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A B S T R A C T

The structural and optical emission properties of Er-doped silicon-rich silica layers containing $10^{21}$ at cm$^{-3}$ of erbium are studied as a function of deposition conditions and annealing treatment. Magnetron co-sputtering of three confocal targets (Si, SiO$_2$ and Er$_2$O$_3$) under a plasma of pure argon was used to deposit the layers at 500 °C. The silicon excess was varied in the layers in the range 7–15 at% by monitoring the power applied on Si cathode. The as-grown samples were found significantly emitting at 1.54 μm under non-resonant excitation. A maximum Er emission was observed after annealing at a moderate temperature (600 °C) for any amount of Si excess, with a highest 1.54 μm photoluminescence (PL) from the sample containing 13 at% of Si. While no nanocrystals were observed in the samples annealed at 600 °C, the sensitizers might, therefore, consist in ‘atomic’ scaled entities (Si agglomerates, for example) considered in recent similar work. The comparison of the emission features of our “best” sample and their counterparts reported so far, shows that the approach used this work allows to increase the fraction of the Er$^{3+}$ ions coupled to Si sensitizers from 3% up to 12% of the total Er content.

1. Introduction

A considerable effort is devoted since many years on silicon-based photonics [1] and, more particularly, on the use of Er ions [2,3] as optical centers emitting at the standard telecommunication wavelength of 1.5 μm. As host materials for Er, either crystalline Si (c-Si) or silica has shown specific drawbacks, such as the detrimental back transfer for c-Si [4] preventing any room-temperature functionality and the low absorption cross section ($\sim 10^{-20}–10^{-21}$ cm$^2$) for SiO$_2$ [5] requiring high-power resonant excitation. These problems have been overcome by the coexistence of Er ions with Si nanoclusters (Si-nc) within a silica matrix. Indeed, these Si-nc were found to induce an indirect excitation of Er [6] that allows an increase of the effective excitation cross section of Er ions by 3–4 orders of magnitude, while maintaining the absence of energy back transfer owing to the large bandgap of both silica and Si-nc [7,8]. The presence of Si-nc provides additional advantages such as the increase of the refractive index for better optical guiding and the improvement of both injection and transport of carriers for electrical excitation. On the other hand, it appeared that either crystalline or amorphous Si-nc are still sensitizers [8,9], provided that their size is lower than about 4 nm [10] and their separating distance from the Er ions is lower than 0.5 nm [10,11]. Such a very short interaction distance is considered as being the main factor limiting the fraction of Er ions coupled to Si-nc to a few % [12,13]. This situation requires, therefore, some nanoengineering of the material composition through the growth of a high density of very small Si-based sensitizer entities to maximize the fraction of coupled Er.

The present paper deals, therefore, with the optimization of such a material. Series of layers were grown by magnetron co-sputtering and subsequently annealed before being submitted to various structural and optical characterizations. The effects of both and either Si excess and annealing treatment have been particularly examined.

2. Experimental details

The series of Er-doped silicon-rich silicon oxide (Er-SRSO) layers were deposited by magnetron co-sputtering of three confocal targets, SiO$_2$, Er$_2$O$_3$ and Si under a plasma of pure argon. This novel approach used by our team differs from that used earlier when a unique SiO$_2$ target topped by chips of Er$_2$O$_3$ is sputtered under a mixture of Ar+H$_2$ plasma [8,11,13]. The content of Er and Si excess can be finely and independently tuned through the monitoring of the RF power applied on each cathode. The total pressure in the chamber and the power density applied onto SiO$_2$ and ErO$_3$ cathodes were fixed at 3 mTorr, 7.4 and 0.74 W/cm$^2$, respectively, while that on the Si target ($P_{RF}^Si$) was changed between 0.74 and 2.07 W cm$^{-2}$. All samples were grown at a
substrate temperature $T_s = 500$ °C on a rotating holder for good homogeneity of both thickness and composition. The deposited layers were subsequently annealed under a nitrogen flux for 1 h at different temperatures $T_a$ between 500 °C and 900 °C. The structural properties were investigated by Fourier transform infrared spectroscopy (FTIR) measurements performed under normal and Brewster ($65^\circ$) incidence in the range $500$–$4000$ cm$^{-1}$, using a Nicolet Nexus spectrometer. The refractive index and the thickness of layer were obtained by M-lines measurements performed in reflection configuration [14–16], using a He–Ne laser emitting at 632.8 nm polarized in transverse electric (TE) mode. The optical properties were examined by photoluminescence (PL) measurements recorded with a non-resonant excitation (476 nm) line delivered by an Ar$^+$ laser, in order to ensure that Er was excited through the Si sensitizers. The Er PL was measured with a Jobin Yvon 1 m single grating monochromator and Northcoast Germanium detector cooled with liquid nitrogen. The PL signal was recorded through a SRS lock-in amplifier (SP830 DPS) referenced to the chopping frequency of light of 20 Hz. The time-resolved Er PL was measured with the already described PL system, the signal being displayed by a Tektronix oscilloscope (TDS 3012B).

3. Results and discussion

The Si excess induced by the variation of $P_{RF}^\text{Si}$ has been estimated from the evolution of the TO$_3$ mode of the FTIR spectra recorded at normal incidence, as shown in Fig. 1, following the method described in Ref. [17]. Briefly summarized, this method is based on a linear relation between the shift of the TO$_3$ peak position for a sub-stoichiometric SiO$_x$ ($1 < x < 2$) matrix, with respect to that of sputtered SiO$_2$, and the stoichiometric coefficient $x$. Fig. 1 shows a gradual shift of the TO$_3$ peak position towards lower values of the wavenumber, when ($P_{RF}^\text{Si}$) is increased. This behaviour implies some proportionality between the oxidation degree of the matrix, or the coefficient $x$, and $P_{RF}^\text{Si}$ responsible of the Si excess. Indeed, the TO$_3$ peak shifts gradually from 1080 cm$^{-1}$ for $P_{RF}^\text{Si} = 0$ (silica film free of Si excess) to 1062 cm$^{-1}$ for the sub-stoichiometric layer grown with $2.07 \text{W cm}^{-2}$. The estimated Si excess was found to increase from about 7 at% to nearly 15 at% when the $P_{RF}^\text{Si}$ was increased from 0.74 to 2.07 $\text{W cm}^{-2}$, as shown in the inset of Fig. 1. Such an increase is corroborated by the evolution of the refractive index shown in the same inset, and which increases with $P_{RF}^\text{Si}$ from 1.489 to 1.646, as deduced from M-lines measurements.

Since the samples should contain the optimum concentration of emitting centers, we have incorporated an Er content as high as $10^{21} \text{cm}^{-3}$ which appears higher than the threshold value ($\sim 2 \times 10^{20} \text{cm}^{-3}$) preventing up-conversion phenomena [18]. However, such detrimental phenomena can be minimized when using moderate values of excitation flux $\phi$ ($10^{18}$–$10^{19} \text{ph cm}^{-2} \text{s}^{-1}$). Indeed, the typical evolution of the PL intensity at 1.54 $\mu\text{m}$ vs $\phi$, displayed in Fig. 2 for both as-deposited and annealed (at 600 °C) samples, shows an almost linear behaviour that is indicative of limited effects of up-conversion phenomena and concentration quenching. To note that the emission observed from

![Fig. 1.](image1.png) Shift of the TO$_3$ peak position as a function of the power density applied on the Si target. The inset shows the evolution of either the Si excess, as estimated from FTIR spectra, and of the refractive index, as determined by M-lines, in function of the power density applied on the Si target.

![Fig. 2.](image2.png) Variation of the PL intensity at 1.54 $\mu\text{m}$ for the as-deposited and annealed samples as a function of the photon flux of the non-resonant excitation line 476 nm.
the as-grown samples is likely originated from the formation of Si-based sensitizers during the growth process. The annealing at relatively low temperature (600 °C) improves the emission by a factor 2, probably due to the formation of additional sensitizers together with the expected improvement of the matrix.

Fig. 3a shows the effect of the annealing temperature \( T_a \) on the evolution of the Er PL for various values of the Si excess. The general trend is almost similar for all values of Si excess: a first slight increase up to a maximum for \( T_a = 600 \) °C, followed by a gradual decrease for higher values of \( T_a \). This evolution is hardly observable for the lowest values of Si excess (7.5–8.3 at%), but it is the most pronounced for 13 at% of Si excess. To note that the level of PL increases with the Si excess, all along the range, reaches a maximum for 13 at% that appears as the optimum value for the best emission and then decreases for the highest value of 14.7 at%.

Such a result might indicate that, for any \( T_a \) value, the highest coupling between Er ions and Si-based sensitizers takes place for 13 at% of Si excess. Concerning the evolution of the PL level against the Si excess, one may suggest the following: (i) the first increase of Si excess is expected to enhance the density of Si-nc sensitizers until probably saturating the available seeds (for a given \( T_a \)) which seems to occur for 13 at% of Si excess; (ii) further increase of Si excess, from 13 to 14.7 at%, might lead to increasing the average size of the formed Si-nc at the expense of their density and then of their coupling with Er ions. Anyway, these Si-nc are all amorphous, since \( T_w < 900 \) °C and then hardly observable by transmission electron microscopy. The emission lifetime \( \tau_{PL} \) is generally increased when \( T_a \) is raised, even though this improvement is very limited, if not absent, for the highest values of Si excess with 1.5 ms < \( \tau_{PL} < 2 \) ms, as shown in Fig. 3b.

The most significant feature concerns the notable emission observed from the non-annealed sample grown at 500 °C (Fig. 2) and the maximum PL recorded after annealing at moderate temperature, \( T_a = 600 \) °C (Fig. 3a), for which no ‘large’ Si-nc might form. These two observations are both indicative of the production of some Si-based entities (agglomerates of atoms, for example) that play the role of sensitizers. These entities might be compared to the ‘atomic’ scaled sensitizers that have been suspected to form at the same moderate temperature (600 °C) in similarly sputtered samples [19]. The ‘atomic’ scaled sensitizers should be dense enough to ensure a high coupling and this could explain the significant PL observed from our as-grown samples, as well as the maximum emission detected after annealing at only 600 °C. To attempt a rough estimate of the proportion of Er ions coupled to Si sensitizers, we compare in Fig. 4, the Er PL spectra of our best sample (13 at% of Si excess) recorded before and after annealing at 600 °C with the PL spectrum of the best sample obtained earlier with the sputtering of a silica target topped by Er2O3 chips [8,11,13]. This latter is considered here as a reference, was annealed at 900 °C and contains, however, lower Er content (5 × 10^20 at cm^-3) and Si excess (7 at%). Among the Er ions contained in the reference sample, only 3% were found coupled to sensitizers [13], i.e. ~1.5 × 10^19 cm^-3. As the spectra of Fig. 4 were recorded with the same photon flux (~10^19 ph cm^-2 s^-1), we assume that the intensity (normalized to the excited volume of the sample) \( I_{PL} \propto \sigma_{eff} N_{Er,c} \tau_{PL}/\tau_{RAD} \) where \( \sigma_{eff} \) is the effective excitation cross section, \( \tau_{PL} \) is the measured lifetime, \( N_{Er,c} \) the concentration of Er ions coupled to Si-nc, which is equal to about 1.5 × 10^20 cm^-3 for the reference sample.

By comparing the \( I_{PL} \) of the reference to that of our present layer (3.7 times higher), the values of their \( \tau_{decay} \) (4.5 vs 1.5 ms), the values of their \( \tau_{rad} \) which is \( \propto n^2 \) (\( n \) is the refractive index and is equal to 1.52 for the reference and 1.56 for the examined sample), can be deduced from Ref. [20] (9 vs 12 ms), and the ratio of their \( \sigma_{eff} \) (5–6 × 10^-18 vs ~10^-17 cm^-2), one could estimate \( N_{Er,c} \) for our best sample to about 1.2 × 10^20 cm^-3. This value represents nearly 12% of the total Er content (~10^21 cm^-3) and might correspond to higher percentage of the optically active Er

![Fig. 3](image_url)  
Fig. 3. (a) Variation of the PL intensity at 1.54 µm for the annealed samples containing the indicated values of Si excess, as a function of annealing temperature \( T_a \). The samples were excited with the non-resonant line (476 nm). (b) Evolution of the corresponding emission lifetime against the annealing temperature \( T_a \).

![Fig. 4](image_url)  
Fig. 4. Comparison of the PL spectra recorded on the best sample of Fig. 3 (13 at% Si excess, annealed at 600 °C) to those of the as-grown sample and the best sample described in Ref. [13] (Si excess: 7 at%, Er content: 5.4 × 10^20 at cm^-3, annealing temperature: 900 °C). The samples were excited with the non-resonant line (476 nm) and a flux of ~10^19 ph cm^-2 s^-1.
ions. Considering the relatively high value of Er content (~$10^{21}$ \text{cm}^{-3}), one should be cautious concerning the concentration-related detrimental effects (up-conversion, concentration quenching, ...), as for their impact on the improvement of the coupled fraction of Er. There is an ongoing work on lower Er content (few $10^{20}$ \text{cm}^{-3}) for further optimization of the material which is necessary to couple the majority of the optically active Er ions and minimize the losses for the achievement of a net gain.

4. Conclusion

In conclusion, the effects of both $T_a$ and Si excess have been examined with the aim of improving the fraction of Er coupled to Si-nc sensitizers. While the as-grown samples at 500 °C were found significantly emitting, the highest emission was detected from the samples containing 13 at% of Si excess and annealed at $T_a = 600$ °C. Such a behaviour with a non-resonant excitation (476 nm) is ascribed to the sensitizing effects of some 'atomic' scaled entities formed at relatively moderate temperature (500–600 °C). A rough estimate of the fraction of coupled Er has allowed to notice that their proportion was improved from about 3% for earlier samples to nearly 12%. Further optimization of the material is still needed, together with processing at moderate temperature, for achieving an amplification silica medium with Er-coupled to Si-based sensitizers.

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