



Exciton interaction with Ce³⁺ and Ce⁴⁺ ions in (LuGd)₃(Ga, Al)₅O₁₂ ceramics



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ARTICLE INFO

Keywords:

Garnet scintillators

Ce⁴⁺

Excitons

Energy transfer

Synchrotron

XANES

ABSTRACT

Scintillators based on Ce-doped garnets are regularly co-doped with Mg²⁺ or Ca²⁺ to form Ce ions in 4+ state and reduce undesired afterglow. However overly high Ce⁴⁺ concentration leads to poor light yield performance. In order to understand the reason for variation in luminescence efficiency of Ce³⁺- and Ce⁴⁺-doped garnets we investigate the differences in energy conversion processes in complex LuGd₂Ga₃Al₂O₁₂:Ce³⁺/Ce⁴⁺ ceramics by means of VUV synchrotron irradiation. At first we have established via transmission spectroscopy and X-ray absorption spectroscopy that LuGd₂Ga₃Al₂O₁₂:Ce, Mg sample contains cerium in the 4+ state only. Then we show with VUV spectroscopy efficient interaction of excitons with Gd³⁺ and Ce³⁺, and lack of exciton absorption edge in LuGd₂Ga₃Al₂O₁₂:Ce⁴⁺ excitation spectrum. Instead, Ce⁴⁺ exhibits charge-transfer absorption band in the range of exciton emission. We suggest that when Ce⁴⁺ concentration becomes too high, the exciton → Gd³⁺ → Ce³⁺ energy transfer path is hindered. It leads to high intensity of Gd³⁺ luminescence in Lu₁Gd₂Ga₃Al₂O₁₂:Ce, Mg ceramics, but lowered Ce³⁺ X-ray excited luminescence. Fine balance between 3+ and 4+ Ce concentrations is necessary to achieve the best performance of garnet scintillators.

1. Introduction

Cerium doped complex garnets are actively investigated for scintillator [1] and persistent phosphor [2] applications. Modification of (Lu, Gd, Y)₃(Ga, Al)₅O₁₂:Ce³⁺ cation composition [3] allows fine-tuning of light yield, emission wavelength, level of afterglow, radiation absorption efficiency, luminescence temperature stability and etc. [4,5]. Adding small amounts of divalent ions like Mg²⁺ or Ca²⁺ can lead to lower levels of afterglow (by an order of magnitude) [6,7], and very fast (<50 ps) luminescence rise time under X-rays [8].

The improvement of timing characteristics of YAG:Ce, Mg [9] and LYSO:Ce, Mg [10] is ascribed to favorable change in charge migration processes by formation of Ce⁴⁺ (charge compensated with Mg²⁺). A

model of e-h interaction with Ce³⁺ and Ce⁴⁺ ions explains the extremely fast rise time kinetics [6] and low afterglow levels. The corresponding processes proceed as follows [11]:



According to processes (1)–(3), Ce³⁺ radiatively relaxes after two sequent events of capturing a hole (1) and afterwards an electron (2). The delay of electron capture by process (1) means that Ce³⁺ is not able to compete with electron traps for the initial volley of CB-electrons [12].

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Contrary to Ce³⁺, equilibrium Ce⁴⁺ (charge compensated) is available for immediate capture of electrons (2) directly leading to luminescence (3) [13]. Furthermore, Ce⁴⁺, as Coulomb-active center in a 3+ lattice, provides efficient competition to electron traps [14]. Diminished amount of trapping then leads to increased light yield for LuAG:Ce, Mg [15]. Interestingly, increased light yield of LuAG:Ce, Mg is observed only for low concentrations of Mg co-doping. Likewise, in GGAG:Ce, Ca [16], GGAG:Ce, Mg [17,18] co-doping with Ca²⁺, Mg²⁺ above 0.1% leads to lower light yield by factor of 1.3–2.

The model depicted above includes only e-h recombination on Ce^{3+/4+} and does not account for excitons formation and their interactions with Ce and Gd ions. In garnets of simpler compositions, e.g. YAG exciton emission is detected as a broad UV emission band around 270 nm [19], while excitons distorted by antisite defects emit at around 300–350 nm [20]. The absorption bands of Ce³⁺ (4f-5d₂) overlap well with excitonic emission. Indeed, doping YAG with RE ions significantly distorts and quenches exciton emission (see e.g. Ref. [21] for Ce³⁺ or [22] for Pr³⁺), indicating energy transfer to impurity RE ions.

In complex solid solutions of (Gd,Y)₃(Al,Ga,Sc)₅O₁₂ [23,24], the excitons emit at similar wavelengths of 250–350 nm, but with higher Gd³⁺ content their emission goes down in intensity. As such, in Gd₃(Ga, Al)₅O₁₂:Ce crystals no emission from excitons was detected so far, while excitation spectra of Gd³⁺ and Ce³⁺ still contained excitation peaks characteristic for excitons [25]. That can be attributed to complete energy transfer from excitons to Gd³⁺ ions.

Ce⁴⁺ also has an absorption band in the UV range due to charge transfer (CT) transition from the O²⁻ states of valence band to the Ce⁴⁺ ground state [13]. The CT absorption of Ce⁴⁺ does not lead to emission. Addition of Ce⁴⁺ potentially creates an exciton quenching channel (by CT absorption) that lowers the probability of exciton → Ce^{3+/4+}/Gd³⁺ energy transfer. Thus we consider it important to study exciton interplay with Ce³⁺, Ce⁴⁺ and Gd³⁺ ions in complex garnets.

In the current work we investigate the differences in energy migration processes in complex Lu₃Ga₃Al₂O₁₂ and LuGd₂Ga₃Al₂O₁₂ garnet ceramics doped with Ce³⁺ or Ce⁴⁺ ions. We first develop an understanding on broad band UV emission in Lu₃Ga₃Al₂O₁₂:Ce garnet ceramics and lack of it in Gd³⁺-containing garnets. With X-ray Absorption Near Edge Structure (XANES) and transmission spectroscopy we establish that Mg²⁺ co-doping fully converts Ce to 4+ state in LuGd₂Ga₃Al₂O₁₂:Ce, Mg samples. Then we study the differences in excitation spectra of LuGd₂Ga₃Al₂O₁₂:Ce⁴⁺ and LuGd₂Ga₃Al₂O₁₂:Ce³⁺ luminescence in the 4.5–10 eV range under synchrotron irradiation.

1.1. Materials and experimental

Luminescence spectroscopy was conducted at photoluminescence endstation [26,27] of the FinEstBeAMS undulator beamline [28] of MAX IV synchrotron (Lund, Sweden) located at the 1.5 GeV storage ring. The range of utilized excitation energy for this work was 4.5–45 eV, while temperature was varied from 7 to 300 K. The excitation spectra were corrected for beamline photon flux by measuring a reference curve with AXUV-100G diode. In order to suppress high orders of the undulator excitation passing through the monochromator a set of the filters (SiO₂, MgF₂ and Al) were selected. Luminescence detection in UV-visible spectral range (200–800 nm) was performed by an Andor Shamrock (SR-303i) spectrometer equipped with 8259-01 Hamamatsu photon counting head. The emission spectra were corrected for the spectral sensitivity of the detection system.

XANES experiments were carried out on the Balder beamline [29] of MAX IV Laboratory (Lund, Sweden) located at the 3 GeV storage ring. The XANES spectra were measured in fluorescence detection mode by a 7-element silicon drift detector. Continuous energy scanning was performed at a speed ~3.5 min/XANES. For each sample, 10 repeats were collected and afterwards accumulated into a resulting spectrum. The reference CeO₂ sample was measured in transmission mode in order to avoid self-absorption distortion.

Absorption spectra were recorded with a double-beam PerkinElmer Lambda 950 UV/vis/NIR spectrometer. The transparent ceramic samples were placed in one optical path of the deuterium/halogen lamp without placing a reference sample in the second path. The transmitted beams were detected by a PMT. Spectra were acquired with a resolution of 0.5 nm within a spectral range of 200–700 nm. Below 330 nm the deuterium lamp was used, which automatically switched to the halogen lamp above 330 nm.

X-ray excited luminescence spectra were measured under continuous X-ray (40 kV, 10 mA, 3 cm distance) excitation. Emission spectra were registered in a reflection geometry using a Lomo Photonica MDR-2 monochromator (0.3 nm resolution) coupled to a Hamamatsu H8259-01 photon counting head. The spectra were corrected for wavelength-dependent transmission of the monochromator and the spectral sensitivity of the PMT.

Ceramic Lu₃Ga₃Al₂O₁₂:Ce 0.2 mol.% and Lu₁Gd₂Ga₃Al₂O₁₂:Ce 0.2 mol.% garnet samples for this study were prepared at Philips Research Eindhoven by mixing the initial oxides Gd₂O₃, Lu₂O₃, Ga₂O₃, CeO₂ and Al₂O₃, purity no less than 99.99% (4 N), with a dispersing agent and distilled water. This slurry is then milled for 100 h on a roller bench in a plastic jar using 2 mm Al₂O₃ balls. After grinding, organic binders were added to the slurry, and the suspension was then dried in a drying chamber. The dried granulate was sieved using a metal sieve with a mesh size of <500 µm and then dry-pressed in a uniaxial press into ‘green-body’ pellets. After pressing, the resulting pellets were heat treated to burn off the organic binders. The pressed green-body pellets were then sintered for 8 h in an Astro Industries Inc. vacuum oven at a temperature of 1600–1750 °C, under high vacuum (10⁻⁵ to 10⁻⁶ mbar) or in oxygen atmosphere. The final ceramics are in the form of pills of 14 mm diameter and 1 mm thickness. Based on the X-ray diffraction patterns it was concluded that all samples consisted of a single garnet phase. One Lu₁Gd₂Ga₃Al₂O₁₂:Ce sample was co-doped 0.2 mol.% Mg. With transmission spectroscopy and XANES we established that the LuGd₂Ga₃Al₂O₁₂:Ce, Mg 0.2% sample had Ce in the 4+ state only, Fig. 1.

The XANES peak shape and Ce L_{III}-edge positions were compared for LuGd₂Ga₃Al₂O₁₂:Ce, LuGd₂Ga₃Al₂O₁₂:Ce, Mg and CeO₂ samples. The LuGd₂Ga₃Al₂O₁₂ singly doped with Ce exhibited an absorption peak at 5726 eV, showing dominant presence of Ce³⁺ [30]. The CeO₂ XANES profile showed two main peaks at 5731 and 5738 eV, caused by the interaction of hybridized 4f, 5d orbitals of Ce with O 2p orbitals of the nearest surrounding [31,32]. The XANES spectrum for LuGd₂Ga₃Al₂O₁₂:Ce, Mg repeated the one for CeO₂ suggesting complete conversion of Ce³⁺ to Ce⁴⁺ in Mg co-doped ceramics. From in-line absorption spectra, Fig. 1b, similar conclusion could be drawn. LuGd₂Ga₃Al₂O₁₂:Ce sample exhibited two absorption bands with maxima at 2.75 eV (450 nm) and 3.65 eV (340 nm) belonging to 4f-5d_{1,2} spin and parity allowed transitions. After co-doping with large concentration of Mg, LuGd₂Ga₃Al₂O₁₂:Ce, Mg sample showed no 5d₁ absorption of Ce³⁺, instead there was a strong CT absorption band of Ce⁴⁺ below 3.55 eV (350 nm) [13,33].

1.2. Results and discussion

1.2.1. Excitons, Ce³⁺ and Gd³⁺ excitation in mixed garnets

We first focus on emission of excitons, Gd³⁺ and Ce³⁺ and respective VUV excitation spectra to determine their interaction with each other. In Fig. 2a the emission spectra of Lu₃Ga₃Al₂O₁₂:Ce 0.2% and LuGd₂Ga₃Al₂O₁₂:Ce 0.2% samples under 6.4 eV excitation by synchrotron irradiation at 7 K are shown. The Lu₃Ga₃Al₂O₁₂:Ce 0.2% spectrum (orange curve) exhibits the double emission band of Ce³⁺ 5d-4f transitions at around 2.5 eV (500 nm) and the UV emission band at 4.6 eV (270 nm). The broad UV luminescence band in YAG and LuAG:Ce has been determined with optical and electron-paramagnetic resonance (EPR) methods as exciton related [19,21,34]. In similar solid solutions of e.g. (Y,Lu)₃(Al,Ga)₅O₁₂ [35], the excitons also emit at energies of 3.5–5 eV. For Gd-containing garnets the emission spectrum shows different features in the UV range. Instead of broad excitonic emission band the

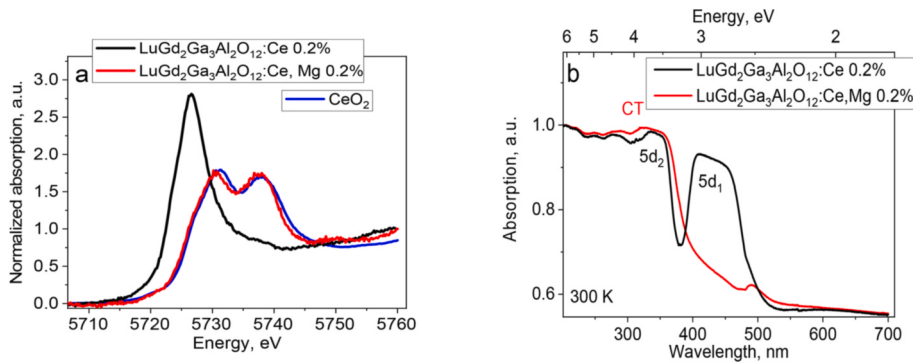


Fig. 1. a) XANES spectra performed at RT for LuGd₂Ga₃Al₂O₁₂:Ce and LuGd₂Ga₃Al₂O₁₂:Ce, Mg samples. For the reference CeO₂ measurement is shown. b) In-line transmission spectra for LuGd₂Ga₃Al₂O₁₂:Ce and LuGd₂Ga₃Al₂O₁₂:Ce, Mg measured at RT.

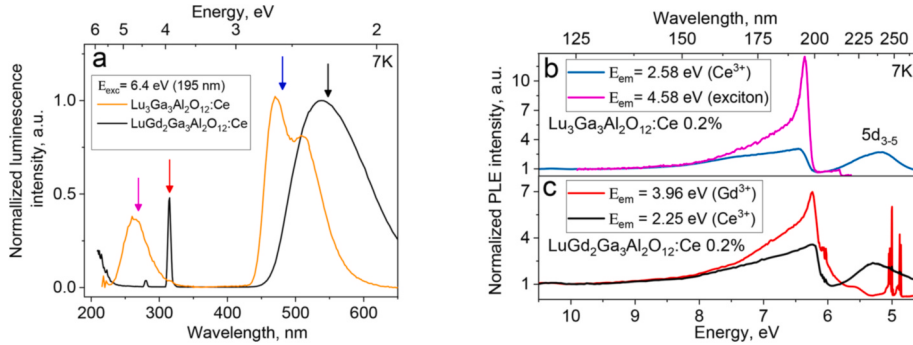


Fig. 2. a) Emission spectra of Lu₃Ga₃Al₂O₁₂:Ce 0.2% and LuGd₂Ga₃Al₂O₁₂:Ce 0.2% under 6.4 eV excitation at 7 K. The arrows indicate the monitored emission energy E_{em} for excitation spectra measurements in Fig. 2b and c b) Excitation spectra of UV emission band (4.58 eV, 270 nm) and Ce³⁺ (2.58 eV, 480 nm) in Lu₃Ga₃Al₂O₁₂:Ce 0.2% measured at 7K. c) Excitation spectra of Gd³⁺ (3.96 eV, 313 nm) and Ce³⁺ (2.25 eV, 550 nm) emission of LuGd₂Ga₃Al₂O₁₂:Ce 0.2% at 7K.

spectrum at 7 K is dominated by $^8S_{7/2} \rightarrow ^6P_J$, $^8S_{7/2} \rightarrow ^6I_J$ f-f transitions of Gd³⁺ 4.51 and 3.96 eV (275 and 313 nm, respectively) for LuGd₂Ga₃Al₂O₁₂:Ce 0.2%, black curve. It has been shown that excitonic emission can be distorted/quenched by the absorption bands of doped species (Ce³⁺ [21], Nd³⁺ [36], Pr³⁺ [22], Gd³⁺ [25,37]), which leads to exciton \rightarrow dopant energy transfer. For temperature dependence of the LuGd₂Ga₃Al₂O₁₂:Ce 0.2% emission spectra and energy transfer between Gd³⁺ and Ce³⁺ please see supporting information.

Fig. 2b shows VUV excitation spectra of Ce³⁺ emission at 480 nm and of UV emission band at 4.58 eV (270 nm) in Lu₃Ga₃Al₂O₁₂:Ce 0.2%. Ce³⁺ spectrum exhibits a 5d₃₋₅ ($^2D_{5/2}$, t_{2g} unresolved triplet state) excitation band [38] and the fundamental absorption edge that continues into the region of interband transitions (above \sim 7 eV, 180 nm). The 4.58 eV emission band is only excited effectively above 6.3 eV with a sharp excitation edge, which is a distinctive shape of excitation spectrum for excitons [39]. For more detail on the dependence of the excitation spectra of excitons on monitored emission wavelength please see supporting information.

In Fig. 2c the excitation spectra for Gd³⁺ and Ce³⁺ in LuGd₂Ga₃Al₂O₁₂:Ce 0.2% are shown. Gd³⁺ excitation spectrum exhibits several lines and a sharp absorption edge at 6.3 eV, while Ce³⁺ spectrum exhibits the same lines of Gd³⁺ transitions, the same absorption edge and an additional excitation band at 5.2 eV. The Gd³⁺ lines at 4.85 eV (255 nm), 5.02 eV (247 nm) and 6.05 eV (205 nm) are due to $^8S_{7/2} \rightarrow ^6D_{9/2}$, $^8S_{7/2} \rightarrow ^6D_{7/2}$ and $^8S_{7/2} \rightarrow ^6G_J$ f-f transitions, respectively [40]. Existence of these lines in the excitation spectrum of Ce³⁺ indicates Gd³⁺-Ce³⁺ energy transfer [41]. The 5.2 eV (240 nm) excitation band of Ce³⁺ belongs to 5d₃₋₅ excitation band [42] (same as Fig. 2b, blue).

The excitation spectra of Ce³⁺ and Gd³⁺ emission in LuGd₂Ga₃Al₂O₁₂:Ce 0.2% from Fig. 2c indicate interaction between excitons and Ce³⁺/Gd³⁺. The sharpness of the absorption edge at 6.3 eV in both

excitation spectra is due to exciton creation and its localization at Gd³⁺ or Ce³⁺ [21,43,44]. At 6.3 eV the contribution from excitons to Gd³⁺ is much more significant than to Ce³⁺. That suggests a stronger interaction of excitons with Gd³⁺ instead of Ce³⁺, most probably in view of Gd/Ce concentration ratio.

To summarise: Gd-free garnets show a UV emission band of significant intensity, the corresponding sharp excitation band at 6.3 eV allows to attribute the UV emission to formation of excitons. In garnets containing Gd³⁺ (LuGd₂Ga₃Al₂O₁₂:Ce) excitonic emission is quenched, while the excitation spectra of Gd³⁺ and Ce³⁺ still show the sharp excitation edge at 6.3 eV. That supports the notion that excitons transfer their energy to Gd³⁺ and Ce³⁺ ions.

1.2.2. Ce³⁺ and Ce⁴⁺ excitation in mixed garnets

Here we discuss luminescence and excitation spectra of LuGd₂Ga₃Al₂O₁₂:Ce, Mg under band-to-band excitation and their difference from those of LuGd₂Ga₃Al₂O₁₂:Ce. In Experimental we have shown with XANES and transmission spectroscopy that the Mg-codoped sample has no Ce³⁺.

In Fig. 3a the emission spectra under 7 eV excitation of LuGd₂Ga₃Al₂O₁₂:Ce 0.2% (black) and LuGd₂Ga₃Al₂O₁₂:Ce, Mg 0.2% (red) are presented. The spectra show the same 2.25 eV Ce³⁺ emission and the lines of Gd³⁺ at 4.50 and 3.96 eV. The spectra are normalized on Gd³⁺ emission and LuGd₂Ga₃Al₂O₁₂:Ce, Mg exhibit three times lower Ce emission intensity probably due to lack of energy transfer between Gd³⁺ and Ce⁴⁺ as opposed to Gd³⁺ and Ce³⁺ interaction.

Excitation spectra of Ce luminescence in the two samples show very different properties, Fig. 3b. For LuGd₂Ga₃Al₂O₁₂:Ce sample excitation spectrum has been explained above (Fig. 2c, black): one can see the band at 5.2 eV (direct excitation of Ce³⁺), Gd³⁺ lines at 4.85 and 5.02 eV (Gd³⁺ \rightarrow Ce³⁺ energy transfer) and contribution from excitons as a sharp

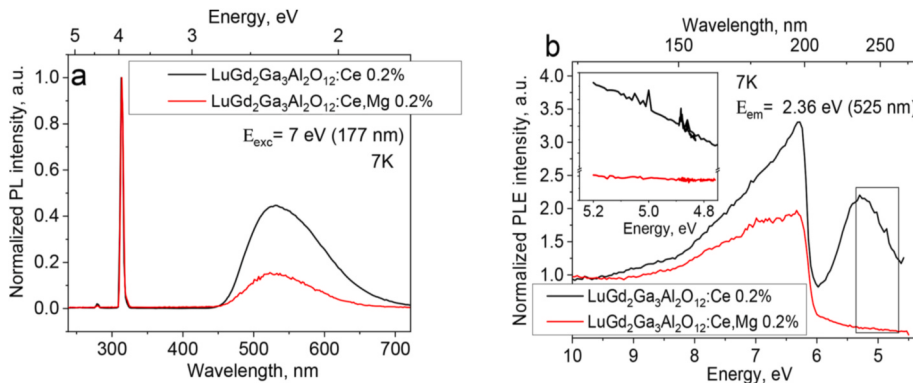


Fig. 3. a) Emission spectra at 7 K of LuGd₂Ga₃Al₂O₁₂ ceramics doped with 0.2% Ce (black) or 0.2% Ce and Mg (red), excited at 7 eV respectively. b) Excitation spectra at 7 K of Ce emission ($\lambda_{\text{em}} = 525 \text{ nm}$) for LuGd₂Ga₃Al₂O₁₂ ceramics doped with 0.2% Ce (black) or 0.2% Ce and Mg (red). The inset shows the presence of Gd f-f transitions in LuGd₂Ga₃Al₂O₁₂:Ce excitation spectrum and lack of those in LuGd₂Ga₃Al₂O₁₂:Ce, Mg excitation spectrum.

edge at 6.3 eV. The excitation spectrum of LuGd₂Ga₃Al₂O₁₂:Ce, Mg sample shows no Gd³⁺ f-f transitions (see inset) and can only be excited above 6.3 eV. The shape of the band-to-band excitation is rather flat, showing small contribution from direct exciton formation [39]. From the difference between excitation spectra of LuGd₂Ga₃Al₂O₁₂:Ce and LuGd₂Ga₃Al₂O₁₂:Ce, Mg we propose that Ce⁴⁺ interacts neither with Gd³⁺ nor with excitons in ways that lead to luminescence.

Now we compare X-ray excited luminescence (XRL) spectra of the two LuGd₂Ga₃Al₂O₁₂:Ce and LuGd₂Ga₃Al₂O₁₂:Ce, Mg samples and their intensity, see Fig. 4. As with PL emission spectra, the shape of XRL emission spectra of the samples is the same: 2.25 eV band of Ce³⁺ 5d-4f transitions. The XRL intensity for LuGd₂Ga₃Al₂O₁₂:Ce, Mg 0.2% is lower by a factor of three, consistent with findings of W. Chewpraditkul et al. [17] on light yield of GGAG:Ce, Mg crystals with 0.1% Mg co-doping. We connect the low XRL intensity (low light yield) of garnets overly co-doped with Mg²⁺ to hindered exciton channel of energy transfer towards Ce ions. Formation of (Ce–Mg)-centers [45] and O²⁻-Mg²⁺ centers [6,14,46] was shown to occur, likely leading to lower scintillation efficiency. Additionally, the re-charging of Ce⁴⁺ to its equilibrium state can be delayed by hole-trapping [6,7] lowering the efficiency of Ce⁴⁺ as recombination center.

Based on the experiments described above we have constructed bandgap diagrams on how the thermalized charge carriers recombine on Ce for two extreme cases of only Ce³⁺ or Ce⁴⁺ present in garnets, Fig. 5a

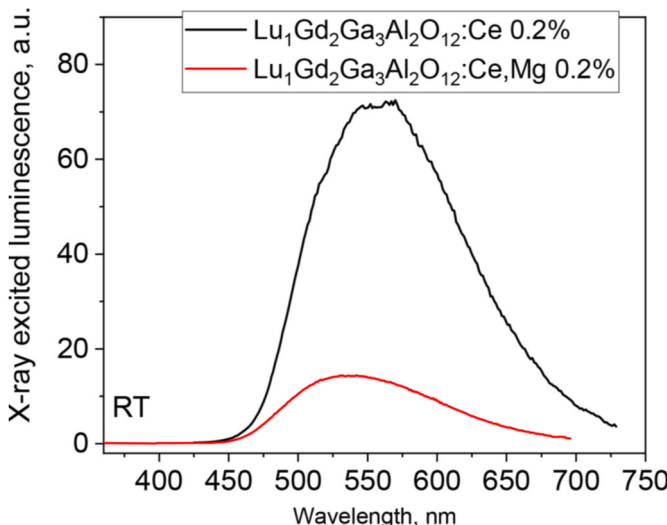


Fig. 4. X-ray excited luminescence spectra at 300K LuGd₂Ga₃Al₂O₁₂ ceramics doped with 0.2% Ce (black) or 0.2% Ce and Mg (red).

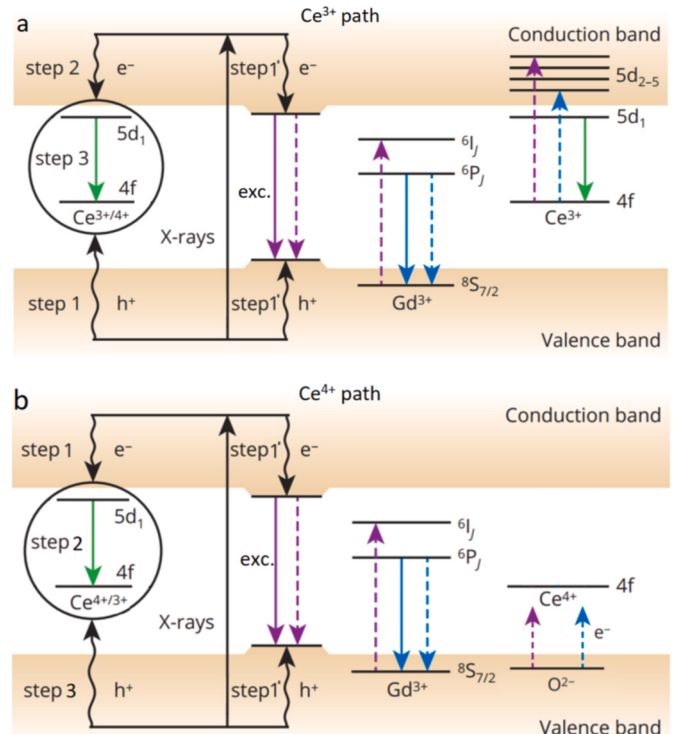


Fig. 5. Bandgap diagrams describing the mechanisms of e-h and exciton capture and transport towards a) Ce³⁺ and b) Ce⁴⁺. ‘X-rays’ stands for creation of electrons (e^-) and holes (h^+) in CB and VB, respectively. The steps (1)–(3) of sequent e-h recombination on Ce³⁺ and Ce⁴⁺ are constructed after [14]. Step (1') is an alternative path of exciton formation. Solid arrows indicate radiative transition: green – Ce³⁺ emission, purple – exciton emission, blue – Gd³⁺ 3.96 eV emission. Dashed arrows indicate resonant energy transfer or re-absorption. Note that in (b) excitonic path does not lead to Ce luminescence.

and b, respectively.

In the diagrams two recombination channels are described, the e-h (h-e) recombination on Ce, steps (1)–(3), and exciton formation, step 1', with sequent energy transfer to Gd³⁺ and Ce³⁺. After absorption of X-ray photon secondary electrons and holes thermalize to the bottom of CB and top of VB respectively. The e-h pair can be captured by Ce³⁺ or Ce⁴⁺ in sequent manner, leading to Ce³⁺ 5d-4f emission (green arrow). The e-h recombination on Ce⁴⁺ is usually distinguished from h-e recombination on Ce³⁺ experimentally with rise-time measurements [8] and transient spectroscopy [9]. In depth these mechanisms are described in Refs. [12,14,16]. Here we are more focused on alternative process of

exciton formation (step 1'), which can then emit (solid purple arrow) or become localized/transfer energy to Gd^{3+} or Ce^{3+} states (dashed purple arrow). The $\text{Gd}^{3+} \text{ }^8\text{S}_{7/2} \rightarrow \text{ }^6\text{I}_1$ and $\text{Ce}^{3+} \text{ }^2\text{F}_{5/2} \text{--} \text{ }^2\text{F}_{7/2} \rightarrow \text{ }^2\text{D}_{5/2}$ transitions overlap well with UV exciton emission [37]. Gd^{3+} can emit on its own as $\text{P}_{J=1} \text{--} \text{ }^8\text{S}_{7/2}$ at 3.96 eV (313 nm) as the last step for exciton path in LuGd₂Ga₃Al₂O₁₂:Ce, Mg sample (blue solid line, Fig. 5b), or Gd^{3+} can transfer energy to Ce^{3+} (Fig. 5a, for details see supporting information or [47]).

In LuGd₂Ga₃Al₂O₁₂:Ce case formation of excitons leads to Ce^{3+} luminescence, while Ce^{4+} in LuGd₂Ga₃Al₂O₁₂:Ce, Mg cannot interact positively with Gd^{3+} or excitons (dashed blue and solid arrow on CT in Fig. 5b). Additionally, CT absorption band of Ce^{4+} can re-absorb excitonic and Gd^{3+} emission further negatively impacting on the light yield of garnet materials.

Ce^{4+} in scintillators is used to rectify the short-comings of Ce^{3+} ability to compete with electron traps [7,12,48], but in case Ce^{4+} concentration becomes too high the number of transport pathways for delocalized charge to reach Ce is diminished.

2. Conclusions

Based on our finding we conclude the following. As Ce^{3+} 5d₃₋₅ band is located in UV range where excitons emit, Ce^{3+} can accept energy from excitons. When part of the lattice is substituted with Gd ions, Ce^{3+} luminescence is enhanced via exciton $\rightarrow \text{Gd}^{3+} \rightarrow \text{Ce}^{3+}$ energy transfer. The energy transfer from excitons is visible in excitation spectra for both Ce^{3+} and Gd^{3+} .

The excitation spectrum of LuGd₂Ga₃Al₂O₁₂:Ce, Mg (Ce^{4+}) luminescence shows no Gd^{3+} f-f transitions and no interactions with excitons. From the difference between excitation spectra of LuGd₂Ga₃Al₂O₁₂:Ce and LuGd₂Ga₃Al₂O₁₂:Ce, Mg we have found that Ce^{4+} interacts neither with Gd^{3+} nor with excitons in ways that lead to luminescence. The exciton $\rightarrow \text{Gd}^{3+} \rightarrow \text{Ce}^{3+}$ energy transfer path is interrupted, which is one of the reasons for lowered Ce^{3+} X-ray excited luminescence in LuGd₂Ga₃Al₂O₁₂:Ce, Mg ceramics.

The combined presence of Ce^{3+} and Ce^{4+} ions allows efficient competition with hole and electron traps, as well as practical channels for host excitations to reach activator ions. Finely-tuned concentration of both Ce^{3+} and Ce^{4+} in the material has led to the highest light yield garnet materials.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

The authors acknowledge the expert help of the staff of MAX IV Laboratory. The research leading to this result has been supported by the project CALIPSOplus under the Grant Agreement 730872 from the EU Framework Programme for Research and Innovation HORIZON 2020. I. V. acknowledges the support of Russian Foundation for Basic Research # 20-52-S52001.

Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jlumin.2021.118150>.

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