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Abstract	This work analyzes t	This work analyzes the influence of the nanocrystals on the luminescence	
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Chapter 15 Luminescence of Er³⁺ Ions in Nanocrystalline Glass-Ceramics

Rolindes Balda, R. Morea, J. Gonzalo, and J. Fernandez

Abstract This work analyzes the influence of the nanocrystals on the luminescence 5 properties of Er^{3+} ions in transparent oxyfluoride tellurite glass-ceramics obtained 6 by heat treatment of the precursor Er-doped glass. The comparison between the 7 fluorescence properties of Er-doped glass and glass-ceramic suggests that a fraction 8 of Er^{3+} ions are forming nanocrystals while the rest remain in a glass environment. 9 The presence of ErF_3 nanocrystals has a strong effect in the upconverted red 10 emission from ${}^4F_{9/2}$ level. The time evolution of the upconverted red emission shows 11 that energy transfer upconversion processes are responsible for the increase of this 12 emission in the glass-ceramic sample.

15.1 Introduction

Rare earth-doped transparent oxyfluoride glass-ceramics have been the subject of 15 several recent investigations since they possess optical properties of interest for 16 photonic applications such as lasers, optical amplifiers, sensors, light converters 17 and so on. In particular, these materials containing one or more crystalline phases 18 embedded in the glass matrix, combine the low phonon energy, optical transparency 19 and rare-earth (RE) ions solubility of fluoride crystals with the good mechanical, 20 thermal, and chemical properties of oxide glasses [1].

The first transparent glass-ceramic (TGC) based on aluminosilicate glass doped ²² with Er^{3+} and Yb^{3+} ions was developed by Wang & Ohwaki in 1993 [2]. In ²³ this glass, cadmiun lead fluoride nanocrystals were nucleated by suitable thermal ²⁴ treatments. It was observed that, exciting Yb^{3+} ions at 970 nm, the green-red ²⁵ upconversion emission from Er^{3+} ions was 100 times more intense than observed ²⁶

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from fluoride glass, which suggested the segregation of Yb^{3+} and Er^{3+} ions in ²⁷ the nanocrystals. Since then, different rare earth doped oxyfluoride glass ceramics ²⁸ mainly based on silicate glass matrices and other fluoride crystal phases, such as ²⁹ LaF₃ [3–5], CaF₂ [6, 7], BaF₂ [8, 9], etc. have been proposed as active media ³⁰ for solid state lasers, optical amplifiers, phosphors or to enhance the efficiency ³¹ of photovoltaic cells [10, 11]. However, the limited RE solubility of silica-based ³² compositions and small nonlinear behavior as well as the low chemical durability of ³³ pure fluoride compositions has motivated the search for other glass precursors for ³⁴ TGC. ³⁵

Among different oxide glasses, tellurite glasses combine good mechanical stability, chemical durability, and high linear and nonlinear refractive indices, with a wide transmission window (typically $0.4-6 \mu m$), which make them promising materials for photonic applications [12]. In fact, broadband Er-doped fiber amplifiers have been achieved by using tellurite-based fibers with a gain of 25.3 dB and a noise figure of less than 6 dB from 1561 to 1611 nm [13] and more recently, efficient laser emission around 2 μm has been demonstrated in a tellurite fiber doped with Tm³⁺ ions [14].

Mixed fluorotellurite glasses which combine the low phonon energies of fluorides 44 with the high chemical durability and thermal stability of tellurites can reduce the 45 OH content which has a great influence in the quenching processes of the radiative 46 emission of excited levels of rare earth ions, which increases fluorescence lifetimes 47 [15, 16]. The transformation, by the adequated heat treatment, of fluordellurite 48 glasses into glass-ceramics in which the RE ions are incorporated in the crystalline 49 phase can increase the maximum cross-sections of the RE ions and the emission 50 efficiency. However, up to now, there are only a few works dealing with Er^{3+} -doped 51 fluorotellurite glass ceramics [17, 18].

In this work, we report the effect of the heat treatment on absorption spectra, ⁵³ near infrared luminescence, and upconversion emission of Er^{3+} -doped transparent ⁵⁴ oxifluoride tellurite glass-ceramics obtained by the heat treatment of the Er^{3+} - ⁵⁵ doped 73.6 TeO₂-17.6 ZnO-8.8 ZnF₂ glass. The comparison of the fluorescence ⁵⁶ properties of Er^{3+} -doped precursor glass and glass-ceramic confirms the successful ⁵⁷ incorporation of the rare-earth into the nanocrystals. ⁵⁸

15.2 Theoretical Background

the number of electrons in the unfilled 4f shell.

The rare-earth ions, referred to as the lanthanides, comprise the series of elements in $_{60}$ the sixth row of the periodic table after lanthanum from cerium to ytterbium. These $_{61}$ atoms are usually incorporated in crystalline or amorphous hosts as trivalent ions $_{62}$ and occasionally as divalent ions. The ground state configuration of the trivalent $_{63}$ rare-earth ions is [Xe]4fⁿ where n varies from 1 (Ce³⁺) to 13 (Yb³⁺) and indicates $_{64}$

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The 4f electrons are shielded by external full outer shells which limits the 66 interaction between the ion and the crystalline field to a small perturbation giving 67 the RE ions their characteristic sharp and well defined spectral features. 68

The optical activity of the RE ions in solids occurs mainly between electronic ⁶⁹ states within the 4f configuration. These transitions are parity forbidden and are ⁷⁰ made partially allowed by crystal field interactions mixing states of different parity. ⁷¹ The result is that the transitions between 4f states are weak, with oscillator strengths ⁷² of the order of 10^{-6} , and radiative lifetimes in the micro and miliseconds range. ⁷³ The long lifetime plays an important role to increase the probability of sequential ⁷⁴ excitations in the excited states of a single ion as well as in permitting ion-ion ⁷⁵ interactions in the excited states to allow energy transfer. ⁷⁶

The lifetime of an emitting level is governed by a sum of probabilities for 77 several competing processes: radiative decay, nonradiative decay by multiphonon 78 (MPH) relaxation and by energy transfer to other ions. MPH relaxation processes 79 in which the energy difference between the emitting level and the next lower level 80 is converted into many lattice phonons are strongly dependent on the host matrix. 81 Generally, the larger the number of phonons needed to convert the excitation energy, 82 the lower is the efficiency of the non radiative processes. Therefore to enhance the 83 emission efficiency hosts with low phonon energy are required. 84

Energy levels in rare-earth ions are labeled according to their angular momentum 85 and spin quantum numbers using terms symbols such as ${}^{4}I_{15/2}$, or ${}^{4}S_{3/2}$. Here the 86 letter refers to the total orbital angular momentum of the ion, the superscript is the 87 number of possible orientation of the total spin of the ions, given as 2S + 1, where 88S is the total spin angular momentum. The subscript refers to the total angular 89 momentum of the ion and is determined using the Russell-Saunders coupling 90 scheme [19]. These levels represent the energy levels of a free ion. However, when 91 the ion is in a host, electron-host interactions further split these levels into Stark 92 sublevels, due to the effect of the electric field of the matrix (crystal field effect). 93 This effect is quite small since the 4f orbitals are shielded from the environment 94 by the filled f of sub-shells. Nevertheless, the crystal field (CF) induced by 95 the host is what determines the shape of the absorption and emission spectra of RE $_{96}$ ions. When comparing the spectra of RE ions in crystals and glasses, the absorption 97 and emission bands are narrow with well resolved Stark components, even at room 98 temperature, in the case of crystals when RE ions occupy a well defined crystal field 99 site. On the contrary, the spectra of RE ions in an amorphous matrix, like a glass, 100 present an inhomogeneous broadening due to the different sites occupied by the ions 101 in the host and thus, different crystal fields. As a consequence the Stark structure of 102 RE spectra in glasses is only partially resolved even at low temperature. 103

On the other hand, the symmetry and strength of the CF also affect the splitting 104 of the levels. Moreover, the probability of radiative and nonradiative transitions is 105 strongly affected by the host matrix.

The radiative transition probabilities between energy levels in rare-earth ions 107 can be calculated by using the Judd-Ofelt theory [20, 21]. In the framework of this 108



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theory, and considering the electric-dipole and magnetic-dipole contributions, the 109 radiative transitions within the $4f^n$ configuration from the initial state |a J>, to the 110 final state |b J'>can be described in terms of the oscillator strength by using the 111 expression, 112

$$f_{cal}(aJ; bJ') = \frac{8\pi^2 m\nu}{3h(2J+1)e^2n^2} \left[\chi_{ed} S_{ed}(aJ; bJ') + \chi_{md} S_{md}(aJ; bJ') \right]$$
(15.1)

where m is the mass of the electron, *h* the Planck's constant, *v* is the frequency of 113 the transition, *J* is the total angular momentum of the initial state, *e* the electron 114 charge, and *n* is the refractive index of the host. $\chi_{ed} = \frac{n(n^2+2)^2}{9}$ and $\chi_{md} = n^3$ are 115 the effective field correction at a well-localized center in a medium of refractive 116 index *n*. *S*_{ed} and *S*_{md} are the line strengths for electric-dipole and magnetic-dipole 117 transitions respectively, expressed by, 118

$$S_{ed} = e^{2} \sum_{t=2,4,6} \Omega_{t} |\langle aJ \| U^{(t)} \| bJ' \rangle|^{2}$$
(15.2)

$$S_{md} = \frac{e^2 \hbar^2}{4m^2 c^2} \left| \left\langle aJ \right\| \overrightarrow{L} + 2\overrightarrow{S} \right\| bJ' \right\rangle \right|^2$$
(15.3)

 $\langle || U^{(t)} || \rangle$ are the double-reduced matrix elements of the unit tensor operators which 119 are considered to be independent of the host matrix and Ω_t are the JO parameters. 120

The calculated oscillator strengths can be compared with the experimental ones 121 obtained from the absorption spectra by, 122

$$f_{\exp} = \frac{mc}{\pi e^2 N} \int_{band} \alpha(v) dv$$
(15.4)

where N is the number of ions per unit volume and $\alpha(v)$ is the absorption coefficient. ¹²³ Since the experimental oscillator strength contains electric-dipole and magneticdipole contributions, the latter has to be subtracted from the experimental one to ¹²⁵ obtain the electric-dipole contribution. The magnetic-dipole contribution, f_{md} , can ¹²⁶ be obtained from the equation $f_{md} = nf'$ [22], where *n* is the refractive index of the ¹²⁷ studied sample and f' is a quantity calculated based on the energy-level parameters ¹²⁸ for lanthanide aquo ions [23].

The JO parameters are derived from the electric-dipole experimental oscillator ¹³⁰ strengths by using a least squared fitting between experimental and calculated ¹³¹ oscillator strengths. The magnetic dipole contribution is only present in transitions ¹³² with $\Delta S = \Delta L = 0$ and $\Delta J = 0, \pm 1$ [23]. ¹³³

The radiative transitions probability for electric- and magnetic-dipole transitions 134 are given by, 135

$$A(aJ;bJ') = A_{ed} + A_{md} = \frac{64\pi^4 v^3}{3h(2J+1)c^3} \left[n\frac{(n^2+2)^2}{9} S_{ed} + n^3 S_{md} \right]$$
(15.5)

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The magnetic-dipole contributions can be calculated using the values given by 136 Weber for LaF₃ [24] and correcting for the refractive index, 137

$$A_{md} = \frac{n}{n'} A'_{md} \tag{15.6}$$

where *n* and A_{md} are the values reported for LaF₃.

The radiative lifetime of an emitting level is related to the total spontaneous ¹³⁹ emission probability from this level by, ¹⁴⁰

$$\tau_R = \left\{ \sum_{bJ'} A\left(aJ; bJ'\right) \right\}^{-1} \tag{15.7}$$

The emission branching ratio of a transition is defined by,

$$\beta = \frac{A \left(aJ; bJ' \right)}{\sum_{bJ'} A \left(aJ; bJ' \right)}$$
(15.8)

The experimental lifetimes and the radiative lifetimes can be related by,

$$\frac{1}{\tau_{\exp}} = \frac{1}{\tau_R} + W_{NR} = \frac{1}{\tau_R} + W_{MPH} + W_{ET}$$
(15.9)

where W_{NR} is the nonradiative probability. This nonradiative probability can be due 143 to multiphonon relaxation, which depends on the host matrix, and/or energy transfer 144 which is dependent on ion concentration. 145

Finally, one important parameter for laser applications and optical amplification 146 is the emission cross-section. The stimulated emission cross-section can be obtained 147 from the absorption spectrum by using the McCumber approach [25], which relates 148 the absorption and emission cross-sections by, 149

$$\sigma_{em}(\nu) = \sigma_a(\nu) \exp\left[\frac{(\varepsilon - h\nu)}{KT}\right]$$
(15.10)

where σ_a and σ_{em} are the absorption and emission cross-sections respectively, ν ¹⁵⁰ is the photon frequency, ε is the net free energy required to excite one Er^{3+} ion ¹⁵¹ from the ⁴I_{15/2} to the ⁴I_{13/2} state at temperature T, h is the Planck's constant, ¹⁵² and K is the Boltzmann constant. The absorption cross-section is the absorption ¹⁵³ coefficient divided by the concentration of active ions and ε can be determined ¹⁵⁴ by using the simplified procedure provided in reference 26. The effective emission ¹⁵⁵ cross-section can also be calculated from the experimental emission spectrum and ¹⁵⁶ radiative transition probability by using the following expression [27], ¹⁵⁷

$$\sigma_{eff} = \frac{\lambda_p^4}{8\pi n^2 c} \frac{1}{\Delta \lambda_{eff}} A\left(aJ; bJ'\right)$$
(15.11)

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142

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where *c* is the velocity of light, λ_p is the peak fluorescence wavelength, $\Delta \lambda_{eff}$ is the 158 effective linewidth, *A* is the radiative transition probability of the emitting level, and 159 *n* the refractive index. The effective linewidth of the emission is defined by, 160

$$\Delta \lambda_{eff} = \int \frac{I(\lambda) \, d\lambda}{I_{\text{max}}} \tag{15.12}$$

15.3 Structural Characterization

Precursor oxyfluoride tellurite glass with a composition of 73.6 TeO₂–17.6 ZnO–8.8 ¹⁶² ZnF₂ mol% doped with 1 wt% of ErF₃, which corresponds to $1.4 \cdot 10^{20}$ ions/cm³, ¹⁶³ was prepared by the conventional melt-quenching technique. To nucleate a glass- ¹⁶⁴ ceramic (GC) phase a two step treatment was performed according to the values of ¹⁶⁵ the glass transition and crystallization temperatures ($T_g \approx 293$ °C and $T_c \approx 390$ °C ¹⁶⁶ respectively) [18]. The first treatment was done at 310 °C for 10 h, followed by a ¹⁶⁷ 3 h treatment at 340 °C, which is below that of the onset of crystallization. ¹⁶⁸

The microstructure of the samples and chemical composition of the nanocrystals 169 have been analyzed by Transmission Electron Microscopy (TEM) and Energy 170 Dispersive X-ray spectroscopy (EDX). TEM analysis clearly demonstrates the 171 formation of nanocrystals in the heat treated sample as it is shown in Fig. 15.1a. 172 They present a dendritic structure with typical sizes of 45 ± 10 nm and clear 173 lattice fringes. The Fast Fourier Transform (FFT) of this image in Fig. 15.1b shows 174 different brilliant spots in the reciprocal space which can be indexed according to the 175 planes of ErF₃ crystals obtained in the [101] direction. This analysis was performed 176 based on the compositional profile obtained from EDX spectroscopy. The EDX 177 analysis of the nanocrystals (Fig. 15.2) shows intense peaks corresponding to F and 178



Fig. 15.1 (a) HRTEM image of the GC sample. (b) Fast Fourier Transform (FFT) image indexed according to the ${\rm ErF}_3$ structure



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Er in addition to those associated to the C layer of the replica and the Cu grid (C, $_{179}$ O, and Cu). The absence of peaks related to Zn or Te suggests that the nanocrystals $_{180}$ are formed by ErF_3 .

15.4 Influence of the Nanocrystals on the Luminescence Properties of Er³⁺ Ions

15.4.1 Absorption Properties

The influence of the nanocrystals on the spectroscopic properties of Er^{3+} ions in 185 glass-ceramic samples based on fluorotellurite glasses has been studied by using 186 absorption and steady-state and time-resolved laser spectroscopy. The absorption 187 spectra of the glass (GS) and glass-ceramic (GC) samples are displayed in Fig. 15.3. 188 The absorption bands are assigned to the transitions from the ${}^{4}I_{15/2}$ ground state to 189 the excited states of Er^{3+} ions. Due to the absorption of the host in the UV, some 190 transitions are not observed. As can be seen, there are some difference between 191 both spectra due to the presence of the nanocrystals (NCs). The absorption edge 192 is shifted to longer wavelengths in the GC sample due to the scattering of short 193 wavelength light by the NCs present in this sample. Moreover, the intensity of 194 the hypersensitive transition ${}^{4}I_{15/2} \rightarrow {}^{2}H_{11/2}$ is strongly reduced in the glass-ceramic 195 sample which indicates that the ligand field around Er³⁺ ions has changed as a 196 consequence of the heat treatment. The strength of the hypersensitive transitions is 197 greatly affected by the environment of rare earth ions. Ligand polarizability as well 198 as an asymmetric surrounding environment increases the intensity of hypersensitive 199 transitions of RE ions [28–30]. Another evidence of the incorporation of Er^{3+} ions 200 in the nanocrystals is the splitting of the inhomogeneously broadened absorption 201 bands in the absorption spectrum of the GC sample. As an example, the inset in 202 Fig. 15.3 shows a detail of the ${}^{4}I_{15/2} \rightarrow {}^{4}I_{13/2}$ absorption band for the glass and GC 203 samples. 204

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 Table 15.1
 Judd-Ofelt parameters and r.m.s. deviation for precursor glass (GS) and glass ceramic (GC) samples

Sample	$\Omega_2 (\times 10^{-20})$	$\Omega_4 (\times 10^{-20})$	$\Omega_6 (\times 10^{-20})$	r.m.s.	
GS	4.71	1.57	1.13	$3.24 \cdot 10^{-7}$	t3.1
GC	1.28	1.04	1.00	$3.47 \cdot 10^{-7}$	t3.2

The absorption spectra have been used to calculate the radiative transition 205 probabilities, the radiative lifetimes, and branching ratios of the emissions by 206 using the JO intensity parameters. From these spectra the experimental oscillator 207 strengths have been calculated and following the procedure described in Sect. 2, the 208 theoretical oscillator strengths and the JO parameters of Er^{3+} ions in the glass and 209 glass-ceramic samples. The values are displayed in Table 15.1. The matrix elements 210 given by Carnall et al [23] have been used in the calculation. The low value of the 211 root mean square (rms) indicates a good agreement between the experimental and 212 calculated oscillator strengths.

The value of Ω_2 in the GC sample is lower than in the GS, which means that the 214 heat treatment induces a reduction in the degree of covalency in the rare-earth site. It 215 is well known that Ω_2 is most sensitive to local structure and its value is indicative of 216 the amount of covalent bonding between RE ions and ligand anions. The decrease of 217 Ω_2 further confirms the incorporation of Er^{3+} ions into the fluoride NCs. Moreover, 218 the sum of the JO parameters decreases for the GC sample due to the decrease of 219 covalency of the chemical bond between the Er^{3+} ion and the ligand anions. This 220 is in accordance with reported results that predict that the sum decrease in the order 221 oxide > oxyfluoride > fluoride. In addition, the Ω_2 parameter is closely related to 222 the hypersensitive transitions. The more intense the hypersensitive transition is, the 223 larger the value of Ω_2 is. The lower value of this parameter in the heat treated sample 224 is in agreement with the decrease of the strength of the hypersensitive transition 225 ${}^{4}I_{15/2} \rightarrow {}^{2}H_{11/2}$ which indicates the variation of the local structure around the Er³⁺ 226 ions and further confirms their incorporation into the nanocrystals. 227

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The JO parameters have been used to calculate the radiative transition probabilities and radiative lifetimes. The radiative lifetimes of level ${}^{4}I_{13/2}$ are 3.23 and 229 3.73 ms for the GS and GC sample respectively. 230

15.4.2 Near Infrared Emission

The room temperature near infrared emission spectra have been obtained by exciting 232 at 801 nm in the ${}^{4}I_{9/2}$ level with a Ti-sapphire ring laser. The fluorescence was 233 analyzed with a monochromator and the signal detected with a photomultiplier 234 and finally amplified by a standard lock-in technique. After excitation of level 235 ${}^{4}I_{9/2}$, the next lower levels are populated by multiphonon relaxation. Two emission 236 bands were observed at around 980 and 1530 nm, which correspond to transitions 237 from ${}^{4}I_{11/2}$ and ${}^{4}I_{13/2}$ levels to the ground state respectively. Figure 15.4 shows 238 the emission spectra corresponding to the ${}^{4}I_{13/2} \rightarrow {}^{4}I_{15/2}$ transition. The spectrum 239 of the GC sample shows a more resolved structure with seven Stark components. 240 Moreover, the effective bandwidth is reduced from 73.4 nm in the Gs to 64 nm in 241 the GC sample probably due to the reduction of inhomogeneous broadening. 242

Figure 15.5 shows the absorption and emission cross-sections obtained for 243 the GC sample by using the McCumber approach together with the normalized 244 measured emission spectrum. The stimulated emission cross-section has a value of 245 $8.9 \cdot 10^{-21}$ cm², similar to the one found in the glass sample ($8.8 \cdot 10^{-21}$ cm²) [16] 246 and higher than those found in other fluorotellurite glass-ceramics [31]. 247

The comparison of the excitation spectra of the ${}^{4}I_{15/2} \rightarrow {}^{4}I_{9/2}$ transition obtained 248 by collecting the luminescence at 1540 nm for the glass and glass-ceramic samples 249 also confirms the incorporation of Er^{3+} ions in the nanocrystals. As can be seen in 250



Fig. 15.4 Room temperature emission spectra obtained under excitation at 801 nm in the glass and glass-ceramic samples





Fig. 15.6 Excitation spectra of the ${}^{4}I_{15/2} \rightarrow {}^{4}I_{9/2}$ transition obtained by collecting the luminescence at 1540 nm in the glass and glass-ceramic samples at 295 K and 9 K

Fig. 15.6, the spectrum corresponding to the GC sample shows well defined peaks $_{251}$ not observed in the excitation spectrum of the glass sample. This is more evident $_{252}$ from the low temperature excitation spectrum of the GC sample which shows $_{253}$ narrow peaks according with the presence of Er^{3+} ions in a crystalline environment. $_{254}$

The experimental decays of luminescence from levels ${}^{4}I_{11/2}$ and ${}^{4}I_{13/2}$ were 255 obtained at room temperature for both samples by exciting at 801 nm in level ${}^{4}I_{9/2}$ 256 by using a Ti-sapphire laser pumped by a pulsed frequency doubled Nd:YAG laser 257 (9 ns pulse width). The decay from level ${}^{4}I_{13/2}$ shows, in both samples, an initial 258 rise time due to the population from the higher ${}^{4}I_{11/2}$ level. Figure 15.7 shows the 259 experimental decay curves for GC and GS samples. The decay of the glass sample 260 can be described by a single exponential function with a lifetime of 3.6 ms, however 261 the experimental decay of the GC sample is well described by a double exponential 262 function with lifetimes of 1.2 and 3.1 ms which suggest contributions from different 263 environments around Er^{3+} ions. In the GC, a fraction of Er^{3+} ions remains in a 264 glass environment while the rest is forming NCs. The reduction in lifetimes after 265 heat treatment has been previously reported [32] and attributed to the segregation 266



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of the rare-earth ions into the crystalline phase which reduces $Er^{3+}-Er^{3+}$ distances. ²⁶⁷ Shorter distances between Er^{3+} ions favour dipole-dipole energy transfer between ²⁶⁸ them, which increases non-radiative rates and reduces the lifetime of level ⁴I_{13/2}. A ²⁶⁹ similar behavior is observed for the lifetime value of the ⁴I_{11/2} level which is reduced ²⁷⁰ from 420 µs in the glass sample to 220 µs in GC sample. ²⁷¹

15.4.3 Upconversion Emission

Up-conversion (UC) is an anti-Stokes process that converts the absorbed lower 273 energy photons, usually in the NIR range, into higher energy photons in UV, visible, 274 or shorter NIR ranges. UC can occur mainly by two fundamental mechanisms: (i) 275 excited state absorption (ESA); (ii) energy transfer upconversion (ETU). In the case 276 of ESA, excitation involves sequential absorptions of pump photons by a single 277 ion. This mechanism requires a nearly equal separation for the first and second 278 steps, as well as the long lifetime of the intermediate level. In an ETU process, 279 two ions interact, one of then loses energy and goes to the ground state whereas 280 the other one gains energy and goes to the upper level. The UC efficiency of an 281 ETU process is strongly dependent on the active ions concentration. In contrast to 282 ETU, the efficiency of an ESA process is independent of the dopant concentration, 283 because ESA involves sequential excitation in the same ion.

Visible upconversion emission has been observed for both glass and GC samples the resonance with the ${}^{4}I_{9/2}$ level. The upconverted emission spectra obtained under cm tasks to remove the pumping radiation. Figure 15.8a shows the upconversion emission the glass sample shows the characteristic green emission attributed 291 cm tasks to remove the glass sample shows the characteristic green emission attributed 291 cm tasks to remove the glass sample shows the characteristic green emission attributed 291 cm tasks to remove the glass sample shows the characteristic green emission attributed 291 cm tasks to remove the glass sample shows the characteristic green emission attributed 291 cm tasks the task tasks the tasks task

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Fig. 15.8 Visible emission obtained (a) under 801 nm and (b) under 488 nm excitation for the glass and glass-ceramic samples

to the transitions from the two thermallized levels (${}^{2}H_{11/2}$, ${}^{4}S_{3/2}$) together with a 292 weak red emission from the ${}^{4}F_{9/2}$ level. However, the spectrum of the GC sample 293 shows in addition to the green and red emissions very weak blue emissions from 294 higher levels which do not appear in the upconversion emission of the glass sample. 295 As can be seen, the emission bands are better resolved in the GC spectrum in 296 agreement with the incorporation of Er^{3+} ions in the crystalline phase. Moreover, 297 the intensity of the red emission increases significantly in the GC sample. This 298 increase of the red emission, not observed in the spectra obtained by exciting at 299 488 nm (Fig. 15.8b), indicates that after NIR excitation, level ${}^{4}F_{9/2}$ is populated by 300 energy transfer processes in addition to multiphonon relaxation from level ${}^{4}S_{3/2}$. 301 This behavior could be associated to a higher concentration of Er^{3+} ions in the 302 nanocrystals, which reduces the Er^{3+} - Er^{3+} distances and increases the probability 303 of energy transfer processes [33, 34].

To investigate the excitation mechanisms for populating the ${}^{4}S_{3/2}$ and ${}^{4}F_{9/2}$ levels 305 after IR excitation in the GC sample, we have measured the upconverted emission 306 intensities for different pumping powers. The upconversion emission intensity (I_{em}) 307 depends on the incident pump power (P_{pump}) according to the relation I_{em} \propto (P_{pump})ⁿ, 308 where n is the number of photons involved in the pumping mechanism. Figure 15.9 309 shows a logarithmic plot of the integrated emission intensity of the upconverted 310 green and red fluorescence as a function of the pump laser intensity. The dependence 311 in both cases is nearly quadratic which indicates a two photon (TP) upconversion 312 process to populate the ${}^{4}S_{3/2}$ and ${}^{4}F_{9/2}$ levels. This process may be associated with 313 excited state absorption and/or energy transfer upconversion. 314

As it is well known, the time evolution of the upconversion luminescence ³¹⁵ after an excitation pulse provides an useful tool in discerning which the operative ³¹⁶ mechanism is. The radiative ESA process occurs during the excitation pulse and ³¹⁷ leads to an immediate decay of the upconversion luminescence after excitation. ³¹⁸ Upconversion by energy transfer leads to a time-dependent emission that shows ³¹⁹ a rise of the upconverted population after the laser pulse, followed by a decay of ³²⁰



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the population, with a lifetime longer than the one after direct excitation. The rise 321 and decay times are determined by both the intermediate and the upper excited state 322 lifetimes. This distinction is possible when the pump pulse width is much shorter 323 than the time constant of the relevant energy transfer step. 324

According to the energy level diagram of Er^{3+} ion in the GC sample, the ³²⁵ ${}^{4}S_{3/2}$ level can be populated by ESA and/or ETU in the case of the green ³²⁶ emission(Fig. 15.10). In the first case, the absorption of an IR photon excites the ³²⁷ electrons to the ⁴I_{9/2} level, then multiphonon relaxation populates the next lower ³²⁸ level, and the absorption of a second IR photon promotes the electrons to the ⁴F_{3/2,5/2} ³²⁹ levels from where ⁴S_{3/2} level is reached by nonradiative relaxation. Part of the ³³⁰ excitation energy in the ⁴I_{11/2} level further relaxes, radiatively and nonradiatively ³³¹ to level ⁴I_{13/2} and ESA from ⁴I_{13/2} to ²H_{11/2} can occur. This last process is nearly ³³² resonant. Another possibility is an energy transfer process which involves the ³³³ energy and reaches level ⁴F_{7/2}, whereas the other one loses energy and goes to the ³³⁵ ground state.

The lifetime behavior of the upconverted green emission suggests that both 337 processes ESA and ETU are present. Figure 15.11a shows the decays of the 338 green emission obtained by exciting at 488 and 801 nm respectively for the GC 339 sample. In both cases, the decays can be described to a good approximation by two components. The fast one has a lifetime of 36 and 41 μ s for the 488 and 341 801 nm excitation respectively but the lifetime of the long component (129 μ s), 342 is longer than the one obtained by direct excitation (107 µs). The observation of 343 the instantaneous rise after the 9 ns pulse excitation indicates the presence of an 344 ESA process; however, the longer lifetime of the slow component after IR excitation 345 indicates that an additional ETU process is also necessary to describe the lifetime of 346 the decay. The lifetime of the long component $(129 \,\mu s)$ is about half the lifetime of 347 the ${}^{4}I_{11/2}$ level (220 μ s), which suggests that the upconversion green emission from 348 level ${}^{4}S_{3/2}$ can be produced by the interaction between two Er^{3+} ions in the ${}^{4}I_{11/2}$ 349 level via the transitions $({}^{4}I_{11/2} \rightarrow {}^{4}I_{15/2})$ and $({}^{4}I_{11/2} \rightarrow {}^{4}F_{7/2})$. 350

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Fig. 15.10 Energy level diagram of Er^{3+} ions in the GC sample and possible upconversion mechanisms



Fig. 15.11 Semilogarithmic plot of the experimental decay curves of the (**a**) green $({}^{4}S_{3/2})$ and (**b**) *red* $({}^{4}F_{9/2})$ emissions under excitation at 488 nm (*black line*) and 801 nm (*red line*) for the GC sample

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Instead, the time evolution of the upconverted red emission confirms that the 351 ${}^{4}F_{9/2}$ level is populated by ETU processes. As an example, the time evolution of 352 the red emission obtained under excitation at 801 nm and 488 nm for the GC 353 sample is shown in Fig. 15.11b. In both cases, the decays can be described by two components. The fast one ($\sim 21 \ \mu s$) is similar for both excitation wavelengths 355 whereas the lifetime of the slow component increases from $89 \,\mu s$ to $183 \,\mu s$ when the excitation wavelength changes from 488 nm to 801 nm. Moreover, there is an initial 357 rise time of about 15 μ s. The presence of a rise time together with the lengthening 358 of the slow component of the decay supports the hypothesis that ETU processes 359 are populating level ${}^{4}F_{9/2}$. According to the energy level diagram of Er^{3+} ions in 360 the GC sample (Fig. 15.10), there are different ETU processes to populate the ${}^{4}F_{9/2}$ 361 level. This level can be populated via $({}^{4}I_{9/2} \rightarrow {}^{4}I_{13/2})$ and $({}^{4}I_{11/2} \rightarrow {}^{4}F_{9/2})$ transitions 362 and/or $({}^{4}I_{11/2} \rightarrow {}^{4}I_{15/2})$ and $({}^{4}I_{13/2} \rightarrow {}^{4}F_{9/2})$. There exists another possible process for 363 populating level ${}^{4}F_{9/2}$ in which two Er³⁺ ions interact, one of them in level ${}^{4}I_{11/2}$ and 364 the other one in level ${}^{4}F_{7/2}$, going both to level ${}^{4}F_{9/2}$. By considering that the lifetime 365 of a higher-energy level excited by ETU reflects those of the intermediate levels from which upward excitation occurs, and that the long component of the decay of 367 the red emission after excitation at 801 nm is very close to the lifetime of level ${}^{4}I_{11/2}$, 368 the $({}^{4}I_{9/2} \rightarrow {}^{4}I_{13/2}); ({}^{4}I_{11/2} \rightarrow {}^{4}F_{9/2})$ and $({}^{4}F_{7/2} \rightarrow {}^{4}F_{9/2}); ({}^{4}I_{11/2} \rightarrow {}^{4}F_{9/2})$ processes seem 369 to be the likeliest ones to explain the population of the ${}^{4}F_{9/2}$ level by ETU. 370

15.5 Conclusions

Transparent glass-ceramics containing ErF_3 nanocrystals have been prepared by 372 suitable thermal treatment of an Er^{3+} - doped 73.6 TeO₂-17.6 ZnO-8.8 ZnF₂ mol% 373 fluorotellurite glass. 374

The comparison of the fluorescence properties of Er^{3+} -doped precursor glass 375 and glass-ceramic confirms the successful incorporation of the rare-earth into the 376 nanocrystals. The strong reduction of the Ω_2 JO parameter and the intensity of the 377 hypersensitive transition ${}^{4}I_{15/2} \rightarrow {}^{2}H_{11/2}$ in the heat treated sample indicate that Er^{3+} 378 ions are in a crystalline environment. The ${}^{4}I_{13/2}$ decay in the glass-ceramic sample 379 can be described by two single exponential functions which suggests that a fraction 380 of Er^{3+} ions are forming nanocrystals while the rest remains in a glass environment. 381 Moreover, the emission lifetime is shorter than in the glass sample probably due to 382 the segregation of the Er^{3+} ions into the crystalline phase which reduces Er^{3+} - Er^{3+} 383 distances and increases energy transfer processes. 384

The presence of ErF_3 nanocrystals has a strong effect in the upconverted red ³⁸⁵ emission from ${}^4\text{F}_{9/2}$ level. This emission increases significantly as compared with ³⁸⁶ the green one in the GC sample. The time evolution of the upconverted red emission ³⁸⁷ suggests that ETU processes are responsible for the increase of this emission in the ³⁸⁸ glass-ceramic sample. ³⁸⁹

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