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# Impact of cerium and lanthanum on the photo-darkening and photo-bleaching mechanisms in thulium-doped fibre

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## A B S T R A C T

Photo-darkening prevents developing of new applications of rare-earth doped silica fibre lasers or amplifiers operating at wavelengths shorter than  $1\ \mu\text{m}$  or at any wavelength at high power. The photo-darkening is characterized by a decrease of the laser intensity during amplification. In this article, we are interested in *Tm*-doped fibres. This ion offers many potential optical transitions spanning from  $0.45$  to  $1.9\ \mu\text{m}$ . Several interesting transitions can be excited via up-conversion, using one pump around  $1\ \mu\text{m}$ . However, this scheme induces a particularly fast and intense photo-darkening, enough to prevent amplification in the short wavelengths region ( $0.45$ – $0.9\ \mu\text{m}$ ). To mitigate this effect, we co-dope the fibres with cerium or lanthanum ions. By characterizing the steady state values of photo-darkening and its characteristic times, we demonstrate the beneficial role of *Ce* and *La* on the bleaching mechanisms.

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**Keywords:**  
Optical fibre  
Thulium  
Photo-darkening  
Silica

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## 1. Introduction

Silica as a glass host for rare-earth (RE) doped fibre lasers and amplifiers offers the best performances in terms of efficiency, power, reliability and cost effectiveness. It is indeed chemically stable, mechanically robust and its optical transparency is high in the visible and the near-infra-red (NIR) range up to  $2\ \mu\text{m}$ . Then, silica is the main glass used, for example, in high power fibre lasers [1]. In the case of *Yb*-doped fibre, this technology reaches kilowatt level in single mode and continuous wave regime. Presently, one of the main issues related to the output power is related to the photo-darkening mechanism. It is characterized by the presence of a broad absorption band in the visible and NIR. It causes the decreasing of the laser power versus time. This induced loss has a significant impact on the threshold and the laser slope efficiency [2]. In addition, photo-darkening is suspected also to trigger mode instability, leading to a degradation of the *M* factor of these lasers [3–6].

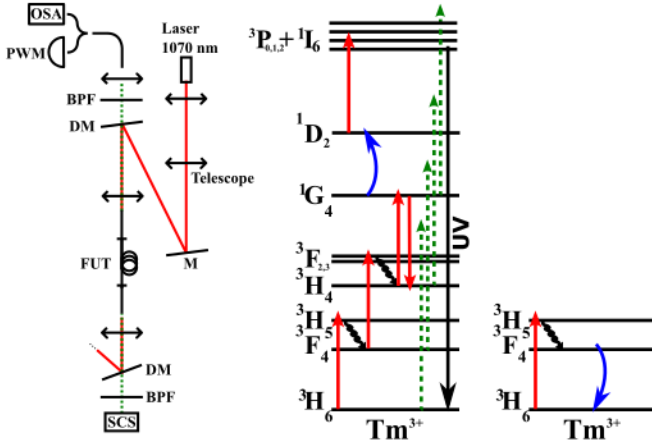
For the development of fibre lasers at wavelengths shorter than  $1\ \mu\text{m}$ , thulium ( $Tm^{3+}$ ) is particularly interesting because it offers many potential optical transitions spanning from  $0.45$  to  $1.9\ \mu\text{m}$ .

Except for the conventional pumping scheme at  $0.79\ \mu\text{m}$  which provides efficient amplification around  $1.9\ \mu\text{m}$  through a down-conversion energy transfer process [7], none of the other transitions are exploited in silica. An interesting up-conversion pumping scheme using a pump source at  $1.07\ \mu\text{m}$  provides up to seven laser emissions (spanning from  $0.45$  to  $1.9\ \mu\text{m}$ ) that were implemented in ZBLAN fibres [8]. If one applies this scheme to a *Tm*-doped silica fibre, two phenomena will hamper amplification: fluorescence quenching by non-radiative decay and transparency degradation by photo-darkening [9,10]. The high phonon-energy of silica glass (as compared with fluorides) induces fast non-radiative decays from most energy levels of thulium, causing a strong reduction of their effective lifetimes and hence the reachable population inversion, even under high pumping. In *Tm*-doped silica, the non-radiative decay from the  $^3H_4$  excited level may be mitigated by high *Al* co-doping, promoting the amplification of the  $0.8\ \mu\text{m}$  and  $1.47\ \mu\text{m}$  emission bands as well as up-conversion to higher energy levels (see Fig. 1 right) [9,11,12]. Because the sought applications necessitate high concentrations of aluminium and thulium, strong photo-darkening is observed [10,13]. Besides, pumping of RE-doped optical fibres may cause the degradation of the transparency of both silica and fluoride glasses, especially when they are heavily doped with RE ions [10,13–17]. In the case of thulium-doped silica fibre pumped at  $1.07\ \mu\text{m}$ , photo-darkening (or photo-induced

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**Fig. 1.** Left: Experimental setup for PIA measurement. M: dielectric mirror, FUT: fiber under test, BPF: band pass filter, DM: dichroic mirror, OSA: optical spectrum analyser, SCS: supercontinuum source, PWM: power meter. Right: Energy level diagram of  $Tm^{3+}$ ; red: pump induced transitions, blue: energy transfer, green:  $0.55 \mu m$  probe. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

attenuation, PIA) is particularly fast and intense, enough to prevent amplification in the extended-visible region ( $0.45\text{--}0.9 \mu m$ ) [10]. The beneficial role of co-doping with cerium or lanthanum ions to mitigate the steady states of photo-darkening has been reported previously [18]. The aim of this article is to study both the steady state of photo-darkening and its characteristic time to discuss the role of Ce and La on the generation and bleaching mechanisms.

## 2. Samples preparation

Preforms were fabricated by the conventional Modified Chemical Vapor Deposition (MCVD) technique [19] and the so-called solution doping technique [20] was applied to incorporate aluminium (Al), thulium (Tm), cerium (Ce) and lanthanum (La) ions. To fabricate a preform, a porous silica layer was deposited inside substrate silica tube. After soaking of the layer with ionic solutions of chloride salts, the solvent was dried and the core layer was sintered down to a dense glass layer. Then the tube was collapsed into a solid rod, referred to as preform, at an elevated temperature higher than  $1800^\circ C$ . Preforms were stretched into  $125 \mu m$  fibres using a drawing tower at temperatures higher than  $2000^\circ C$  under normal conditions. Three series of samples were prepared with various concentrations of thulium, lanthanum and cerium ions. The concentrations of Al, Tm, Ce and La were measured by EPMA (Electron Probe Micro-Analysis) in the fibres. All the concentrations are given in atomic ppm and reported in Table 1. In all samples, the concentration of aluminum remained almost constant, typically around  $8000 \text{ ppm}$ . In the Tm series, the concentration of thulium spans from 0 to  $600 \text{ ppm}$ . In the La series, the concentration of lanthanum varies from 350 to  $7000 \text{ ppm}$  whereas the concentrations of thulium remains almost constant ( $190 \pm 30 \text{ ppm}$ ). In the Ce series, the concentration of cerium varies from 0 to  $1300 \text{ ppm}$  whereas the concentration of thulium remains almost constant ( $260 \pm 40 \text{ ppm}$ ).

**Table 1**  
Concentrations of all elements incl. oxygen (ppm.at).

Series	[Tm]	[La]	[Ce]	[Al]
Tm	0–600	–	–	~ 8000
La	$190 \pm 30$	0–7000	–	~ 8000
Ce	$260 \pm 40$	–	0–1300	~ 8000

## 3. Experimental setups

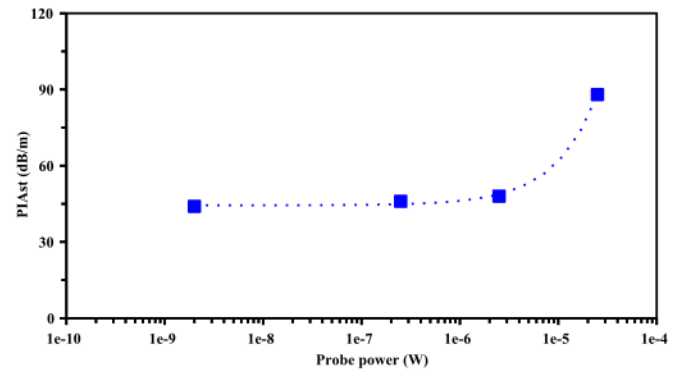
Photo-darkening, fluorescence lifetime and fluorescence measurements were performed in this study.

### 3.1. Photo-darkening

Fig. 1 describes the experimental setup used to measure the PIA. The pump laser at  $1.07 \mu m$  is a continuous wave Yb-doped fibre laser (Keopsys, KPS-CUS-OEM-1064). It was coupled into a commercial passive fibre (Corning, HI-1060, referred to as “input fibre”) through a telescope, a high reflection dielectric mirror, a dichroic mirror and an aspherical lens. The dichroic mirror is highly reflective at  $1.07 \mu m$  and transparent from  $0.55$  to  $1.0 \mu m$ . The input fibre end was cleaved at right angle. The fibre under test (FUT) samples were spliced to the input fibre. The typical length of samples was about 2 cm, short enough to neglect the pump depletion (loss  $< 1 \text{ dB}$  at  $1.18 \mu m$ ). Another passive HI-1060 fibre was spliced at the other end of the FUT. A super continuum source (SCS, Leukos SM-30) was coupled into the passive fibre in the counter-propagative direction relative to the pump beam, using two metallic mirrors (not represented) and an aspherical lens. The remaining pump was rejected by a second dichroic mirror, so that the SCS was protected against laser damage. Two band pass filters at  $0.55 \mu m$  with a  $2 - nm$  bandwidth were placed before and after the FUT. Their extinction ratio is higher than  $40 \text{ dB}$ . Next, the green beam is called the probe.

The injected pump was measured from the end of the input fibre by a power meter before the splicing of the FUT. The time-resolved PIA measurement was recorded at  $0.55 \mu m$  while pumping at  $1.07 \mu m$ . This probe wavelength was selected because the PIA is strong whereas there is no absorption band of  $Tm^{3+}$  ions at  $0.55 \mu m$ . The band pass filters were placed, the SCS was turned on, and the PWM (Si photo-diode) was used to continuously measure the transmitted probe while pumping. The recording was continued after the laser was turned off, in order to measure bleaching dynamics caused by the probe.

The probe power can affect the measurements. For example, Fig. 2 shows steady state values of photo-induced attenuation for the Tm-doped sample containing the highest concentration of Ce ( $1300 \text{ ppm}$ ) as a function of the probe power. The steady state values of the PIA ( $PIA_{st}$ ) span from  $44 \text{ dB/m}$  to  $90 \text{ dB/m}$  when the probe power increases from  $2 \text{ nW}$  up to  $25 \mu W$ . The same trend was observed in the La series. To minimize the effect of the probe in the case of Ce or La doped samples, it is necessary to use a probe power lower than  $\sim \mu W$ . Therefore photo darkening



**Fig. 2.**  $PIA_{st}$  values obtained for various probe powers spanning from  $2 \text{ nW}$  to  $25 \mu W$ , for pump power of  $1 \text{ W}$  measured in the  $1300 \text{ ppm}$  cerium-doped sample. Dot: experimental values, dotted line: linear fitting.

measurements should be performed at these low probe power in order to quantify only the effect of the pump and rare-earth ions concentrations. In the case of singly  $Tm$ -doped samples, we have checked that a higher probe power ( $25 \mu W$ ) does not disturb the measurements.

### 3.2. Fluorescence spectra

The experimental setup used to measure the fluorescence spectra of  $Tm^{3+}$  is described in Fig. 1. The pump wavelength was  $1.07 \mu m$ . For these measurements, the SCS and the filter were not used. The contra-propagative fluorescence was collected and measured by an optical Fourier transform analyser (Thorlabs OSA-201C). The spectra were acquired from  $400 nm$  to  $1 \mu m$  with different pump powers. All spectra present a parasitic peak at  $632.8 nm$  which is due to the intrinsic  $He-Ne$  reference inside the device. When the pump power is lower than  $250 mW$ , photo-darkening can be neglected for few minutes. When the pump power is higher than  $500 mW$ , the intensity of the emission spectrum is impacted by the photo-darkening particularly in the short-wavelengths domain.

### 3.3. Fluorescence lifetimes

Fig. 3 describes the experimental setup used to measure the fluorescence lifetime of the  ${}^3H_4$  level of  $Tm^{3+}$ . The pump laser was a fibre coupled laser-diode emitting at  $785 nm$ . A 50/50 coupler was used to collect the backward fluorescence. The fibre under test was spliced on one of the coupler arms. Both ends are immersed in index liquid to prevent any reflection of parasitic light. The contra-propagative fluorescence was collected, then filtered by a spectrometer tuned to  $810 nm$  (resolution  $\sim 0.1 nm$ ). The output light was detected by an amplified avalanche silicon photodiode (APD, EG&G SPCM AQR-14-FC) operated in the photon-counting mode. The TTL electrical pulses from the APD were counted by a Stanford SR400 photon counter synchronized by the laser diode modulation signal. Decay curves were registered using a time-gate scanning across one pump modulation period. In order to minimize errors (laser fluctuations,...) and increase the S/N ratio, the signal was normalized in real time by the signal from a fixed time-gate integrating the signal over a full  $1 - ms$  modulation period, and each data point was averaged 5000 times. A decay curve contains typically 500 data points.

## 4. Rate equation model

In this section, we present the model used to discuss the darkening and the bleaching phenomena. It is well-known that light can be absorbed by precursors present in the glass, leading to the formation of color-centres absorbing in the visible. In silica, many different color-centres have been identified. For the sake of simplicity in this model, we consider the formation of one generic

color-centre which can be photo-bleached. In this case, the normalized density  $n$  of color-centres is ruled by the generation ( $g$ ) and the bleaching ( $b$ ) rates. The time derivative of the number of color-centres  $\dot{n}$  has two contributions: (i)  $g \cdot (1 - n)$  which is the generation term requiring an exhaustible source, (ii)  $-b \cdot n$  which is the bleaching term requiring a color-centre to bleach. These considerations lead to the differential equation (1).

$$\dot{n} = g \cdot (1 - n) - b \cdot n \quad (1)$$

According to the Beer-Lambert law, the generation of one color-centre, characterized by the absorption coefficient  $\alpha_{cc}$ , leads to the photo-induced absorption  $PIA = \alpha_{cc} \cdot n$  (in  $dB/m$ ). The resolution of equation (1) gives the temporal form of  $PIA(t)$ :

$$PIA(t) = \alpha_{cc} \cdot \frac{g}{g+b} \left( 1 - e^{-(g+b) \cdot t} \right) \quad (2)$$

Equation (2) exhibits two parameters:  $PIA_{st} = \alpha_{cc} \cdot \frac{g}{g+b}$  and  $\tau_{1/e} = \frac{1}{g+b}$  which are the steady state value of  $PIA$  and the characteristic time, respectively. Both parameters can be determined experimentally. Reversed relations are:  $\alpha_{cc} \cdot g = \frac{PIA_{st}}{\tau_{1/e}}$  and  $(g+b) = \frac{1}{\tau_{1/e}}$ . As the value of  $\alpha_{cc}$  is unknown,  $g$  and  $b$  cannot be determined separately.

The measurements of  $PIA_{st}$  and  $\tau_{1/e}$  allow to discuss on the increase or the decrease of the photo-darkening and the photo-bleaching rates. Indeed, when  $PIA_{st}$  increases and  $\tau_{1/e}$  decreases, the phenomenon is ruled by the increasing of  $g$ . When both  $PIA_{st}$  and  $\tau_{1/e}$  increase, the phenomenon is ruled by the decreasing of  $b$ . When both  $PIA_{st}$  and  $\tau_{1/e}$  decrease, the phenomenon is ruled by the increasing of  $b$ . Finally, when  $PIA_{st}$  decreases and  $\tau_{1/e}$  increases, the phenomenon is ruled by the decreasing of  $g$ . Table 2 synthesises the link between the measured parameters ( $PIA_{st}$  and  $\tau_{1/e}$ ) rates of photo-darkening and photo-bleaching.

This model is based on the assumption that only one kind of color-centre is implied. To discuss this statement, Fig. 4 shows one typical decay curve of the  $PIA$  fitted with a single, a double and a stretched exponential. The experimental curve was measured in an  $Al-Tm$  doped sample pumped at  $1 W$  and with a probe at  $550 nm$  (power:  $25 \mu W$ ).

The best curve fittings correspond to double and stretched exponential cases. Stretched exponentials are commonly used to describe non-exponential decay curve, but there is no easy interpretation of the stretch parameter. The two-exponentials fitting can be explained by the formation of two color-centres. However, the experimental values of interest ( $PIA_{st}$  and  $\tau_{1/e}$ ) are well fitted even with the single exponential decay curve. The single exponential curve fails to fit the intermediate part, which is out of interest in this study. For these reasons, we consider here the formation of one generic color-centre.

## 5. Results

### 5.1. Effect of thulium concentration on the photo-darkening

In this subsection, we consider the  $Tm$  samples series. The effect

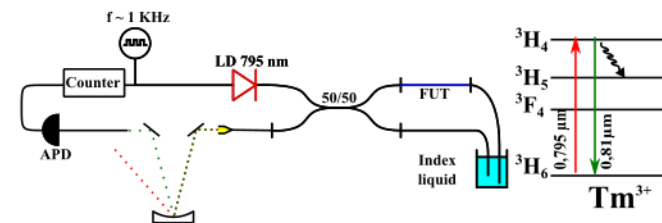
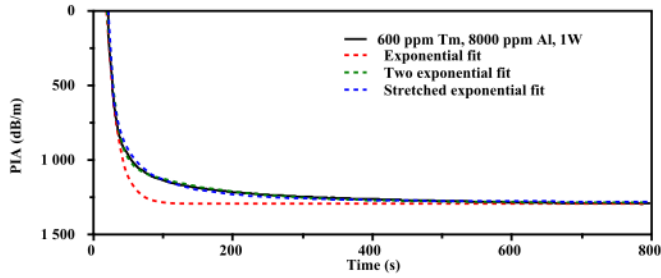


Fig. 3. Experimental setup used to measure the fluorescence lifetime of the  ${}^3H_4$  level of  $Tm^{3+}$ . APD: avalanche photo-diode. FUT: fibre under test.

Table 2

Effects of the increase and decrease of  $PIA_{st}$  and  $\tau_{1/e}$  on the generation and bleaching rates.

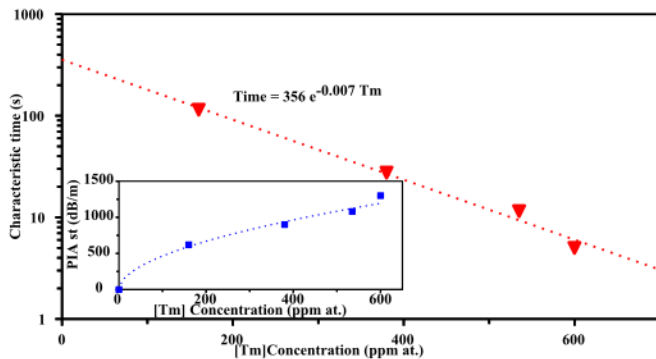
$PIA_{st}$	↑	↑	↓	↓
$\tau_{1/e}$	↓	↑	↓	↑
Ruling rate	$g \uparrow$	$b \downarrow$	$b \uparrow$	$g \downarrow$



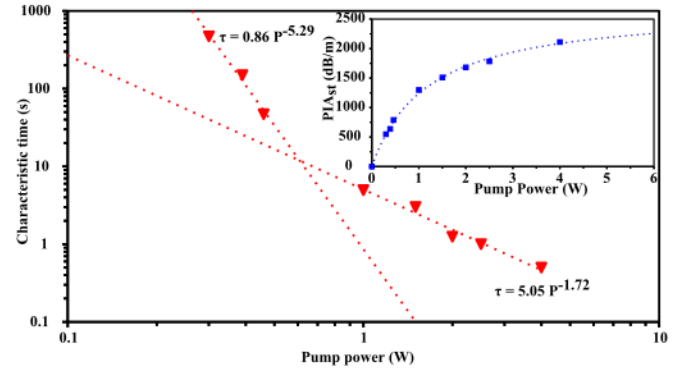
**Fig. 4.** Time resolved PIA for a sample containing 600 ppm Tm and 8000 ppm Al and three fitting curves. Pump power: 1 W, probe power: 25  $\mu$ W.

of thulium concentration on the  $PIA_{st}$  and the characteristic time  $\tau_{1/e}$  is reported in Fig. 5. As reported previously, when Tm concentration increases from 0 to 600 ppm,  $PIA_{st}$  increases from 0 to 1300 dB/m. The data can be fitted with a power law, with an exponent equals to 1/2 for the dependence on Tm concentration (dotted line in the inset of Fig. 5) [18]. The same trend was observed in Yb-doped silica with varying Tm concentration and probe light at 633 nm [13]. The characteristic time decreases from 115 to 6 s when Tm concentration increases from 160 to 600 ppm. For this range of Tm concentration, the characteristic time can be fitted with an exponential law (dotted line in Fig. 5). At concentrations lower than 150 ppm, it is expected that  $\tau_{1/e}$  behaviour deviates from this exponential law. Indeed, when Tm concentration tends to 0,  $PIA_{st}$  tends to 0 and  $\tau_{1/e}$  tends to infinity. According to the data reported in Fig. 5, in the 200–600 ppm range,  $\tau_{1/e}$  is very sensitive to the Tm concentration as it varies by a factor 30 while the  $PIA_{st}$  varies by a factor 2 only.

The inset of Fig. 6 reports the  $PIA_{st}$  values for the sample doped with the highest content of Tm and for pump power ( $P$ ) varying from 300 mW to 4 W.  $PIA_{st}$  values increase with the pump power. These data can be fitted with this law:  $PIA_{st} = PIA_{st}^{sat} \frac{A \cdot P}{1 + A \cdot P}$  where  $A = 0.86 \text{ W}^{-1}$  and  $PIA_{st}^{sat} = 2700 \text{ dB/m}$ . The variation of the characteristic time  $\tau_{1/e}$  versus  $P$  is reported in Fig. 6. These values are fitted with a power law whose exponent is  $-5.29$  and  $-1.72$  when pump power is lower or higher than 0.7 W, respectively. In the case of Yb-doped silica fibres, a power-law dependence was reported and the exponent was associated to the number of photons implied in the photo-darkening process [21]. Based on this consideration, at low pump power, the phenomenon would be ruled by the absorption of 5 or 6 photons while at higher pump power only 1 or 2 photons would be involved. These two processes can be tentatively



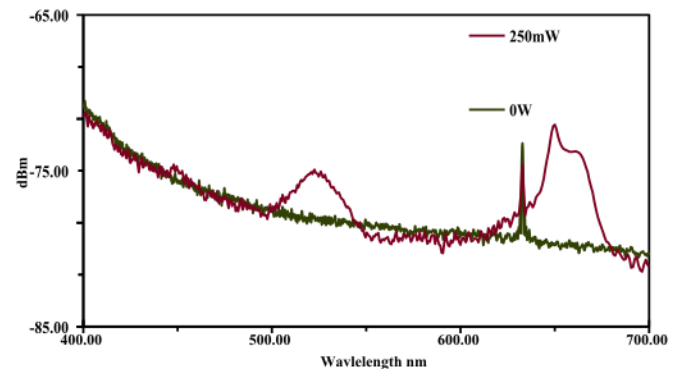
**Fig. 5.** Characteristic time for various thulium concentrations, from 0 to 600 ppm and 8000 ppm Al. Measured at 1 W pump power and 25  $\mu$ W of 550 nm probe. Dot: experimental data, dotted lines: exponential fitting. Inset:  $PIA_{st}$  corresponding, Dot: experimental data, dotted line: square-root fitting.



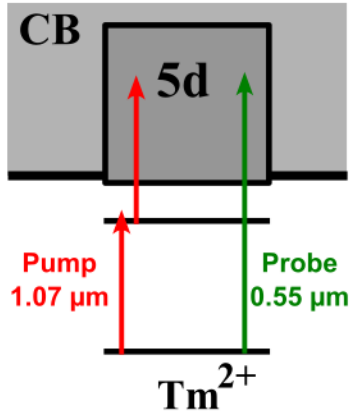
**Fig. 6.** Characteristic times for the 600 ppm Tm and 8000 ppm Al sample, for pump powers spanning from 300 mW to 4 W, measured at 550 nm. Dot: experimental data, dotted lines: power law fittings. Inset:  $PIA_{st}$  values for the same measurements. Dot: experimental data, dotted lines: fitting with saturate linear.

interpreted as follows.

For the sample with the highest Tm concentration, the emission spectra were measured under 1.07  $\mu$ m pumping scheme with pump power varying between 0 and 4 W. A characteristic emission spectrum is reported in Fig. 7 and corresponds to the pump power of 250 mW. For higher pump power, the emission spectra in the visible domain are difficult to measure due to the photo-darkening effect. The emission spectrum exhibits three emission bands at 470, 530 and 650 nm. These emission bands are attributed to  $Tm^{3+}$  ions. Blue and red emissions can be attributed to de-excitations from the  $^1G_4$  manifold. This level is populated through up-conversion mechanism (3-photon absorption) described in Fig. 1. The green emission band is emitted from energy levels higher than the  $^1G_4$ . The  $^1D_2$  level can be populated through energy transfer between two  $Tm^{3+}$  ions: a  $Tm^{3+}$  ion absorbs a pump photon to reach the  $^3H_5$  level and decays non-radiatively to the  $^3F_4$  level. Then, the energy transfer ( $^3F_4, ^1G_4$ )  $\rightarrow$  ( $^3H_6, ^1D_2$ ) takes place as already observed in fluoride glass [22]. From the  $^1D_2$  level, the  $Tm^{3+}$  ion can absorb another pump photon to reach the  $^3P_{0,1,2}$  and  $^1I_6$  levels. According to these processes, reported in Fig. 1, the absorption of 5 pump photons leads to the emission of UV light. These UV photons are expected to be involved in the photo-darkening process because they can ionize precursor defects such as the Al-E' and NBOHC centres [14,23–25]. In this case, electrons are released in the conduction band and holes in the valence band. These charges can then recombine on traps to form color-centres: the AlOHC in the case of



**Fig. 7.** Visible-NIR fluorescence under 1.07  $\mu$ m pumping, for the sample doped with 600 ppm Tm and 8000 ppm Al. Res: 1 nm, peak at 632.8 nm: artefact from OSA.



**Fig. 8.** Schematic representation of bleaching mechanisms by  $Tm^{2+}$  ion. Red: pump photon, green: probe. CB: Conduction band. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

hole, or, in the case of electron, on a  $Tm^{3+}$  to form a  $Tm^{2+}$ .  $AlOHC$  and  $Tm^{2+}$  absorb in the visible [26]. As a conclusion, at low pump power, the variation of the characteristic time is associated to the generation of color-centres due to the UV emission induced by the 5-photon absorption mechanism of  $Tm^{3+}$  (see Fig. 8).

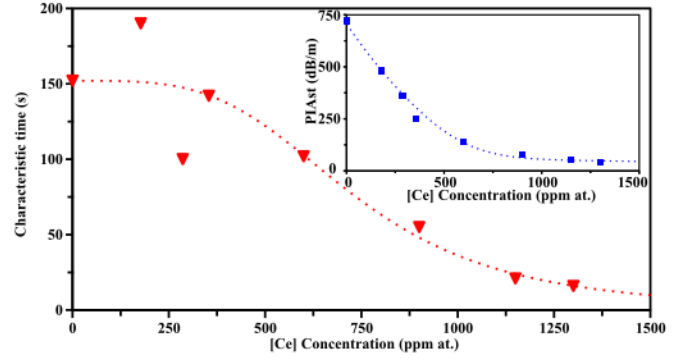
For pump power higher than 0.7 W, thulium ions undergo a similar effect as the induced transparency. Indeed, there is a competition between the absorption of pump photon on the  $^3H_4 \rightarrow ^1G_4$  transition and the stimulated emission of the opposite transition. Then, at high pump power, populations of the  $^3H_4$  and  $^1G_4$  levels are saturated and the ratio of these populations is the ratio of the absorption cross-sections and associated emission cross-sections. If these populations are saturated, it can be deduced that the pump has no more interactions with  $Tm^{3+}$  ions and therefore the generation of color-centres saturates.

For pump power higher than 1 W,  $PIA_{st}$  starts to saturate and the characteristic time evolves differently as the exponent of the power law is between 1 and 2. The ground level of the  $Tm^{2+}$  ions, initially present in the glass or induced by UV emission, is located close to the bottom of the conduction band ( $\sim 1.9$  eV) [27,28]. Moreover, the  $4f-4f$  transition of  $Tm^{2+}$  occurs at  $\sim 1$  eV [26,29,30]. Then, only one probe photon ( $\sim 2.25$  eV) or two pump photons ( $\sim 1.16$  eV each) suffice to ionize  $Tm^{2+}$  ions (see Fig. 8). Once the  $Tm^{2+}$  ion is ionized, it becomes a  $Tm^{3+}$  and an electron is released in the conduction band. This electron can recombine on a  $Tm^{3+}$  ion, a trap defect or return to its ionized precursor. This last case allows to bleach the degradation.

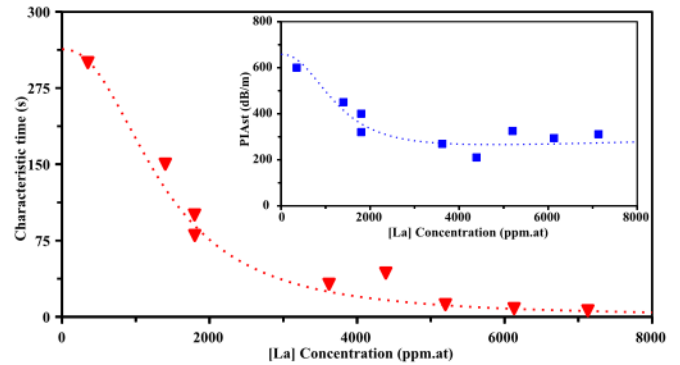
## 5.2. Effect of cerium and lanthanum concentrations on the photo-darkening under pumping

The characteristic times and the  $PIA_{st}$  values for different concentrations of  $Ce$  and  $La$  are reported in Figs. 9 and 10, respectively. In both cases, as the rare-earth ion concentration increases,  $PIA_{st}$  and characteristic time decrease.  $Ce$  reduces  $PIA_{st}$  from 720 to 42 dB/m and the characteristic time from 150 to 15 s.  $La$  reduces  $PIA_{st}$  from 675 to 250 dB/m and the characteristic times from 250 to 6 s.

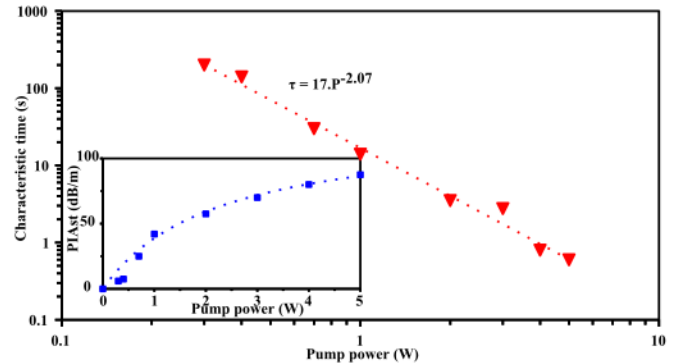
The characteristic time and  $PIA_{st}$  values have been measured for the sample with the highest concentration of  $Ce$  for different pump powers. The data are reported in Fig. 11. As already observed in the case without  $Ce$  (cf Fig. 6), the  $PIA_{st}$  increases with the pump power. Data were fitted with the law  $PIA_{st} = PIA_{st}^{sat} \frac{AP}{1+AP}$  where



**Fig. 9.** Characteristic times and  $PIA$  steady state values (inset) as a function of cerium concentration. Pump power: 1 W, probe power: 2 nW. Dot: experimental values, dotted lines: fittings explained in the text.



**Fig. 10.** Characteristic times and  $PIA$  steady state values (inset). Pump power: 1 W, probe power: 2 nW. Dot: experimental values, dotted lines: fittings explained in the text.



**Fig. 11.** Characteristic times and  $PIA$  steady state values (inset) for the 1300 ppm cerium-doped sample as a function of pump power. Dot: experimental data, dotted lines: fitting in saturate linear for  $PIA_{st}$  and with a power law for the characteristic times.

$PIA_{st}^{sat} = 125$  dB/m and  $A = 0.45$   $W^{-1}$ . The fitting is good except for some data at low pump power. Indeed, at low pump power the equilibrium level is low. Then, a longer length of fibre ( $\sim 8$  cm) was required, and the equilibrium values were undervalued. In comparison with cerium-free samples, the  $PIA_{st}^{sat}$  value is significantly reduced by a factor of 22, and the parameter  $A$  is reduced by a factor of  $\sim 2$ . The values of characteristic time as a function of the pump power are fitted by a single power law, with an exponent of  $\sim 2$ . On the contrary to the case of  $Ce$ -free samples (cf Fig. 6), there is no trend with an exponent of 5 or 6 at low pump power. So, this

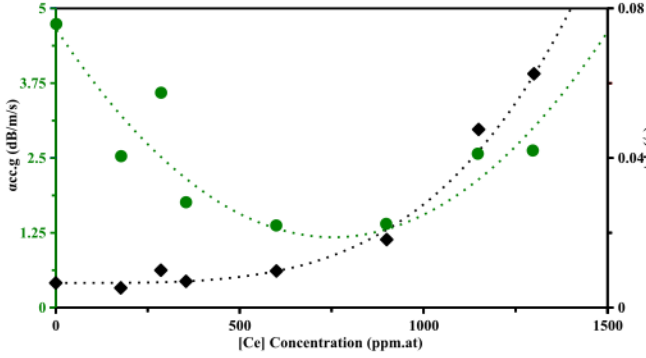


Fig. 12.  $\alpha_{cc.g}$  et  $g+b$  as a function of Ce concentration. Symbol: experimental values, dotted lines: fittings.

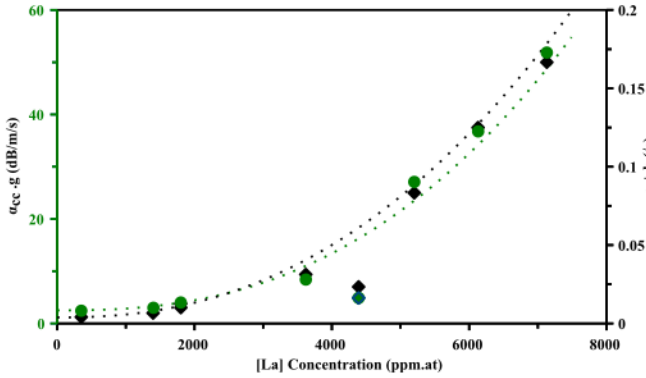


Fig. 13.  $\alpha_{cc.g}$  et  $g+b$  as function of the lanthanum concentration. Symbol: experimental values, line: adjustments in power laws.

strongly suggests that the photo-darkening dynamics would be ruled by bleaching, namely the photo-bleaching by the pump.

The characteristic times and  $PIA_{st}$  values are then interpreted. The  $\alpha_{cc.g}$  and  $g+b$  parameters as a function of Ce and La concentrations are shown in Figs. 12 and 13, respectively.

In the case of Ce-doped samples,  $\alpha_{cc.g}$  parameter has a non-monotonic behaviour. From 0 to 750 ppm of cerium, this parameter decreases from 4.8 to 1.25 dB/m/s while it increases when cerium concentration is higher than 750 ppm. These measurements are fitted with a second order polynomial function and a satisfying match is found (green dotted curve in Fig. 12). The parameter  $g+b$  is quite stationary at low Ce concentration, then it increases. These data are fitted with a power law (the exponent is 3.71) and a good match is also found (black dotted curve in Fig. 12). At low concentration of cerium,  $g$  decreases and  $g+b$  stays constant, meaning that  $b$  increases. At higher concentration of cerium,  $g$  and  $g+b$  increase. Consequently it is impossible to determine the variation of  $b$ .

For La-doped samples, the value of  $\alpha_{cc.g}$  increases by one order of magnitude, from 5 to 50 dB/m/s as the lanthanum concentration increases from 0 to 7000 ppm. For the same range of concentration, the parameter  $g+b$  changes from 0.005 to 0.17 /s. These data were fitted by two different power laws. The exponents are 2.5 and 2.3 for  $\alpha_{cc.g}$  and  $g+b$  curves, respectively. The difference between these two exponents may seem low but no simultaneous satisfying agreement is found for the four charts (Figs. 10 and 13) with equal exponents. As both  $g$  and  $g+b$  increase, it is not possible to discuss confidently on the variation of each parameter.

An inverse mathematical transformation is applied to derive the fitting of  $PIA_{st}$  and  $\tau_{1/e}$  data (dotted curves in Figs. 9 and 10). A good

matching is found for all the data except for two concentrations of Ce ( $\sim 250$  ppm). These two samples have different concentrations of Tm compared to the other samples of the series: Tm concentration is 240 and 350 ppm in the '180 ppmCe' and '260 ppmCe' samples, respectively. As already indicated in Fig. 5, any change in Tm concentration strongly impacts the characteristic times and, to a less extent,  $PIA_{st}$  values.

### 5.3. Effect of cerium and lanthanum concentrations on the photo-darkening after pumping with high probe power

The variations of PIA versus time for three concentrations of Ce and La are reported in Figs. 14 and 15, respectively. Before "Pump stop", the pump laser is on and its power is 1 W. At "Pump stop", the pump laser is switched off and the probe power remains the same.

After the pump is turned off, the rare-earth ion (Ce or La) concentrations influence the recovery of the photo-darkening. The higher the rare-earth ions concentrations, the higher the recovery. One can note that at  $t = 3600$  s (Fig. 14), the PIA absolute value increases suddenly for the sample without cerium. We assume that this increase is due to the vanishing of the green light emitted by  $Tm^{3+}$  under pumping.

Cerium-doped samples exhibit stronger photo-bleaching dynamics than the La ones. However, the bleaching is not complete, even after a long time. Such bleaching effects have been observed previously in cerium-doped fibre under UV excitation [31] and more recently in ytterbium doped fibre [32–34] and ytterbium-cerium co-doped fibre under pumping at 976 nm [35,36]. But the role of cerium on these effects was not discussed and probe powers were not specified.

### 5.4. Fluorescence lifetimes of the ${}^3H_4$ : $Tm^{3+}$ -level

Fluorescence lifetime measurements were performed on all samples series. The monitored emission issued from the  ${}^3H_4$  level of  $Tm^{3+}$  is very sensitive to the phonon energy. For all measurements, the decay curves were non-exponential. Then, we considered the  $1/e$  values for the lifetimes, reported in Fig. 16. For all these samples, the aluminum concentration remains constant around  $\sim 8000$  ppm. So, we assume that the variation of fluorescence lifetimes are only caused by rare-earth ion concentrations.

For Tm samples series, the fluorescence lifetime remains constant (about 19.5  $\mu$ s) when the concentration of Tm varies between 160 and 600 ppm. Despite a quite high concentration of Tm, no quenching was reported in these samples. Interestingly, Ce and La have opposite effect: Ce decreases the fluorescence lifetime ( $-2.8$   $\mu$ s/1000 ppm) while La increases it ( $+1.5$   $\mu$ s/1000 ppm).

The increase of the  $Tm^{3+}$  lifetime is discussed in the light of previous studies related to the effect of the local phonon energy on

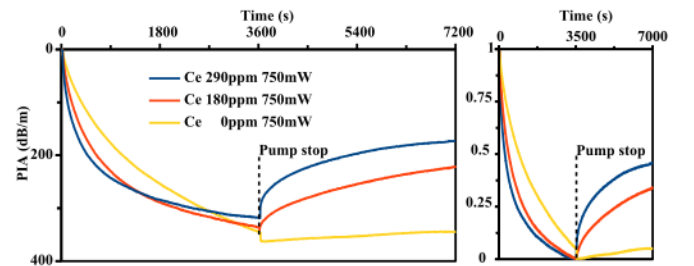


Fig. 14. Left: Time resolved PIA measurements for various Ce concentrations. Pump power before "Pump stop": 750 mW (0 W after "Pump stop"), probe power: 25  $\mu$ W. Right: same measurements with normalized PIA curves.

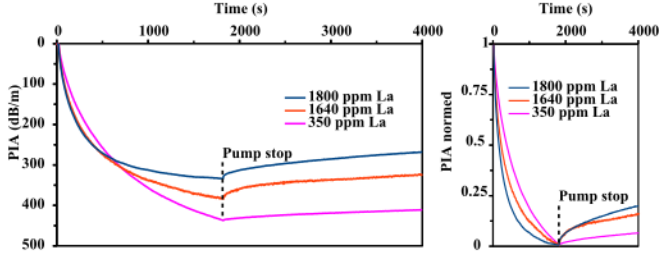


Fig. 15. Left: Time resolved PIA measurements for various *La* concentrations. Pump power before “Pump stop”: 750 mW (0 W after “Pump stop”), probe power: 25  $\mu$ W. Right: same measurements with normalized PIA curves.

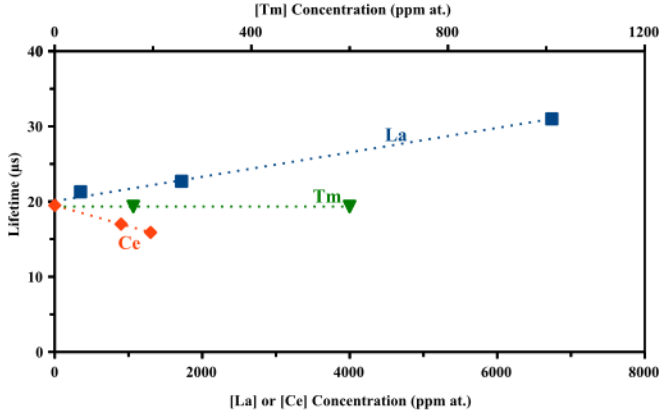
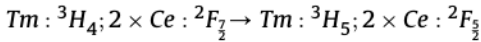


Fig. 16. Fluorescence lifetimes versus concentrations of *Tm*, *La* and *Ce*.

the  ${}^3H_4$  level lifetime [11,12,37]. The improvement of the *Tm*-fluorescence lifetime can be interpreted in the frame of the change of its local phonon energy ( $E_p$ ) as previously reported for aluminium [11]. Indeed, a higher content of *La* in the *Tm* environment may decrease the local phonon energy ( $E_p(La_2O_3) = 400cm^{-1}$ ,  $E_p(Al_2O_3) = 870cm^{-1}$ ,  $E_p(SiO_2) = 1100cm^{-1}$ ), then enhance the population in the  ${}^3H_4$  level, as describe in Ref. [37]. Due to the lower phonon energy of  $La_2O_3$  compared to that of  $Al_2O_3$ , *La* would be more efficient to increase the *Tm*-fluorescence lifetime. The same mechanism would occur with *Ce* but with this ion the lifetime decreases. The decrease of the lifetimes has been reported previously for the  ${}^4I_{1/2}$  level of erbium in tellurium and sulfur based glasses [38–40]. It was explained by an energy transfer mechanism between erbium and cerium ions. As the energy difference between the  ${}^3H_4$  and the  ${}^3H_5$  energy levels of  $Tm^{3+}$  ( $\sim 4000 cm^{-1}$ ) and between the  ${}^4I_{1/2}$  and the  ${}^4I_{13/2}$  energy levels of  $Er^{3+}$  ( $\sim 3600 cm^{-1}$ ) are very close, we hypothesize this energy transfer:



involving one  $Tm^{3+}$  ion and two  $Ce^{3+}$  ions (the energy difference between the  ${}^2F_{5/2}$  and  ${}^2F_{7/2}$  levels of  $Ce^{3+}$  is about  $2000 cm^{-1}$ ). Following this energy transfer, both cerium ions quickly de-excite by emitting two phonons each, and the thulium ion falls into the  ${}^3F_4$  level.

## 6. Discussions

In *Tm*-doped fibres, the photo-induced attenuation is due to the balance between the generation and the bleaching mechanisms.

The generation of color-centres is caused by the ability of  $Tm^{3+}$  ions to emit UV photons through up-conversion mechanisms when it is pumped at  $1.07 \mu m$ . This UV photon ionizes a precursor and, depending on the centre, an electron is released in the conduction band and can be trapped by a  $Tm^{3+}$  ion to form a  $Tm^{2+}$  ion, or a hole is released in the valence band and it can be trapped to form, for example, an *AlOHC* centre. These color-centres can be bleached by absorbing two pump photons or one probe photon. For example,  $Tm^{2+}$  ion recovers its initial trivalent state. By adding *Ce* or *La*, both  $PIA_{st}$  and characteristic time values decrease. This indicates that these ions would act favourably on the photo-bleaching mechanism (see Table 2).

Cerium addition brings an almost total solution to darkening problem. Indeed, the equilibrium levels drop drastically. An addition of 1300 ppm *Ce* induces a drop of 95 % of degradation level. Cerium ions can decrease *g* for three reasons:

- as fluorescence lifetime decreases (Fig. 16), there are less  $Tm^{3+}$  ions in the higher energy levels, then less UV photons are emitted, this can barely affect the generation.
- cerium is known to be present also in its tetravalent state in silica [15,41].  $Ce^{4+}$  can absorb UV photon (charge transfer band) emitted by  $Tm^{3+}$  ions to form  $Ce^{3+}$  ion which is stable over several months [42,43]. These two valence states of cerium are transparent in the visible [44]. Hence, this phenomenon is in competition with the generation of color-centre and therefore reduces the photo-darkening.
- cerium ion has two valence states, it can trap electron ( $Ce^{4+} \rightarrow Ce^{3+}$ ) and hole ( $Ce^{3+} \rightarrow Ce^{4+}$ ) to the detriment of the formation of color-centres.

Moreover, when  $RE^{3+}$  ion is added into aluminosilicate glasses, it acts as a charge compensator if  $[RE_2O_3] \leq \frac{[Al_2O_3]}{3}$  (or  $[REO_2] \leq \frac{[Al_2O_3]}{4}$  for  $Ce^{4+}$ ) [45]. As a consequence, RE is located preferentially close to the tetra-coordinated  $Al^{IV}$  species. *Ce* and *Tm* compete to be located close to  $Al^{IV}$  species. Therefore, *Ce* tends to reduce the probability of *Tm*-cluster formation and hence the emission of UV photons. When cerium concentration is above the threshold of the charge compensator effect, it acts as network modifier and depolymerizes the silica network. This depolymerization may lead to the formation of new precursors defects [46]. This may explain why *g* increases at higher content of cerium.

The bleaching mechanism is related to the ionization of color-centres. In this case, an electron is released in the conduction band. The presence of cerium ions offers a new opportunity for this electron to be trapped by  $Ce^{4+}$ , to the detriment of  $Tm^{3+}$  ions. The same stands for the ionization of the *AlOHC* centre as the hole released in the valence band can be trapped by  $Ce^{3+}$  ions.

On the contrary to cerium ions, the addition of lanthanum tends to increase the fluorescence lifetime which would be a detrimental effect on the photo-darkening. Indeed, this would promote the population of  $Tm^{3+}$  in the high energy levels, then the emission of UV photon and the generation of color-centres. However, lanthanum ions allow to reduce photo-darkening. Compared to cerium, lanthanum ion does not absorb UV photon and has only one stable valence state,  $La^{3+}$ . The divalent and tetravalent states are not stable in silica [27,28]. An electron has the possibility to recombine on ionized precursor,  $Tm^{3+}$  and  $La^{3+}$ . If it recombines on  $La^{3+}$ , the new valence is not stable and the electron will move away until reaching a stable recombination. Hence, lanthanum addition decreases the probability of recombination on a thulium, and keeps constant the probability of recombination of an ionized precursor.



Therefore, the addition of lanthanum would lead to the increase of the parameter  $b$  because it would increase the probability for an electron to return to an ionized precursor. Moreover, as already discussed in the case of cerium ions, at low content of  $La$ , this ion could limit the formation of  $Tm$ -clusters [47], while at high content, this ion would also act as a network modifier, increasing the numbers of precursor defect [46].

## 7. Conclusion

Three sets of samples were investigated, all doped with thulium and aluminium and co-doped with cerium or lanthanum. In each series, the impact on photo-darkening of rare-earth ions was studied. The photo-darkening was characterized by its steady state value ( $PIA_{st}$ ) and also its characteristic time. Thanks to a rate equation model, these parameters were related to the generation and sum of generation and bleaching rates. While  $PIA_{st}$  increases with the  $Tm$  concentration, the addition of  $Ce$  or  $La$  allows the  $PIA_{st}$  and the characteristic time to reduce. Most importantly, we reveal that the main impact of cerium and lanthanum on photo-darkening is due to their effect on the bleaching mechanisms. This can be explained by the ability of cerium ions to trap holes or electrons. In the case of lanthanum co-doping, the probability for a charge to return to an ionized precursor would be increased by the instability of the divalent and tetravalent states of lanthanum.

Cerium and lanthanum reduce the steady state degradation because they have a strong impact on photo-bleaching. This an increment on the comprehension of darkening process. These results could be transposed to other systems like  $Yb$ - or  $Er$ -doped fibres in order to enhance actual devices and create new devices and new applications.

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