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Cathodoluminescence and photoluminescence comparative study of erbium-doped silicon-rich silicon oxide

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Abstract. We present a study on erbium (Er)-doped silicon (Si)-rich silicon oxide thin films grown by the magnetron cosputtering of three confocal cathodes according to the deposition temperature and the annealing treatment. It is shown that, through a careful tuning of both deposition and annealing temperatures, it is possible to engineer the fraction of agglomerated Si that may play the role of sensitizer toward Er ions. To investigate the different emitting centers present within the films according to the fraction of agglomerated Si, a cathodoluminescence experiment was made. We observe in all samples contributions from point-defect centers due to some oxygen vacancies and generally known as silicon-oxygen deficient centers (SiODC), at around 450–500 nm. The behavior of such contributions suggests the possible occurrence of an energy transfer from the SiODCs toward Er$^{3+}$ ions. Photoluminescence experiments were carried out to characterize the energy transfer from Si nanoclusters toward Er$^{3+}$ ions with a nonresonant wavelength (476 nm) that is unable to excite SiODCs and then exclude any role of these centers in the energy transfer process for the PL experiments. Accordingly, it is shown that structural differences have some effects on the optical properties that lead to better performance for high-temperature deposited material. This aspect is illustrated by the Er-PL efficiency that is found higher for 500°C-deposited, when compared to that for RT-deposited sample. Finally, it is shown that the Er-PL efficiency is gradually increasing as a function of the fraction of agglomerated silicon.

Keywords: erbium; silicon-nanoclusters; thin films; energy transfer; photoluminescence; cathodoluminescence.

1 Introduction

The integrated circuit was invented fifty years ago and has continuously improved ever since. Microelectronics industries can now routinely integrate one billion transistors on a single chip. This tremendous progress is now facing several major issues lying mainly in increased signal delay and higher power dissipation, referred to as “interconnection bottleneck.” Such problems associated with the electrical and thermal parasites could be overcome if the present metallic interconnections are replaced by photonics devices. Recently, a fully silicon-based photonics device was demonstrated for interchip communications.1 Nevertheless, for intrachips communications, interconnections must be integrated into microprocessors and a smart choice would be to use the “less-absorbed wavelength in silica,” such as the standard 1.5 μm to transport the
information. For this purpose, erbium (Er) ion in silica is a good candidate as an emitting center at the standard 1.5 \( \mu \text{m} \), corresponding to the \( ^4I_{13/2} \rightarrow ^4I_{15/2} \) transitions within a 4f shell. Unfortunately, this rare earth is very weakly excited in silica, with an absorption cross section \( \sigma \) as low as \( 10^{-20}–10^{-21} \text{ cm}^2 \) that prevents the achievement of optical gain within compact devices. However, it was shown and demonstrated that this low absorption can be effectively increased by a factor as high as \( 10^4 \), when Er\(^{3+} \) ions are excited through the neighboring Si-nanoclusters (Si-nc) that absorb the excitation photons along a broad band with a cross section close to \( 10^{-16} \text{ cm}^2 \). Such findings led the scientific community to consider the erbium-doped silicon-rich silicon oxide (SRSO:Er) as a promising material for integrated photonics, although few studies reported some internal optical gain. Two main factors are considered as limiting the achievement of a net gain: 1. the low fraction of Er coupled to Si-nc that stems from the distance-dependent energy transfer between Er\(^{3+} \) ions and Si-nc sensitizers with a critical distance lower than 1 nm, and 2. the low fraction of optically active Er\(^{3+} \) ions. Recently, our group demonstrated a way to increase the fraction of coupled Er, from 2–3% to about 22%, by carefully optimizing the concentrations of Si-excess and erbium, and also showed the importance of the temperature of deposition for increasing the number of optically active Er\(^{3+} \) ions. Such advances are still, however, insufficient to achieve a net optical gain, which require the coupling of the majority of Er ions to the sensitizers. The latter are not well understood, especially as the Si-entities able to sensitize Er\(^{3+} \) ions appear in different states, from atomic-scale centers to well-defined Si-nanocrystals as well as amorphous Si-nanoclusters. The purpose of this paper is to investigate the nature of the Si-based sensitizers considered as responsible of an energy transfer toward Er\(^{3+} \) ions. By successively tuning the deposition and annealing temperatures, we were able to obtain a wide range of different state of Si-based sensitizers. A combination of emission dispersive x ray spectroscopy (EDX) and Fourier-transform infrared (FTIR) spectroscopy allows us to characterize the Si-based entities in term of percentage of agglomerated Si. Thanks to cathodoluminescence (CL) spectroscopy, the different emitting centers present in the films were identified, such as silicon-oxygen deficient centers (SiODCs) that were observed in all of our samples. To avoid any influence from SiODCs on the energy transfer, photoluminescence (PL) experiments were carried out using a 476 nm excitation wavelength that is nonresonant for Er\(^{3+} \) ions and also below the excitation threshold for SiODCs.

2 Experimental Details

The samples used for this study were deposited by the magnetron cosputtering of three confocal cathodes: SiO\(_2\), Si, and Er\(_2\)O\(_3\). The power density applied on each of these cathodes can be independently tuned and are fixed at: \( P_{\text{SiO}_2} \approx 8.88 \text{ W/cm}^2 \), \( P_{\text{Si}} \approx 1.63 \text{ W/cm}^2 \), and \( P_{\text{Er}_2\text{O}_3} \approx 0.44 \text{ W/cm}^2 \). The total plasma pressure was kept constant at 2 mTorr during all deposition runs. The sample holder can be intentionally heated during the deposition and thus the deposition temperature, \( T_d \), was varied from room temperature (RT, i.e., \( \sim 20^\circ \text{C} \)) to 700°C. The substrate holder is rotated during the growth process, in order to ensure a good homogeneity for both composition and thickness. The duration for all depositions was kept constant at 10 h. Some samples were submitted to a postdeposition annealing during 1 h under a flux of pure nitrogen. The annealing temperature, \( T_a \), was varied from 600 to 1100°C.

The thickness was determined by the M-lines technique. The Er content and the stoichiometry of SiO\(_x\) matrix were measured by EDX. The FTIR absorption spectra were recorded under normal incidence to investigate the evolution of the TO\(_3\) peak, which provides some insights on the structural properties, such as the degree of phase separation and a rough estimate of Si excess.

CL spectra were obtained using a scanning electron microscope with CL system under high vacuum system. The electron beam is provided by a thermal field emission scanning electron microscope (Hitachi S4200) equipped with a CL system whose beam diameter was around tens of nm. The electron energy and beam current were kept constant at 10 kV and 4 nA, respectively, hence allowing appropriate comparison between the different samples. The light emitted by the sample is focused to an optical fiber and guided through the monochromator.
(Jobin-Yvon Triax 320). The signal is thus detected by a charge coupled device of 2048 channels (CCD, Jobin-Yvon Spectrum One). The excitation time was kept the same (10 s) for all samples and the recorded spectra were automatically converted into counts per seconds (cps).

PL measurements were carried out using the nonresonant 476 nm wavelength of an Ar$^+$ laser, in order to ensure that the Er$^{3+}$ ions are indirectly excited. The spot of the laser beam was carefully determined by a “moving knife-edge” method and was found to be around 3 mm$^2$ at $1/e^2$ of the maximum intensity. For the visible PL assumed to originate from the Si nanoclusters, the emitted light is detected by photomultiplier tube (Hamamatsu). For the Er-related PL in the IR range (1.53 μm), the emission is guided through a 1 m single-grating monochromator (Jobin Yvon) and collected by a liquid-nitrogen-cooled Germanium detector (Northcoast). All PL measurements were made using the standard lock-in technique with a chopper setting the reference frequency at 10 Hz.

3 Results and Discussion

3.1 Compositional Characteristics and Structural Properties

The influence of the deposition temperature, $T_d$, on the properties of 300–400 nm-thick films was already analyzed in our previous recent study. However, the present deposited films are much thicker (1.3–1.8 μm) what may result in several quantitative differences. Figure 1 shows the evolution of the thickness, the Er content and the stoichiometry parameter, i.e., the ratio $x = [O]/[Si]$, in function of $T_d$.

It appears that the thickness is almost constant for $T_d \leq 300$°C, before progressively decreasing for $T_d > 300$°C. In parallel, the Er content continuously decreases for $T_d$ higher than 300°C, and loses one order of magnitude when $T_d$ reaches 700°C. Concerning the composition of the as-deposited samples investigated by EDX, the stoichiometry parameter $x$ decreases by about 10% when $T_d$ exceeds 300°C, reaches a minimum of about 1.4 for $T_d = 500$°C, and then slightly increases.

Besides, $T_d$ is likely to also influence the surface mobility of the deposited species during the deposition process, hence favoring the formation of Si agglomerates that may play the role of sensitizers toward the Er ions, when $T_d$ is increased. Though this phenomenon was assumed

![Graph](image_url)
to occur in our previous study, it was not investigated. To fulfill this need, we made use of two complementary methods: FTIR method able to uniquely detect the Si bonded to O (see below) and the EDX that is capable of measuring the total silicon excess from the following relation:

$$SiO_x \rightarrow \frac{x}{2}SiO_2 + (1 - \frac{x}{2})Si,$$

with $x$ being the stoichiometric parameter (i.e., the ratio $[O]/[Si]$ as measured by EDX). We can, therefore, estimate the total Si excess in atomic percent by

$$\%Si_{Excess} = 100 \times \frac{1 - (x/2)}{1 + x}. \quad (2)$$

However, to get the fraction of agglomerated Si, we used the FTIR spectroscopy method that is based on the evolution of the TO$_3$ peak, which is related to the amount of Si unbonded to another Si atom, but rather bonded to oxygen. With this method, as soon as some Si-excess atoms are agglomerated, the “stoichiometric” parameter $y$ (i.e., the ratio $[O]/[Si]$ obtained by FTIR) is different from the parameter $x$ obtained by EDX. Consequently, the Si-excess content, as measured by FTIR, is underestimated when agglomeration of Si atoms occur (i.e., $y > x$).

The parameter $y$ corresponds therefore to a SiO$_y$ matrix composed of isolated atoms of Si excess bonded to oxygen within SiO$_2$ matrix. This parameter $y$ of SiO$_y$ is linked to the wavenumber value of the TO$_3$ peak ($v_{TO3}^{SRSO_{-Er}}$) by the following linear relationship:

$$v_{TO3}^{SRSO_{-Er}} = A \cdot y + v_{TO3}^{Si}, \quad (3)$$

For SiO$_2$, $y = 2$, we obtain

$$A = \frac{v_{TO3}^{refSiO2} - v_{TO3}^{Si}}{2}. \quad (4)$$

The parameter $y$ can then be deduced from

$$y = 2 \cdot \frac{v_{TO3}^{SRSO_{-Er}} - v_{TO3}^{Si}}{v_{TO3}^{refSiO2} - v_{TO3}^{Si}}. \quad (5)$$

Knowing the value for $v_{TO3}^{Si}$ (960 cm$^{-1}$ as measured by FTIR on Si substrate), and measuring $v_{TO3}^{refSiO2}$ and $v_{TO3}^{SRSO_{-Er}}$ for each temperature of deposition (see Fig. 2), one can obtain the value

![Fig. 2 FTIR absorption spectra of the as-deposited samples for all T$_d$. The blue line indicates the shift of the TO$_3$ peak toward high wavenumber when T$_d$ is raised. Spectra were vertically shifted for clarity.](image-url)
of \( y \) for each \( T_d \). With this value \( y \) measured by FTIR, we obtain the atomic percentage of “isolated” silicon atom by the following formula:

\[
\% \text{Si}_{\text{isolated}} = 100 \times \frac{1 - y}{2 \left( 1 + y \right)}. \tag{6}
\]

Note that in Ref. 14, the estimated Si-excess represents a percentage of deviation from the stoichiometry of SiO\(_2\) and not an atomic percentage.

Any difference between values obtained by EDX and FTIR should thus be linked to an agglomeration phenomenon. Therefore, the comparison of the amounts of Si excess obtained by EDX and FTIR enables the determination of the fraction of agglomerated Si that can be estimated from the following relation:

\[
\text{SiO}_x \rightarrow \frac{x}{y} \text{SiO}_y + \frac{y-x}{y} \text{Si}. \tag{7}
\]

The fraction of agglomerated Si is thus obtained by the following definition:

\[
\% \text{Si}_{\text{agglomerated}} = 100 \times \frac{y-x}{1+x}, \tag{8}
\]

which verify the equation

\[
\% \text{Si}_{\text{Total Excess}} = \% \text{Si}_{\text{agglomerated}} + \% \text{Si}_{\text{isolated}}. \tag{9}
\]

The evolution of this fraction against \( T_d \) [Eq. (8)] is compared in Fig. 3 to that of the total Si excess, as determined by EDX [Eq. (2)].

It is noteworthy here that we are dealing with the atomic percentage of Si excess whose atoms are linked to each other within Si agglomerates. Such a fraction is obviously different from the number of sensitzers or Si-nc that should be lower than the percentage of Si atoms bonded only to other Si atoms. As expected, the fraction of agglomerated Si-excess is always lower than the total excess-silicon, while both show a clear increase when \( T_d \) is raised beyond 300\(^\circ\)C until reaching a maximum for \( T_d = 500\)\(^\circ\)C. The values of both parameters join at about 10\% for the highest \( T_d \) value (700\(^\circ\)C). At this stage, several observations can be made: 1. increasing the deposition temperature leads to a progressive agglomeration of the Si-excess, 2. the difference

![Fig. 3 Compared evolution of the total excess-silicon as measured by EDX (empty circles) and the agglomerated excess-silicon (full square).](image-url)
Fig. 4 Compared evolution of the agglomerated fraction of Si normalized to the total Si-excess for RT-deposited (squares) and 500°C-deposited (circles). Inset: agglomerated-Silicon atoms in atomic percent.

between total Si-excess and agglomerated Si-excess is reduced more and more as \( T_d \) is raised, and 3. at \( T_d = 700^\circ C \), the agglomerated Si reaches the value of the total excess silicon, indicating a complete phase separation between Si and SiO\(_2\).

This method can also be used to estimate the evolution of the fraction of agglomerated Si excess as a function of the annealing temperature \( T_a \). Two \( T_d \) values (RT and 500°C) were chosen to examine the evolution of the fraction of agglomerated Si, as depicted in Fig. 4.

It can be seen in the inset of Fig. 4 that the agglomeration of Si is progressively raising in function of \( T_d \) for both kinds of samples, until reaching their highest value after 1100°C. Indeed, after annealing at 1100°C, both kinds of samples (\( T_d \) for RT and 500°C) reach their limits, 7 at.% and 13 at.%, respectively, corresponding to the complete agglomeration of total Si-excess (see inset Fig. 4). The absolute value of % Si\(_{agglo}\) in at.% for RT-deposited is, for each annealing temperature, twice lower than the corresponding one for 500°C-deposited. This value happens to be governed by the total Si excess content (see Fig. 3).

To compare these two behaviors, we normalize these % Si\(_{agglo}\) (at.%) to their respective total Si-excess (7% for RT; 13% for 500°C) to obtain the agglomerated fraction (Fig. 4). Both curves show a similar trend toward \( T_a \), with a progressive increase of agglomerated Si and they eventually reach the value of 100% after 1100°C-annealing. This does not mean that all excess-Si atoms are located within a Si-nc but any excess Si atom is bonded to at least another Si atom. We remark that the fraction of agglomerated Si is always lower for RT-deposited sample because of its lower as-deposited value of agglomeration compared to the corresponding one for 500°C-deposited.

3.2 Cathodoluminescence Spectra

The CL technique was used for a tentative identification of the optically active centers present within our films. Indeed, since the incident electron beam is able to excite over a wide range of energy, the CL method is able to provide a “mapping” of the different emitting centers.\(^{22}\) It is known that even pure SiO\(_2\) presents CL emissions originating from some radiative-defect centers generated by the imperfect local arrangement of atoms, hence deviating from the perfect SiO\(_4\) tetraedra.\(^{23}\) Therefore, to analyze our SRSO:Er samples, it appears essential to compare their CL characteristics to those of sputtered SiO\(_2\), SiO\(_2\):Er, and SRSO samples. The corresponding CL spectra recorded in the visible range are displayed in Fig. 5.
We can see that the sputtered SiO₂ layer, considered as a reference here, present a broad CL spectra between 300 and 800 nm. This large band is the sum of different contributions, such as oxygen vacancies [in this case oxygen deficient center (ODC)] and nonbridging oxygen hole centers (NBOHC). This kind of CL spectrum was already observed by other research groups and recently by our group. When excess silicon is introduced, as for a silicon-rich silicon oxide (SRSO) sample, the corresponding CL spectrum presents two peaks centered at 450 and 480 nm, typically due to ODC. The Si excess in this material is liable to favor the creation of Silicon-ODC (SiODC), but seems unfavorable for the formation of NBOHC that are, indeed, not detected here. When Er ions are incorporated, as for the SRSO:Er sample, the two SiODC peaks are still emitting at the exact same wavelengths, in addition to two peaks at 520 and 550 nm. As the only difference between this sample and SRSO is the presence of Er³⁺, these peaks are likely to originate from Er³⁺ levels’ transitions. This hypothesis is further confirmed by the CL spectra of SiO₂: Er that presents the same transitions that are still distinguishable in spite of the wide overlapping of SiODC contributions. One can, therefore, argue that these two Er-related emissions are respectively arising from the \( ^2H_{11/2} \rightarrow ^4I_{15/2} \) and the \( ^4S_{3/2} \rightarrow ^4I_{15/2} \) transitions.

Another important feature deserves to be noted: the Er-related emissions are predominant in SRSO:Er sample, at the expense of the SiODC contributions. Indeed, these SiODC emissions in a SRSO:Er are about two orders of magnitude lower than the corresponding ones in a SiO₂:Er sample free from Si excess. This counterbalancing phenomenon is reminiscent of the earlier mentioned and well known coupling between the Er³⁺ ions and the Si-based sensitizers. One can therefore consider that we are dealing with a similar energy transfer from SiODC toward Er³⁺ ions, especially as this is accompanied by a drastic decrease of the emissions from SiODC. These emitting centers appear very similar to the so-called Luminescent Centers of Savchyn et al., described as oxygen vacancies and which play a predominant sensitizing role toward Er³⁺ ions. Besides, such an energy transfer was already shown in SRSO from Si-based-entities to neodymium ions, these Si-based entities have absorption energies very close to those of SiODC. We should also note that a so-called “inverse energy transfer” from Er³⁺ ions toward Germanium oxygen deficient centers (GeODC) was reported by Kanjilal et al., indicating that energy transfer between erbium and ODC is possible.

To check the evolution of all emitting centers in function of both \( T_d \) and \( T_a \), we made specific CL analyses on our samples. Figure 6 reports typical evolution of the CL spectra...
First of all, we can see that all samples present the 1.5 μm Er3+ emission due to $^4I_{13/2} \rightarrow ^4I_{15/2}$ radiative transition, which similarly evolves for both RT-deposited and 500°C-deposited samples: an intensity increase by a factor of 3–4 until an optimum $T_a = 600^\circ$C, followed by a strong decrease after annealing at 1100°C. Note that the maximum CL intensity (after 600°C annealing) for the RT-deposited sample is twice higher than that for 500°C-deposited one, which can be partly explained by the 20% less Er in the latter sample (see Fig. 1). But the most significant difference between the two kinds of samples is that the 500°C-deposited sample contains a total Si-excess twice higher than the RT-deposited one. Thus, the origin of the lowering of Er3+ emission at 1.5 μm may be linked to this difference.

By comparing the two CL spectra in the visible range [left panels of Figs. 6(a) and 6(b)], one can observe the following: 1. the Er-related emissions ($^2H_{11/2} \rightarrow ^4I_{15/2}$ and $^4S_{3/2} \rightarrow ^4I_{15/2}$) from the RT-deposited samples are 80–90% higher than those from the 500°C-deposited ones, 2. the CL contribution assigned to quantum-confinement in Si-nc and observed in 500°C-deposited samples (at $\sim 650$ nm for $T_a = 900^\circ$C and at $\sim 720$ nm for $T_a = 1100^\circ$C) is absent for RT-deposited samples, and 3. the visible emissions from Er3+ ions (520 and 550 nm) are still detected after annealing at 1100°C of RT-deposited sample, whereas they are not distinguishable for the 500°C-deposited one. These observations well confirm that the weak emission of erbium in 500°C-deposited samples (both in Vis and IR range) is concomitant with a significant emission
from Si-nc, whereas the RT-deposited samples present high emission from Er$^{3+}$ but no Si-nc emission. We thus argue that the Si-nc influence the CL of Er$^{3+}$ ions, in a way that differs from the PL experiments dealing with the usual energy transfer from Si-nc toward Er that enhances the Er$^{3+}$ PL. The picture here is more complex as the layers are excited by hot electrons, thus enabling the simultaneous excitation of SiODC, Si-nc, and Er$^{3+}$ ions. These concurrent impact excitations are likely to provoke simultaneous energy transfer from Si-nc toward Er$^{3+}$ ions as well as energy back-transfer from Er$^{3+}$ ions toward Si-nc, thus reducing the overall emission from Er$^{3+}$ ions.

On the other hand, the SiODC contribution becomes predominant in CL spectra for both RT- and 500°C-deposited samples after a 1100°C-annealing. In both cases, the huge increase of SiODC emission is concomitantly occurring with a strong decrease in the Er$^{3+}$ ions emission at 1.5 μm. This counterbalancing behavior of the emissions from SiODC and Er ions is indicative of the possible active role of SiODC as sensitizers for Er$^{3+}$ ions. In addition, the emission from those point-defect centers in the visible range overlaps with the Si-nc absorption band, and this strongly suggests that they may also act as sensitizers for Si-nc. The highest emission of SiODC (after 1100°C-annealing) occurs when the Si excess is completely agglomerated (see Fig. 4), indicating that the SiODC emission is not due to isolated Si atoms. Besides, at this high annealing temperature providing a complete phase separation between Si and SiO$_2$, we assume that SiODC are mainly located at the substoichiometric interface between Si-nc and SiO$_2$ matrix.

These different processes of energy transfer, whether demonstrated or likely to occur in CL experiment, are sketched in Fig. 7 with a proposed scheme of excitation, together with the band diagrams of all considered emitting centers (SiODC, Si-nc, and Er$^{3+}$ ions).

These CL results give very valuable insights on the presence of all the emitting centers within SRSO:Er layers, as well as the possible processes of energy transfer between the active

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**Fig. 7** Schematic band diagrams of all detected emitting centers within SRSO:Er layers. In the upper right part a model of interaction between luminescent entities is depicted.
emitting centers. To distinguish between these processes, we carried out PL experiments, as complementary to CL studies.

3.3 Photoluminescence Spectra

In this part, all PL spectra were obtained using an excitation wavelength (476 nm) that is nonresonant for Er$^{3+}$ ions and is well beyond the excitation threshold wavelength for SiODC. This ensures that all emitting Er$^{3+}$ ions are excited by only Si-nc, ruling out any eventual energy transfer from SiODC to Er$^{3+}$ ions. In this way, the following interpretations for the energy transfer would be much simpler. Note that an interference pattern is observed for all of the following PL spectra of samples in the range 600–900 nm, because of the reflections occurring at the interfaces of these relatively thick layers.30

In order to thoroughly analyze the evolution of the Si-PL emission of SRSO:Er versus $T_a$, a first study was carried out for PL emission of SRSO. Figure 8 shows the evolution of Si-nc-PL emission for SRSO layer free from any Er$^{3+}$ ions. It can be seen that the influence of the annealing temperature on Si-nc PL emission is not straightforward. The PL intensity is slightly increased from as-deposited to 600$^\circ$C annealing, followed by a decrease after 900$^\circ$C annealing and eventually a fifty-time enhancement after 1100$^\circ$C-annealing.

Since no significant shift is observed in the wavelength of the PL spectra for as-deposited and 600$^\circ$C-annealed samples, we assume that, although more optically active Si-nc are formed with annealing, their average size remains unchanged. On the contrary the annealing from 600 to 900$^\circ$C leads to a slight shift of the PL spectra, together with a global decrease of the PL intensity. This might be due to some increase of the average size at the expense of their density. This latter phenomenon is strongly increased after 1100$^\circ$C annealing, giving rise to well crystallized Si-nc that were observed by electron microscopy (not shown), responsible of the strong enhancement of Si-nc emission.

Figure 9 displays the PL spectra (normalized to the thickness and to the total Si-excess percentage) in the visible range for both RT- [Fig. 9(a)] and 500$^\circ$C-deposited [Fig. 9(b)] samples, subsequently annealed at the indicated temperatures. This visible PL is assumed to originate from the Si-nc not coupled to Er$^{3+}$ ions, and then not concerned by the energy transfer. Thus, the evolution of the visible band with $T_a$ can be due to a variation in the number of emitting Si-nc and also to an evolution of the coupling between Si-nc and Er$^{3+}$. For the as-deposited [blue lines in Figs. 8, 9(a), and 9(b)], we can see that the visible PL intensities for RT-deposited
samples of SRSO:Er and SRSO deposited at 500°C are comparable, whereas this intensity is twice lower for the SRSO:Er deposited at 500°C. This indicates that the energy transfer toward Er³⁺ ions is more efficient for this latter sample, reducing the overall visible PL band.

The evolution of Si-nc PL against \( T_d \) for both RT- and 500°C-deposited SRSO:Er samples, is qualitatively similar to that of SRSO. We thus assume that the evolution of Si-nc PL intensity is mainly governed by the density of Si-nc, and not by their coupling with Er³⁺ ions. In any case, the Si-nc PL intensity is lower for 500°C-deposited sample than for the RT-deposited one throughout the annealing temperature range, and this suggests that the energy transfer is always more efficient for 500°C-deposited sample.

Let us examine the evolution against \( T_d \) and \( T_a \) of the corresponding Er-PL that is governed by the excitation through the Si-nc.

Figure 9(c) displays the evolution with \( T_a \) of the Er-PL efficiency, defined as the ratio \( I_{Er^{PL}}^E/\text{[Er]} \), \( I_{Er^{PL}}^E \) being the Er-PL intensity. It can be seen that the PL efficiency evolves similarly for the two samples deposited at RT and 500°C. They show both a gradual increase up to a maximum for \( T_d = 900°C \), before collapsing together when \( T_d \) reaches 1100°C. This evolution with \( T_a \) is qualitatively similar to that of CL (Fig. 6), except for the optimum value of \( T_a \): 900°C for PL and 600°C for CL. As we are dealing with the same samples for both studies, one can assume that the different optimum \( T_a \) may be due to the different excitation sources, electrons for CL and photons for PL. To note, however, that this PL efficiency is systematically higher for the 500°C-deposited sample, which is quite consistent with earlier observations on samples as-deposited at different \( T_d \) values¹⁴ and with the above-mentioned lower Si-nc emission from 500°C-deposited samples. While the annealing at the optimum \( T_a = 900°C \) improves by a factor of four, the PL efficiency for the 500°C-deposited sample, the corresponding improvement for the RT-deposited layer is improved by a factor of 18, but still insufficient to reach the 500°C-deposited Er-PL efficiency. The increase of the PL efficiency is due to an enhanced coupling between the Er ions and the Si-nc sensitizers whose formation is increasingly favored by the annealing up to 900°C, as claimed above. The earlier reported
increase of the optically active ions with the thermal budget provided an additional support to our explanation.\textsuperscript{14} For the highest value of $T_d$ (1100°C), two processes compete to the observed drastic decrease of the PL efficiency: first, the tendency of Er to agglomerates\textsuperscript{31,32} second, the coalescence of Si-nc in large crystallized nanograins, as confirmed by the HRTEM observations of our present samples (not shown). It is worth noting that, although the samples deposited at two different temperatures show generally similar evolutions of Er-PL and Er lifetimes (not shown), the sample grown at 500°C shows much better Er emission for each annealing temperature. The origin of this performance is due to the fact that much more Si atoms are agglomerated during the growth, compared to RT deposited layer. As shown in Figs. 34, the fraction of agglomerated Si is increasing with $T_s$ and $T_d$. To have a deeper insight on the influence of this fraction on the sensitization of Er\textsuperscript{3+} ions, in Fig. 10 we display the evolution of the Er-PL efficiency in function of the fraction of agglomerated Si.

We can see that whatever the parameters, such as the total Si-excess and the temperatures of deposition and annealing, the Er-PL efficiency seems to be governed by the fraction of agglomerated Si. Note that the last point shows a collapse of the Er-PL efficiency that is explained by the agglomeration of Er\textsuperscript{3+} ions,\textsuperscript{31,32} occurring after a 1100°C-annealing. This result well demonstrates that the fraction of agglomerated Si is a key factor for the enhancement of the erbium-PL. Future studies will be needed to have a deeper insight on the agglomeration of Si, especially the size and density of Si-nc must be investigated to understand their respective influence on the Er-PL.

4 Conclusion

The optical and structural properties of Er-doped silicon rich silicon oxide thin films grown by magnetron cosputtering were investigated in function of both deposition and annealing temperatures. The temperature of deposition was shown to have several crucial effects on the properties of the deposited thin film. When the deposition temperature is raised, an increase of the fraction of agglomerated Si is observed concomitantly to a lowering of the Er content. Besides, the annealing temperature was shown to increase the fraction of agglomerated Si until complete phase separation at 1100°C, regardless of the deposition temperature and the total Silicon-excess content. The cathodoluminescence technique was used to investigate the emission of optically-active centers. It was shown that silicon-oxygen deficient centers emit around 450–500 nm in all kinds of samples and are likely to contribute to the energy transfer toward both Si-nc and Er\textsuperscript{3+} ions. To rule out any influence of the SiODC on the coupling of Er\textsuperscript{3+} ions, the nonresonant 476 nm wavelength was used for photoluminescence...
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experiments. It was shown that the Er-PL efficiency is systematically higher for the 500°C-deposited samples compared to the RT-deposited ones. It was also observed that the fraction of agglomerated-Si governs the Er-PL efficiency, whatever the value of deposition or annealing temperature.

References


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