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To cite this version:

Vera Pukhkaya, Philippe Goldner, Alban Ferrier, Nadège Ollier. Impact of rare earth element clusters on the excited state lifetime evolution under irradiation in oxide glasses. Optics Express, 2015, 23 (3), pp.3270-3281. $10.1364/\text{OE}.23.003270$. hal-01275486

HAL Id: hal-01275486 <https://hal.sorbonne-universite.fr/hal-01275486v1>

Submitted on 17 Feb 2016

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Impact of rare earth element clusters on the excited state lifetime evolution under irradiation in oxide glasses

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Abstract: Rare earth doped active glasses and fibers can be exposed to ionizing radiations in space and nuclear applications. In this work, we analyze the evolution of ${}^{2}F_{5/2}$ excited state lifetime in Yb³⁺ ions in irradiated aluminosilicate glasses by electrons and γ rays. It is found that the variation of lifetimes depends on the Yb^{3+} clusters content of the glasses for irradiation doses in the $10^2 - 1.5 \cdot 10^9$ Gy range. In particular, glasses with high clustering show a smaller decrease in lifetime with increasing radiation dose. This behavior is well correlated to the variation in paramagnetic defects concentration determined by electron paramagnetic resonance. This effect is also observed in Yb^{3+} doped phosphate and Er^{3+} doped aluminosilicate glasses, inferring that clustering plays an important role in irradiation induced quenching.

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OCIS codes: (140.3615) Lasers, ytterbium; (160.5690) Rare-earth-doped materials; (160.2750) Glass and other amorphous materials; (260.3800) Luminescence; (350.5610) Radiation.

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1. Introduction

 Yb^{3+} ion is an essential dopant in glasses and fibers for IR-lasers due to its highly-efficient emission at \sim 1 µm. Owing to its unusual energy level scheme with only two manifolds separated by about 10000 cm⁻¹, Yb^{3+} offers interesting spectroscopic properties for optical applications such as absence of excited state absorption and relatively long emission lifetime. The lifetime of Yb^{3+} excited state ${}^{2}F_{5/2}$ is an important spectroscopic parameter because it determines the laser pump saturation intensity $I_{sat} \sim 1/\tau_{IR}$ which should be as low as possible [1].

To increase the IR-laser gain, Yb^{3+} concentration can be increased but results in the formation of clusters, especially in pure $SiO₂$ glasses where the network rigidity is not suitable to host a high content of rare earth ions. In glasses containing modifier ions, the local environment of Yb^{3+} ions is defined preferentially by Non-Bridging Oxygens. Their concentration impacts the Yb clusters content in oxide glasses as was demonstrated in aluminosilicate, phosphate and aluminoborosilicate glasses [2, 3].

Active fibers such as Yb^{3+} or Er^{3+} doped ones can be exposed to ionizing radiations in nuclear or space applications [4]. Besides quantifying the Radiation Induced Absorption (RIA) and analyzing the corresponding point defects, it is also necessary to estimate the influence of ionizing radiations on rare earth element spectroscopic properties in these fibers. Only a few papers have reported such studies up to now. It has been shown that ionizing radiations (electron, γ rays) can affect Yb³⁺ spectroscopic properties in aluminoborosilicate glasses such as the ${}^{2}F_{5/2}$ excited state lifetime and the cooperative emission intensity [5]. This was correlated to the formation of hole centers type defect under irradiation, in this case, Non-Bridging Oxygen Hole Centers (NBOHC) [5]. In Er-doped fibers, a similar decrease of 4 ⁴I_{13/2} lifetime has been observed under γ-irradiation [6, 7] and ion radiation [8]. This was attributed to non-radiative energy transfers from the $\frac{4}{1_{13/2}}$ level to radiation-induced point defects [6, 7].

In the present study, we investigate the evolution of Yb^{3+} exited state lifetime under ionizing radiations (electrons, γ rays) for a broad dose range (10²-10⁹ Gy) in a series of aluminosilicate glasses. The composition of these glasses was chosen to study the influence of $Yb³⁺$ clustering on spectroscopic changes. This was motivated by our previous observation of Yb^{3+} clustering impact on the thermal recovery of radiation induced paramagnetic point defects [9]. Here, we show that Yb^{3+} excited state lifetime decreases at a lower rate as a function of the irradiation dose in glasses with high clusters amount. A good agreement is also found with the variation of the defect concentration in these glasses. The same excited state lifetime evolution is also observed in Yb^{3+} doped phosphate glasses and Er^{3+} doped aluminosilicate glasses, suggesting that this behavior is independent on the particular rare earth-glass combination.

2. Experimental details

Four Yb-doped and two Er-containing aluminosilicate glass compositions were synthesized. For that, appropriate amounts of dried powders of SiO_2 , Al_2O_3 , Na_2CO_3 Yb_2O_3 or Er_2O_3 were mixed thoroughly to obtain a well-homogenized powder and slowly heated (during 10 hours) in a Pt-Au crucible from room temperature up to 1600°C (melting temperature) in a furnace under air during 4 hours. Slow heating facilitated the $CO₂$ release process taking place at \sim 800 $^{\circ}$ C. In addition, it promoted higher homogeneity of the melt. Afterwards, glasses were quenched at room temperature and annealed at 580° C (below T_g) in order to reduce the residual thermal stresses. The nominal aluminosilicate glass compositions are presented in Table 1. Samples are labeled according to the Aluminum Saturation Index $(ASI = Al/Na)$, which characterizes the Non-Bridging Oxygen concentration. Actual glass compositions were

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determined with microprobe analysis giving <1 mol. % of error compared to nominal compositions.

Phosphate glass samples were prepared starting from $NH_4H_2PO_4$, MgO , Na_2CO_3 and $Yb₂O₃$ (5 wt. %). The initial powders were mixed and melted in a silica crucible and then slowly heated from room temperature up to 900°C. As in the case of aluminosilicate glasses, slow heating favored $CO₂$ release and homogenization. The melted glass was quenched rapidly into a Pt-Au crucible on an electric plate at 300°C to avoid a high cooling rate, and immediately annealed at 350°C to reduce residual thermal stresses. The nominal phosphate glass compositions are presented in Table 2.

Sample name	SiO ₂ mol. $%$	Al_2O_3 mol. $%$	Na ₂ O ₂ mol. $%$	ASI Al/Na	
ASI03 Yb5 ASI06 Yb5	74 68	6 12	20 20	0.3 0.6	
ASI09 Yb5 ASI11 Yb5	62 58	18 22	20 20	0.9 $1.1\,$	$+5$ wt. % Yb ₂ O ₃
ASI09 Yb05 ASI09 Yb8	62	18	20	0.9	$+0.5$ wt. % Yb ₂ O ₃ $+8$ wt. % Yb ₂ O ₃
ASI03 Er	74	6	20	0.3	$+0.5$ wt. % Er ₂ O ₃
ASI09 Er	62	18	20	0.9	$+0.5$ wt. % Er ₂ O ₃

Table 1. Nominal Yb3+ and Er3+- doped Aluminosilicate Glass Compositions

Bulk glass samples $(5 \times 5 \text{ mm}^2, 500 - 700 \text{ }\mu\text{m}$ thickness) were continuously irradiated at SIRIUS electron accelerator (Laboratoire des Solides Irradiés, Palaiseau, France) with the 2.5 MeV electron beam with a dose rate close to 25 MGy/h. The sample holder was maintained around 40°C with a water-cooling system. The achieved integrated doses were 10^5 , 10^6 , $3 \cdot 10^6$, 10^7 , 10^8 and $1.5 \cdot 10^9$ Gy. Samples were also irradiated with γ-rays of 1.25 MeV from Co⁶⁰ source in **"**Centre National des Sciences et Technologies Nucléaires" in Tunisia (*CNSTN*). The dose rate in this case was 5.64 kGy/h. The achieved doses were 10^4 , 10^5 , 10^6 and $3 \cdot 10^6$ Gy. Two lower doses 100 Gy and 1000 Gy were achieved with γ -rays of 1.25 MeV from Co⁶⁰ source in "Institut de Radioprotection et de Sûreté Nucléaire" (IRSN, Fontenay-aux-Roses, France). The dose rate was 2 Gy/min. The error bar on the radiation dose after electron irradiation is considered as 10% because of the backscattered electrons that were not taken into account in the dose calculation". Concerning gamma rays irradiation, at 100 Gy the error is 5.5%.

Table 2. Nominal Yb3+-doped Phosphate Glass Compositions

Sample name	P_2O_5 mol. $%$	MgO mol. $%$	Na ₂ O, mol. $%$	
MP50	50	16.7	33.3	
PIP45	45.4	18.2	36.4	$+5$ wt.% Yb ₂ O ₃
P ₁ P ₄₀	40	20	40	

 Yb^{3+} emission ${}^{2}F_{5/2} \rightarrow {}^{2}F_{7/2}$ in the IR-region was recorded on a HORIBA Jobin Yvon spectrofluorimeter under Xe lamp excitation at 975 nm with a 1200 gr/mm grating and a RG-830 filter. The lifetime of Yb^{3+} excited state ${}^{2}F_{5/2}$ was measured at 1000 nm with a time resolution of 100 µs. The lifetime of $Er^{3+4}I_{13/2}$ excited state was measured using pulsed laser excitation at 975 nm (Ekspla optical parametric oscillator NT342B) and detection by an InGaAs detector. Yb^{3+} cooperative luminescence was excited at 975 nm by a laser diode, dispersed by a Jobin-Yvon H25 spectrometer and detected by a photon counting photomultiplier tube. In order to perform accurate cooperative luminescence measurements, the samples were grounded into powders of well-controlled particle size in the range 100-125 µm [2].

Electron Paramagnetic Resonance (EPR) experiments were carried out on a Bruker Xband EMX spectrometer. Paramagnetic point defects were analyzed at room temperature and $Yb³⁺$ signal was recorded at 4K. The microwave frequency was 9.8 GHz. Spectra were

normalized by attenuator gain and sample mass to allow quantitative comparisons. Radiationinduced point defects were also analyzed by time-resolved photoluminescence at room temperature. We used a frequency-doubled pulsed Nd:YAG laser for excitation and a Shamrock SR-303i spectrometer and an ANDOR ICCD camera for detection. The delay was 500 ns and the gate width was $100 \mu s$.

3. Results and discussion

3.1 Yb3+ doped aluminosilicate glasses

3.1.1 Optical and EPR spectroscopy of non-irradiated samples

The lifetimes τ_{IR} of ${}^{2}F_{5/2}$ Yb³⁺ excited state in non-irradiated aluminosilicate glasses are displayed in Table 3. Although the concentration of Yb^{3+} was maintained at 5 wt. % of Yb_2O_3 , we observed that τ_{IR} decreased from 2.14 ms in ASI03_Yb5 glass to 0.94 ms in ASI11 Yb5 glass. We attribute this to changes in $Yb³⁺$ environment in the different glasses, resulting in variations of the ${}^{2}F_{5/2} \rightarrow {}^{2}F_{7/2}$ transition oscillator strength [10].

Table 3. Yb3+ 2F5/2 Lifetimes in Non-irradiated Aluminosilicate Glasses

Glass sample	${}^{2}F_{5/2}$ lifetime (ms)
ASI03 Yb5	2.14
ASI06 Yb5	1.97
ASI09 Yb5	1.15
ASI11 Yb5	0.94

 $Yb³⁺$ environment changes are also clearly seen in the normalized emission spectra shown in Fig. 1. In particular, the peak located at 1006 nm in ASI03_Yb5 and ASI06_Yb5 shifts to 1027 nm in ASI09_Yb5 and ASI11_Yb5 glasses. This could reflect a larger Stark splitting in the latter glasses due to a stronger crystal field. These changes in Yb^{3+} environment were confirmed by EPR spectroscopy (see below).

Fig. 1. Normalized Yb^{3+} emission spectra in non-irradiated Yb^{3+} doped aluminosilicate glasses under 975 nm excitation.

 Yb^{3+} clustering was probed by cooperative luminescence [2]. Corresponding spectra for non-irradiated glasses are shown in Fig. 2. Spectra were normalized by the factor $(\alpha_D * \tau_{IR})^2$, where α_D is the absorption coefficient calculated by integrating the product of the absorption spectrum of the sample by the emission spectrum of the diode (normalized at maximum). Since excitation absorption was low in the powder geometry, the cooperative emission probability, which reflects Yb^{3+} cluster concentration, was proportional to the area under the

normalized cooperative emission spectra [3]. The cooperative emission probability increases strongly from ASI03_Yb5 glass to ASI11_Yb5 (Fig. 2) and the glass series can be divided into two groups: $\overline{ASI03}$ Yb5-ASI06 Yb₅ (low Yb³⁺ cluster amount) and ASI09 Yb5-ASI11 Yb5 (high Yb³⁺ cluster amount). The high clustering in ASI09 and ASI11 glass compositions can be explained by a lower amount of NBO due to an increasing $Al³⁺$ content that must be compensated by Na⁺ ions [2]. We checked by MAS NMR of ²⁷Al the fourfold coordination of Al^{3+} in those glasses [9] in order to exclude any modifications of Al^{3+} coordination number.

Fig. 2. Normalized cooperative luminescence spectra of non-irradiated Yb-doped aluminosilicate glasses.

To further probe Yb^{3+} environment and clustering, EPR spectra were recorded at 4K (Fig. 3). The broad line in the whole range of magnetic field corresponds to well-diluted Yb^{3+} ions in glass [5, 11, 12]. In ASI09 Yb5 and ASI11 Yb5 glasses, the EPR signal is much lower despite the same amount of Yb^{3+} in all four glasses (Fig. 3, inset). This decrease indicates that part of Yb^{3+} ions is not detected, probably because of too short spin relaxation time T_1 . This can be due to the formation of Yb^{3+} ion clusters [11], in agreement with the cooperative luminescence results. Besides, in both glasses, a shoulder at $g = 12.8$ can be seen, which we assign to Yb^{3+} clusters [16]. Indeed, the theoretical g-value for isolated Yb^{3+} ions does not exceed 2Λ·Μ where Μ is the maximum of the projection of the total angular momentum and Λ is the Landé factor. The ground multiplet of $\overline{Y}b^{3+}$ is ² $F_{7/2}$, corresponding to $\Lambda = \frac{8}{7}$ and $M = \frac{7}{7}$. Thus, $\alpha = 8$ for isolated Xb^{3+} ions [111], suggesting that the $\alpha = 12.8$ shoulder corres $^{7}/_2$. Thus, g_{max} 8 for isolated Yb³⁺ ions [11], suggesting that the g = 12.8 shoulder corresponds to clusters.

 Yb^{3+} EPR spectra were analyzed in more details by assuming an axial site symmetry. Determining the exact parameters of the g-tensor is difficult in glasses because of inhomogeneous broadening. The value of $g\perp$ is estimated to 3.2 in ASI03 Yb5 and ASI06_Yb5 and 4.1 in ASI09_Yb5 and ASI11_Yb5 glasses. We did not perform simulations to obtain g∥ parameters but we estimate g∥ ~1.3-1.4 in ASI03 Yb5 and ASI06 Yb5 and g∥ \sim 1.1 in ASI09 Yb5 and ASI11 Yb5. The variation in g⊥ with glass composition is in agreement with the emission spectra and reflects different environments for $Yb³⁺$ ions in low and high ASI index glasses. It should also be noted that the $g\perp$ factors obtained in Yb-doped aluminoborosilicate (ABS) glasses with low clustering content were in the range 3.17 to 3.32 [12], close to the values found here for ASI03_Yb5 and ASI06_Yb5, in which clustering is low too.

Fig. 3. EPR spectra normalized at maximum signal of non-irradiated Yb-doped aluminosilicate glasses $(T = 4K)$. Inset: normalization by mass and gain only.

3.1.2 Evolution of $Yb^{3+2}F_{5/2}$ lifetime under irradiation in aluminosilicate glasses

First, we studied the evolution of lifetimes after irradiation for delays up to 4 months. Despite irradiation, all fluorescence decays were well fitted by a single exponential. Figure 4 displays the obtained τ_{IR} values in irradiated glasses at 10⁵ Gy at different times after irradiation. The values before irradiation are labeled by larger symbols. In all samples, irradiation results in a clear decrease (40-50%) of the ${}^{2}F_{5/2}$ lifetime. Then, for glasses ASI09_Yb5 and ASI11_Yb5 there is almost no variation of τ_{IR} for delays up to 4 months. In opposition, for glasses ASI03_Yb5 and ASI06_Yb5, one can observe a slight increase of the τ_{IR} . In all glasses, the lifetime of Yb^{3+} excited state ${}^{2}F_{5/2}$ did not reach its pre-irradiation value on the time scale investigated.

Fig. 4. ${}^{2}F_{5/2}$ lifetimes of Yb³⁺ excited state recorded at different times after electron irradiation $(10⁵$ Gy dose). The non-irradiated glasses are shown with larger symbols.

This behavior can be understood by considering that the ${}^{2}F_{5/2}$ level is quenched by the defects induced by the irradiation. Indeed, the concentration of paramagnetic point defects induced by irradiation decreases with time, with a slower rate in glasses with higher $Yb³⁺$ clustering [9]. This is explained by the ability of clustered Yb^{3+} ions to efficiently trap charges under a stable form during the irradiation process. As a consequence, point defects thermal recovery is slowed down, as its τ_{IR} increase.

We also analyzed the dose effect on τ_{IR} lifetime after 6 months to reach stable values. Figure 5 displays τ_{IR} as a function of lg(dose) obtained for both electron and gamma irradiation. First, we see that irradiation at 100 Gy has no impact on the ${}^{2}F_{5/2}$ lifetime in glasses with high or low Yb^{3+} cluster content. The first effect can be seen at 1000 Gy. This result is meaningful for space applications where cumulated doses reach 300 Gy in 10 years, although the dose rate is much lower than the one used in our experiments. Consequently a direct extrapolation of this result to space application is not straightforward.

Fig. 5. ²F_{5/2} lifetimes for e⁻ and γ irradiated Yb³⁺-doped aluminosilicate glasses (measured 6 months after irradiation). Solid (open) symbols: electron (gamma) irradiation; larger symbols: non-irradiated values. Error bars are shown for ASI03_Yb5 glasses only for easier reading.

For glasses with lower ASI parameters – ASI03_Yb5 and ASI06_Yb5 – in both the lower dose range (<10⁵Gy) and in the higher dose range (>3·10⁶Gy) clear linear decrease is seen (Fig. 5). However, little or no variation occurs in the $10^5 - 3 \cdot 10^6$ Gy dose range. In contrast, for the other two glass compositions with higher ASI parameter, the linear variation in lifetime τ_{IR} presents smaller slopes in the lower and stronger doses regions and there is little evidence for a plateau in the $10^5 - 3 \cdot 10^6$ Gy dose range.

We also observed no difference in lifetimes between e^- - and γ -irradiated glasses (Fig. 5). This indicates that the concentration and nature of the defects involved in ${}^{2}F_{5/2}$ quenching are similar in both cases, in the limit of our observations (three dose levels).

The τ_{IR} lifetime evolution was further analyzed in ASI09 aluminosilicate glasses doped with various Yb^{3+} concentrations: 0.5, 5 and 8 wt. % (Fig. 6). At low Yb^{3+} concentration, the behavior of low clustering glasses ASI03 and ASI06 is reproduced, whereas at high $Yb³⁺$ concentration, the lifetime evolution follows that of ASI09 and ASI011. Taking into account that we checked that clustering is increasing with increasing $Yb³⁺$ concentration, we conclude that Yb³⁺ clustering, and not glass composition, modifies the evolution of τ_{IR} as a function of irradiation dose.

Fig. 6. Variation of the ${}^{2}F_{5/2}$ decay time value of Yb^{3+} as a function of the logarithmic dose. The irradiation has been performed with e⁻-on aluminosilicate glasses ASI09_Ybx with different Yb concentration. Empty symbols are used for gamma irradiation

3.1.3 Correlation between ${}^{2}F_{5/2}$ lifetime and radiation-induced point defects

In order to understand the ${}^{2}F_{5/2}$ lifetime variation, we measured the paramagnetic point defects nature and concentration, and their evolution under an annealing treatment. Figure 7 displays the total amount of paramagnetic point defects as a function of irradiation dose. This was determined from double integration of the EPR spectra in the $g = 2$ region [9]. A plateau (10⁵-3·10⁶ Gy) is followed by an increase of defects with increasing dose, and then further followed by a decrease at the highest radiation exposure level. Moreover, glasses with the higher cluster content (ASI09_Yb5 and ASI11_Yb5) show a smaller variation of defects concentration with dose. This is in qualitative agreement with the limited ${}^{2}F_{5/2}$ lifetime variation with irradiation dose (Fig. 5). For glasses with lower clustering (ASI03_Yb5 and ASI06_Yb5), a similar correlation is observed.

Fig. 7. Total amount of paramagnetic point defects in irradiated Yb³⁺-doped aluminosilicate glasses.

In order to examine the possible role of particular point defects in lifetime decrease, we carried out annealing treatments on two $Yb³⁺$ -doped aluminosilicate glasses with opposite $Yb³⁺$ cluster content: ASI03 Yb5 and ASI09 Yb5. As a matter of fact, bleaching temperature is unique for each defect since it is associated to its energy activation, so an annealing treatment is a tool to separate defect contribution.

The total defect concentration of two Yb^{3+} -doped aluminosilicate glasses as a function of annealing temperatures is presented in Fig. 8. The precise analysis of paramagnetic defect in those glasses was reported in details elsewhere [9]. In ASI03_Yb5, the lifetime of Yb^{3+} excited state ²F_{5/2} is already back to its pre-irradiation value \sim 2.14 ms after annealing at 244 °C [Fig. 8(b)]. However, there are still defects in the glass, which are partly identified as HC type (HC1 and HC2) from the EPR spectrum [9]. Moreover, at the same temperature, we observed under 532 nm laser excitation, a broad band centered at 725 nm that we attribute to NBOHC [Fig. 8(b)]. Even if the position is deviated from 620 nm as in $SiO₂$ glass [13], we assume that the presence of Al can lead a red-shift the position band [14]. We conclude from these results that neither HC defects nor NBOHC defects are solely responsible for the decrease of the ${}^{2}F_{5/2}$ lifetime.

Fig. 8. ${}^{2}F_{5/2}$ lifetimes and total concentration of paramagnetic defects (left), and photoluminescence spectra of NBOHC (right) in ASI09_Yb5 (a) and ASI03_Yb5 (b) glasses obtained after annealing at various temperatures. Electron irradiation at 10^8 Gy.

Contrary to ASI03_Yb5, τ_{IR} in ASI09_Yb5 is not back to the pre-irradiation value after annealing at 400° C [Fig. 8(a)]. At this temperature, peroxy radicals were evidenced by EPR, whereas neither Al-OHC nor NBOHC defects were detected [Fig. 8(b)]. As the absorption band of peroxy radicals centered at 1.97 eV (630 nm) [13] extends to Yb^{3+} emission range, they could be responsible for the decrease of ${}^{2}F_{5/2}$ lifetime.

3.2 Extension to other systems

We first studied ${}^{2}F_{5/2}$ lifetime as a function of irradiation dose and cluster content in Yb³⁺ doped phosphate glasses. As in aluminosilicate glasses, one can observe a multi-step τ_{IR} variation in poly-phosphate glasses, in which the cluster amount is low (Fig. 9), opposed to a monotonically decrease in meta-phosphate glasses with higher Yb^{3+} clustering. The cluster concentration was determined by cooperative luminescence [2]. This result on phosphate glasses first confirms the role of clusters and second suggests that lifetime behavior is irrelevant to the glass composition and consequently on the type of point defect formed.

Fig. 9. ${}^{2}F_{5/2}$ lifetimes of Yb³⁺ doped-phosphate glasses.

We also investigated $Er³⁺$ doped glasses, in order to further assess the role of clusters. Figure 10 displays the lifetime of ${}^{4}I_{13/2}$ excited state in ASI03 and ASI09 glasses as a function of $lg(dose)$. In ASI03_Er05 glass with low Er^{3+} cluster content, the $^{4}I_{13/2}$ lifetime evolution is similar to that of glasses with low Yb^{3+} cluster content, whereas in ASI09 Er05 glass the lifetime variation is lower, as in high $Yb³⁺$ clustering glasses (see Fig. 5). This suggests that the de-excitation channels via point defect formation are similar for Yb^{3+} and Er^{3+} ions.

Fig. 10. ${}^{4}I_{13/2}$ lifetimes of Er^{3+} doped-aluminosilicate glasses.

4. Conclusion

We have studied Yb^{3+} doped aluminosilicate glasses with varying clusters content, which was evidenced by cooperative luminescence and EPR spectroscopy. Yb^{3+} excited state lifetime was recorded as a function of irradiation dose in a 10^2 -1.5⋅10⁹ Gy range (electron and γ rays) and delays after a 10^5 Gy dose irradiation. Ionizing radiations were found to decrease lifetimes with rates depending primarily on Yb^{3+} clustering and not glass composition. In particular, glasses with high clustering show a smaller decrease in lifetime with increasing radiation dose. This behavior was in agreement with the variation in paramagnetic defects

concentration. The nature of quenching defects was studied by annealing, suggesting that peroxy radicals could have a larger effect than other types of hole centers. In addition, clustering was found to impact lifetimes in Yb^{3+} doped phosphate glasses as well as in Er^{3+} doped aluminosilicate ones. This suggests that this effect could be observed in a broad range of rare earth doped glasses.

Acknowledgments

We thank B. Boizot and V. Metayer for electron irradiations (LSI, SIRIUS, France). The SIRIUS accelerator is a part of National Network of Accelerators Dedicated to Materials Irradiation. We acknowledge A. Mejri for gamma irradiations (Tunisia) and P. Aschehoug (ENSCP, France) for Er and Yb lifetime measurements. We are grateful to F. Trompier (IRSN, France) for lower dose gamma-irradiations.