).

For thermal sensor reliability assessment, calibration measurements were conducted in the 26-1000 °C temperature range for RFBG-I with four repetitions [Fig. 6(d)]. Calibration curves, which are crucial for FBG sensors intended for industrial applications, indicate high repeatability. Wavelength measurements at 11 different temperatures for each calibration cycle are the average values of around 100 measurements. The maximum wavelength shift for a certain temperature was determined to be around 30 pm which corresponds to 2 °C. Hence, temperature values as a function of wavelength can be determined with maximum ± 2°C error [Fig. 6(d)].
4.2. Bragg wavelength shifts

The total change in wavelength was determined from $\lambda_B$ values (in nm) at RT before ($\lambda_i$) and after regeneration ($\lambda_f$). Figure 5(a) shows the overall change in wavelength ($\Delta\lambda = \lambda_f - \lambda_i$) variation at different $T_{final}$. A constant blueshift of ~1 nm (± 0.2 nm), independent of the isothermal temperature, occurs upon regeneration. The difference in $\lambda_B$ before and after the heating cycle depends on the depth of the FBG as expressed in decibel (dB) of peak reflectivity. A continuous blueshift (as presented in Fig. 2) was also observed in thermal decay studies conducted isothermally at very low temperatures ($T < 400$ °C) without the involvement of the regeneration process [22, 23]. The decrease in wavelength at constant temperature, which is attributed to the decay of the average refractive index of the fiber core, obeys a power-law [22, 23]. Discrepancies between experimental data and model were observed at higher temperatures ($T > 400$ °C) where a redshift occurred and was attributed to the presence of Boron in the cladding [22, 23]. Nevertheless, since the spectral shifts cannot be neglected during the ramp [Fig. 2], extracting accurate quantitative information from an analysis of this wavelength decay can lead to erroneous conclusions.

On the other hand, in a regeneration study conducted on fibers with 14-15% $B_2O_3$ and 10 mol% GeO$_2$, a continuous blueshift during isothermal treatment as well as a total $\Delta\lambda$ of ~2.1 nm (blueshift) were observed. This wavelength shift was attributed to the relaxation of core/cladding stress resulting in the reduction of the effective optical index [19]. On the contrary, Lai et al. related the blueshift of the wavelength to the inhibition of the structural relaxation in the core of the fiber via an increase in the $T_g$ of glass fibers [24].

In the present study, the most important result obtained for measured spectral shifts is related to their temperature-independence, indicating that the difference between initial and
stabilized Bragg wavelengths is not correlated with the regeneration efficiency. We associate this feature with a global increase of the tensile stress between the fiber core and the cladding, resulting from a structural densification of the whole fiber core during thermal regeneration. Such a variation of the fiber core volume induces changes of both the effective refractive index ($\Delta n_{\text{eff}}$) and the stress condition ($\Delta \sigma_z$), which can be expressed as follows [25]:

$$\Delta n_{\text{eff}} = n_{\text{eff}} \left( \frac{\lambda_i}{\lambda_f} - 1 \right)$$  \hspace{1cm} (4)

$$\sigma_z = \frac{2\Delta n_{\text{eff}}}{3C_1 + C_2}$$ \hspace{1cm} (5)

where $\lambda_i$ and $\lambda_f$ are the initial and final Bragg wavelengths, respectively, and $C_1$ and $C_2$ are the axial stress confining coefficients. From an effective refractive index reduction of about 0.07% obtained in this study, the change of axial stress can be estimated to be around 30.0 kg/mm² from stress optic coefficients $C_1 = 4.102 \times 10^{-5}$ mm²/kg and $C_2 = 7.42 \times 10^{-6}$ mm²/kg [25]. A similar axial stress value was reported for a fiber with 125 µm in diameter [25].

4.3. Activation energy

We report the temperature dependence of ‘recovery time’, $\tau_{\text{rec}}$, defined as the duration between the disappearance of the reflectivity in the background noise ($t_a$) and the appearance of the reflectivity ($t_b$), to determine the activation energy of recovery ($E_{\text{rec}}$). This approach is used to replace more complex curve-fitting functions using decay time dependence to varying temperatures with a simple time-temperature relationship [13, 26–28]. We assume that the time at which the reflectivity disappears in the background noise after $t_a$ is a reference point similar to all experiments conducted at different $T_{\text{final}}$. Hence, even though the regeneration process has not yet started at $t_a$, $\tau_{\text{rec}} = (t_b - t_a)$ can be considered as the incubation period for the onset of the regeneration process. As shown in Fig. 3, $\tau_{\text{rec}}$ depends on the temperature and its dependence is further investigated. As an FBG consists of periodic refractive index modulations in the fiber core, the thermal process responsible for the so-called ‘regeneration’ should be active between these periods at a fixed distance. Therefore, the basic characteristic diffusion length [Eq. (6)] can be used during $\tau_{\text{rec}}$.

$$\chi = \sqrt{Dt}$$ \hspace{1cm} (6)

$$A - \ln t = -\frac{E}{k_B T}$$ \hspace{1cm} (7)

where $D = D_0 e^{-\frac{\chi}{A t}}$, $\chi$ is the fixed length and $A$ is constant ($A = \ln \chi^2 - \ln D_0$).

Equation (7) is derived using Eq. (6) and accordingly $\ln \tau_{\text{rec}}$ vs. $1000/T$ plot was drawn covering the all temperature range (700-1000°C). A linear relationship was found, as shown in Fig. 7. The activation energy of recovery, $E_{\text{rec}}$, from the slope of the linear curve ($\ln \tau_{\text{rec}}$ vs. 1000/T) was determined as 2.64 (± 0.13) eV. The most significant error in measuring $\tau_{\text{rec}}$ from the reflectivity is due to the scattering in the initial background noise level that depends on various factors, including the quality of splicing, the deviation of the cut fiber surface in the open end and the structural variation in the fiber itself. The maximum scattering in the background is found to be ~0.7 dB. Hence, the error bars indicated in Fig. 7 are calculated from the variation of background levels and subsequent recovery time, labelled as $\tau_{\text{rec}}'$. 
Although activation energy values should depend on both fiber composition and FBG specifications, similar values have already been reported by different research groups using mainly the curve fitting approach [26–28]. We found that such an activation energy was consistent with the model proposed by Erdogan et al. [26], who associated the grating decay phenomenon with a de-trapping mechanism of electrons based on a distribution of activation energies peaked at around 2.80 eV (± 0.19 eV) for an Er-doped FBG. Another study based on a similar approach showed two different activation energy distributions for type-I (2.14 eV) and type-IIA (2.45 eV) gratings written on Boron (B)-Germanium (Ge) co-doped silica fibers [27]. Other thermal decay studies also reported activation energies ranging from 2.2 to 3.1 eV on FBGs with different fiber compositions [29, 30]. More recently, Holmberg et al. [13] determined two activation energy values of 3.05 eV and 0.89 eV via the rate of change in refractive index modulation and Bragg wavelength changes, respectively. It was stated that these energy values are due to diffusion-reaction dynamics of molecular water in silica glass [13]. On the other hand, a diffusional study conducted by Zhou et al. [31] reported an activation energy of 2.63 eV (similar to $E_{\text{rec}} = 2.64$ eV determined in this present study) for the removal of hydroxyl from silica glass in a temperature range of 700–900 °C.

To obtain complementary information regarding possible structural changes during regeneration, we performed PL measurements and optical imaging on FBG before and after regeneration at 775 °C. The measurements were carried out at RT, using an experimental approach that allowed us to connect the periodic refractive index modulation of the fibre core with local variations of oxygen-related defect emission [32]. Figure 8 displays the PL spectra recorded for a laser excitation spot focused inside the fiber core indicating a broad PL emission around ~560 nm (2.2 eV) which originates from multiple sources of oxygen defects in silica matrix. Oxygen defects (Non-Bridging Oxygen Hole Centers (NBOHC) ~517-620 nm (2-2.4 eV), Peroxy radicals ~629 nm (1.97 eV) and Si Oxygen Deficiency Centers (ODC) ~440-460 nm (2.7-2.8 eV)) [33–35] generated during the FBG writing disappear during the regeneration process. Moreover, FBG modulation seen in optical images [Fig. 8] before regeneration was not observable after thermal regeneration at 775 °C. The removal of oxygen defects through diffusional transport is consistent with the densification of the whole fiber core (evoked in sec. 4.2), which reduces the concentration of chemical dangling bonds. It can also be related to the disappearance of initial FBG refractive index modulation during thermal regeneration process, thus suggesting that the modulation before and after regeneration.
regeneration may have different origins. This is in agreement with the fact that the thermal stability of peak reflectivity increases significantly upon the completion of regeneration.

![Graph showing PL intensity vs wavelength before and after thermal regeneration process]

**Fig. 8.** PL spectra and optical images obtained before and after thermal regeneration process at 775 °C.

### 5. Comments on the regeneration mechanisms

Several studies associated the thermal regeneration with mechanisms of different origins, involving complex atomic diffusion or stress induced densification of the fiber core [4, 12, 13, 19]. Thermal diffusion related phenomena were suggested by Fokine *et al.* [36, 37] who developed CCG systems with refractive index modulations resulting from spatially periodic variations of fluorine (F) and oxygen (O) dopant concentrations. However, since the activation of these diffusion mechanisms was stated to be dependent on the presence of hydroxyl (OH) groups, such a scenario should not permit to regenerate B/Ge doped silica fibers after Helium loading [18], a remarkable feature demonstrated by Canning *et al.* [15] who correlated the main mechanism responsible for thermal regeneration with the occurrence of stress induced densification effects. In this latter approach, it is suggested that UV inscription induces a periodic pressure modulation inside the fiber core, which promotes the formation of a cristobalite phase during thermal annealing. Higher thermal stability of cristobalite compared to amorphous silica causes stronger gratings that can extend the temperature range of FBG operation up to 1295 °C [15].

This study presents regeneration experiments performed within a temperature range of 700-1000 °C, showing that lower temperatures \((T_{\text{final}})\) improve the regeneration efficiency as well as the long-term stability of RFBG sensors [Fig. 6]. As described earlier in section 4.3, the thermal regeneration process dynamics during ‘recovery time’ was found to follow an Arrhenius type behaviour as verified by the linear dependence of \(\ln \tau_{\text{rec}}\) vs. 1000/T, giving an activation energy of 2.64 (± 0.13) eV. The derivation of the characteristic diffusion length [Eq. (6)] was used on the assumption that if diffusional processes play a role in the thermal regeneration process as reported in previous studies [36, 37], diffusion should take place between equally spaced grating periods of about 500 nm. The linearity between \(\ln \tau_{\text{rec}}\) and 1000/T shown in Fig. 7 is an indication that the assumption is feasible and diffusional processes involve in the thermal regeneration process. A single activation energy, consistent with the reported activation energy values obtained in previous studies by thermal decay and regeneration investigations conducted on FBGs with varying compositions and thermal treatment, shows that one type of diffusional transport is dominantly active during thermal regeneration between 700 and 1000 °C. In addition, the disappearance of oxygen-related defects upon thermal regeneration indicates that diffusion takes place in this process.
Nevertheless, a regeneration theory based only on diffusion related mechanisms does not adequately describe the increase in recovery rate at lower regeneration temperatures, because higher thermal activation should promote atomic motion.

On the other hand, models referring to stress induced densification as the main mechanism for thermal regeneration are not consistent with our experimental findings regarding the overall wavelength shifts. The fact that these wavelength shifts do not depend on the regeneration temperature indicates that stress relaxations are not quantitatively correlated with final peak reflectivities upon regeneration. Therefore, we attributed this overall blueshift to a release of tensile stress between core and cladding. Its continuous evolution during thermal regeneration suggests a continuous increase in the $T_g$ of the fiber core, which is compatible with glass densification. This assumption is partially consistent with scenarios in which the main mechanism responsible for the thermal regeneration process is the crystallization of amorphous silica into the cristobalite phase [15], but our results evidence here that such mechanisms cannot be the primary cause of the increase in recovery rates. Even if we cannot exclude that the densification of the core can strongly affect the periodic mechanical constraints at the core/cladding interface, resulting from the Bragg grating inscription, such a scenario is insufficient to describe the evolution of the peak reflectivity over the whole duration of the regeneration process.

As a consequence of these experimental findings, we infer that both diffusional phenomena and stress induced mechanisms take part in the thermal regeneration process. For the moment, none of the proposed theories can describe the dependence of recovery efficiency of peak reflectivity on the regeneration temperature. This necessitates further investigations based on a new theoretical approach, which takes into account several contributions of different origins to the periodic refractive index modulation.

6. Conclusions and perspectives

In this work we examined the role of annealing temperature on the thermal regeneration process. We determined an inverse relationship between the temperature of regeneration and recovery efficiency reaching 85.2% at 700 °C. The long-term stability of regenerated reflectivity over 500 hours at 1000 °C exhibits great potential to be used as thermal sensors for space applications. Calibration measurements showed the high repeatability and reliability of RFBG sensors up to 1000 °C. Process dynamics was found to follow an Arrhenius type behaviour during recovery time where no reflectivity is detectable. The experimental data suggest the involvement of both diffusion dependent processes as well as stress evolution due to the densification of the fiber core. Hence, the role of both mechanisms should be considered for a clear understanding of the thermal regeneration process. Based on these results, new theoretical models capable of explaining both the diffusion and stress related mechanisms should be developed. Further investigations aiming to optimize parameters such as seed-grating strength and fiber photosensitivity on regeneration dynamics may shed light into the origins of the thermal regeneration process. These studies will allow us to understand the limits of this process for developing optimal performance FBG sensors.

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