Band-gap variation and a self-redox effect induced by compositional deviation in ZnxGa2O3+x:Cr3+ persistent phosphors
Band-gap variation and self-redox effect induced by compositional deviation in Zn$_x$Ga$_2$O$_{3+x}$:Cr$^{3+}$ persistent phosphors

Yixi Zhuang,*a Jumpei Ueda,b Setsuhisa Tanabe,c and Pieter Dorenbosd

We made a systematic investigation on the optical properties of non-doped and Cr-doped Zn$_x$Ga$_2$O$_{3+x}$ (0.98 $\leq$ x $\leq$ 1.02) spinel crystals. Absorption, photoluminescence excitation, and persistent luminescence excitation spectra indicated that the bottom of conduction band (CB) was affected (i) by compositional deviation from stoichiometry or (ii) by atmosphere control in a similar way. By constructing an energy level diagram, a broad distribution of trap depth in a Cr$^{3+}$-doped Zn$_x$Ga$_2$O$_{3+x}$ sample with composition of Zn excess or in one prepared in O$_2$ atmosphere was attributed to lowering of the CB bottom, which resulted in faster decay of persistent luminescence in these two samples. Defect chemical reactions revealed that the anti-sites species of Zn$'$Ga may be the reason for lowering of the CB bottom. A self-redox effect of Cr ions induced by the compositional deviation was found and explained by the defect reactions. The investigation indicated that the composition with slight Zn deficiency is important to obtain a narrow distribution of trap depths and longer persistent luminescence, however the self-reduction effect of Cr ions in the composition of Zn deficiency also need to be considered.

1 Introduction

Persistent phosphors, which continue light emission after removal of the excitation source, have long been of scientific interests due to their important applications as luminous paints, identification markers, or optical memories.1,5 Recently, another attractive application in in vivo medical imaging system by using nano-sized persistent particles as optical probe was proposed by a French group, which focused more attention on the persistent phosphors.6,7 One of the key requirements for in vivo imaging application is that the luminescence wavelength of persistent particles should be inside the biological window from 650 to 1300 nm where the optical signal shows rather high transmittance through the biological tissues.8 This spectral requirement rules out most Eu$^{2+}$-activated phosphors showing excellent persistent performance in the blue or green regions.9,10 Until now, only few Eu$^{2+}$-activated compounds were known to exhibit persistent luminescence beyond 650 nm. Most of them are hygroscopic sulfides and unsuitable for biological application.11,12 Alternatively, several Cr$^{3+}$-activated gallate phosphors, especially ZnGa$_2$O$_4$-based spinel crystals, were reported showing long-lasting persistent luminescence from 680 to 1000 nm, which were considered as promising optical materials for in vivo imaging application.13-26

Bessière et al. firstly reported red persistent luminescence at around 695 nm in Cr$^{3+}$-doped ZnGa$_2$O$_4$ crystals in 2011.16 Pan and Alix et al. greatly improved the red persistent luminescence by partially substituting Ge to form Zn$_{1+x}$Ga$_{2-2x}$Ge$_x$O$_4$ solid solution. The persistent luminescence was detectable by night vision equipment up to 360 h after the excitation source was removed.17,18 We also found that by codoping Bi$_2$O$_3$ into Zn$_{0.98}$Ga$_2$O$_{3.98}$ crystals, the intensity of red persistent luminescence can be improved by 10 times. The absolute intensity (radiance) of persistent luminescence in Zn$_{0.98}$Ga$_2$O$_{3.98}$:Cr,Bi phosphors was comparable to that in SrAl$_2$O$_4$:Eu,Dy phosphors.19 Recently, Abdukayum et al. succeeded to demonstrate in vivo imaging experiments by using Zn$_{1+x}$Ga$_{2-2x}$Ge$_x$O$_4$:Cr,Pr nano-particles. The feasibility, including Cr$^{3+}$ persistent luminescence duration, particle biocompatibility, water solubility etc. was proved in this study.20 Although notable progress on intensity improvement or nanoparticulation has been made, the Cr$^{3+}$-doped ZnGa$_2$O$_4$ crystals need further investigation in order to understand the energy levels governing the red persistent luminescence. As mentioned by Bessière et al. in their milestone paper, a sample with Zn deficiency in the composition showed more intense persistent luminescence than other samples.16 The composition of Zn deficiency was adopted by many researchers. However, the reason for the composition dependency has not been given.
A systematic study on persistent luminescence induced by compositional deviation may give much information on structure variation, and offer theoretical guidance for the design of better persistent phosphors.

In this paper, we introduce methods to discuss the persistent luminescence mechanism by building an energy level diagram (Sec. 2.1) and to discuss structural variation by using a defect formation equation (Sec. 2.2), which are applied in the following sections. In Sec. 3, synthesis conditions (nominal composition and applied atmosphere) of prepared samples and important details about optical measurements are described. The results are summarized and discussed in Sec. 4. It starts from a discussion on band-gap variation in non-doped Zn$_x$Ga$_2$O$_{3+x}$ samples prepared in different conditions (either composition or atmosphere) (Sec. 4.1), and moves to optical characterizations in various Cr$^{3+}$-doped Zn$_x$Ga$_2$O$_{3+x}$ samples by discussing the change of valence state in Cr ions (Sec. 4.2) and the variation of energy level diagram as well as Cr$^{3+}$ persistent luminescence performance (Sec. 4.3). Some conclusions are made in Sec. 5.

2 Methodology

2.1 Building an energy level diagram to discuss the persistent luminescence mechanism

It is indispensable to understand the mechanism of persistent luminescence for successful material synthesis. However, it was a bigger challenge to give a clear explanation for the observed persistent phenomenon, rather than to discovered an excellent material (sometimes accidently) in the past.\(^9\)\(^,\)\(^27\) Several models, generally by using a simple energy level diagram to describe the charge carrier transfer process between active centers and trap centers, were proposed by different researchers.\(^4\) Two major controversial points in these models are (i) the species of charge carrier from the luminescent center (whether a negative electron or a positive hole) captured by the trap center and (ii) the path of the trapping process (whether through the conduction band or through the valence band or by a quantum tunnelling effect in the band-gap). The persistent luminescence mechanism can be correctly determined only when the relative energy locations of (i) the valence band (VB) and the conduction band (CB) of the host, (ii) the ground state (GS) and excited states (ES) of the luminescent center as well as, (iii) the trap energy levels are clear. It is significant to determine the relative locations of these energy bands/levels in the research of persistent phosphors, by using various optical characterizations.

### Fig. 1 Schematic energy level diagram showing typical electron trapping processes.

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<th>Energy (eV)</th>
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<tr>
<td>GS</td>
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<tr>
<td>Lumin. Center</td>
</tr>
<tr>
<td>ES$_1$</td>
</tr>
<tr>
<td>ES$_2$</td>
</tr>
<tr>
<td>ES$_3$</td>
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<tr>
<td>CB</td>
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<tr>
<td>Trap</td>
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In recent publications, most experimental results confirm that an electron from a luminescent center captured by a trap center through the CB is the most efficient path for charge carrier trapping process, although the contribution of quantum tunnelling cannot be completely excluded. Possible trapping processes are shown in Fig. 1. The electron can be excited to the CB in three ways: (i) from GS to ES above the bottom of CB; (ii) directly from GS to CB; (iii) from GS to ES below the bottom of CB. In the third case, the electron can “jump” to CB with thermal energy, which was named thermally assisted photoionization process. The specific trapping process is determined by the relative locations of the energy bands/levels.

Dorenbos has developed an empirical model to determine the GS and ES (f and d orbitals) of R$^{2+}$ and R$^{3+}$ (rare earth ions in +2 and +3 valence states) relative to CB and VB, which was a very useful tool to discuss the persistent properties in many rare-earth-activated phosphors.\(^28\)\(^-\)\(^31\) We consider that the same research approach can be applied to transition-metal-activated materials. Firstly, a host referred binding energy (HRBE) scheme is adopted, where the energy of the VB top is defined as zero. The HRBE of the CB bottom can be determined by the band-gap energy from absorption spectra of a non-doped sample. In the second step, we need to know the relative locations of energy levels of luminescent center (GS and ES) relative to the energy bands of the host (CB and VB), Photoluminescence excitation (PLE) spectra, persistent luminescence excitation spectra (PersLE), or photoluminescence excitation (PLE) spectra measurements are useful to evaluate the energy of electronic transition from the GS to the bottom of CB (arrow ii in Fig. 1). Also, the energy values for 3d-3d intra-transitions from GS to various ES can be read from absorption spectra or PLE spectra in a transition-metal-doped sample. Finally, the depth of trap levels ($\Delta \varepsilon$, thermal activation energy) can be determined in the measurement of thermoluminescence (TL) glow curves.
2.2 Using Kröger-Vink notation to discuss structural variation

In many persistent phosphors, synthesis condition (compositional deviation from stoichiometry or atmosphere control) show either positive effect or negative effect on the persistent luminescence performance. It is helpful if we know what kind of structural change occurs, i.e., which species of defects are created in a real synthesis process. Since direct detection of the defects is generally difficult or unavailable, a theoretical method by writing defect chemical reactions in Kröger-Vink notation can be used. The key defects responsible for the optical properties could be inferred by connecting the applied synthesis condition with the obtained optical properties, although experimental proof is usually required to verify this. For example, the great improvement of persistent luminescence in SrAl₂O₄:Eu²⁺ phosphor by Dy³⁺ codoping was as attributed to the formation of DySr⁺ defect as following:

\[ \text{Dy}_2\text{O}_3 + 2\text{Al}_2\text{O}_3 \rightarrow 2\text{Dy}_2\text{Sr}^+ + 4\text{Al}^3+ + 8\text{O}_2^- + 2e^- + \frac{1}{2} \text{O}_2 \uparrow \] (1)

3 Experimental Procedures

3.1 Sample preparation

Poly-crystalline ceramics were prepared by using a solid-state reaction method. The chemical reagents, ZnO (4N), Ga₂O₃ (4N), and Cr₂O₃ (3N), were used as starting materials. The starting materials were ground in an alumina mortar to form homogeneous and fine powder mixtures. Pellets (0.8 g and φ15 mm) were shaped from the powder mixtures with a uniaxial loading of 50 MPa in a stainless steel mold. The compacts were carefully moved to an electric furnace and fired at 1300 °C for 5 h.

Nominal composition (the molar ratio of Zn to Ga) and applied atmosphere (O₂ partial pressure) were used as two variables to investigate their effects on optical properties. Various synthesis conditions for non-doped ZnₓGa₂O₄₋ₓ samples are shown in Table 1. Firstly the atmosphere was fixed as Air and the composition was varied from 2% Zn deficiency (Zn-d), stoichiometric (Zn-s), to 2% Zn excess (Zn-e) to obtain samples non-d-Air, non-s-Air, and non-e-Air, respectively. Then the composition was kept as stoichiometric and the gas flow was changed from pure N₂ (P₂O₂ = 0%), Air (P₂O₂ = 21%), to pure O₂ (P₂O₂ = 100%) to obtain samples non-s-N₂, non-s-Air, and non-s-O₂. In Table 2, Cr-doped ZnₓGa₂O₃₋ₓ samples were prepared in the same synthesis conditions except for 0.5 mol% substitution of Cr₂O₃ for Ga₂O₃.

<table>
<thead>
<tr>
<th>Comp. (Zn/Ga ratio)</th>
<th>Atmosphere (P₂O₂)</th>
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<tbody>
<tr>
<td>Zn₀.₀₉Ga₂O₄₋ₓ: Zn-d</td>
<td>N₂, non-d-Air</td>
</tr>
<tr>
<td>Zn₁.₀₆Ga₂O₄₋ₓ: Zn-s</td>
<td>non-s-O₂, non-s-Air, non-s-N₂</td>
</tr>
<tr>
<td>Zn₁.₀₆Ga₂O₄₋ₓ: Zn-e</td>
<td>non-e-Air</td>
</tr>
</tbody>
</table>

| ZnₓGa₂O₃₋ₓ poly-crystalline ceramics. The samples were notated as (dopant)-(composition)-(atmosphere). d, s, and e were referred to 2% Zn deficiency, stoichiometric, and 2% Zn excess in the composition. N₂, Air, and O₂ corresponded to gas flow of pure N₂, Air, and pure O₂.

Table 1 Nominal compositions and applied atmospheres of non-doped ZnₓGa₂O₃₋ₓ poly-crystalline ceramics. All the synthesis conditions were the same with the non-doped samples except for using 0.5 mol% Cr₂O₃ as dopant.

Table 3 Optical Measurements

Diffuse reflectance spectra of the samples were measured by using a scanning-type spectrophotometer (Shimadzu, UV3600) with a BaSO₄-based integrating sphere. The scanning wavelength region was from 850 to 200 nm. Photoluminescence (PL), photoluminescence excitation (PLE), and persistent luminescence excitation (PersLE) spectra were measured in a fluorescence spectrophotometer (Shimadzu, RF5000). Persistent luminescence excitation spectra were obtained by plotting Cr³⁺ persistent luminescence intensity in 60 s after excitation for 60 s at each excitation wavelength. Excitation wavelength was changed from 500 to 200 nm with a step of -5 nm. Before moving to the next excitation wavelength, a waiting time of 10 min was set to reduce the effect of the last excitation. Thermoluminescence (TL) glow curves of Cr³⁺ emission (around 695 nm) were recorded using a silicon photodiode (Electro-Optical System Inc., S-025-H). A band-pass filter (350-750 nm) and a long-pass filter (> 580 nm) were fixed before the silicon photodiode to cut off the noise. The samples were first cooled to 100 K (-173 °C) and exposed to a 300 W xenon lamp (Asahi Spectra, MAX 302) in an ultra-violet (UV) mirror module (250 to 380 nm) for 5 min. After another 5 min of waiting time, the samples were heated to 600 K (327 °C) in a heating rate of 10 K/°C. The persistent luminescence decay curves at room temperature (RT, 20 °C) were recorded by using the same silicon photodiode after the samples were excited by the same xenon lamp in the UV module for 5 min. Then, the afterglow intensity was converted to an absolute intensity (radiance, in unit of mW·sr⁻¹·m⁻²) by using a CCD spectrometer (B&W Tek, Glacier X) calibrated by Konica-Minolta company.
4 Results and Discussion

4.1 Band-gap absorption and self-activated emission in non-doped ZnxGa2O3+x

In order to evaluate the optical band-gap, the diffuse reflectance \( (R_\infty) \) of the non-doped ZnxGa2O3+x pellets were converted to a Kubelka-Munk function \( F(R_\infty) \) according to:\(^3\)

\[
F(R_\infty) = S \times \frac{(1 - R_\infty)^2}{2 \times R_\infty}.
\]

As an approximation, the diffusion coefficient \( S \) was treated independent on wavelength. Then a \( (\hbar v^* F(R_\infty))^{-2} - \hbar v \) plot was made according to an expression proposed by Tauc:\(^3\)

\[
(\hbar v \times F(R_\infty))^{-2} = A \times (\hbar v - E_g).
\]

The values of the band-gap energy can be read from the intercepts of fitted straight lines.

The fitting results of the three samples with different compositions (molar ratio of Zn to Ga) are given in Fig. 2(a). The sample with Zn deficiency (blue) showed a band-gap of 4.90 eV. The band-gap energy decreased to 4.88 and 4.58 eV when the molar ratio of Zn to Ga was varied to stoichiometric (black) and Zn excess (red), respectively. The effect of atmosphere (\( P_{O_2} \)) on the band-gap energy is shown in Fig. 2(b). The band-gap decreased from 4.90, 4.88 to 4.74 eV when the applied atmosphere for the solid state reaction was varied from \( N_2 \) (blue), Air (black), to \( O_2 \) (red).

In Fig. 3(a) and (b), the PL and PLE spectra of the non-doped ZnxGa2O3+x are given. The PLE spectra in the non-d-Air (right and top, blue) and non-s-N2 samples (right and bottom, blue) showed excitation bands maximizing at approximately 5.00 eV (248 nm) due to optical band-gap transitions. The excitation bands red-shifted to 4.96 eV (250 nm) in the non-s-Air sample (right, black), and further moved to 4.43 eV (280nm) in the non-e-Air sample (right and top, red) and 4.59 eV (270 nm) in the non-s-O2 sample (bottom and right, red). The variation of PLE bands is consistent with the results in the absorption spectra (Fig. 2). The PL spectra (left part of Fig. 3) of the non-doped ZnxGa2O3+x samples showed self-activated emissions in the UV and visible regions. The emission bands shifted towards lower energy in the sample with Zn excess (left and top, red) or in the one prepared in \( O_2 \) atmosphere (left and bottom, red) compared with other samples.

Schematic energy level diagrams of the non-doped ZnxGa2O3+x samples are depicted in Fig. 4. The HRBE concept was used where the top of the VB was defined as zero of energy. According to the results of absorption, PLE and PL spectra, the energies of band-gap absorption and self-activated emission in the non-d-Air and non-s-N2 samples were approximately 4.9 and 3.5 eV, respectively; the energies decreased and covered broader regions in 4.4–4.9 eV for band-gap absorption and 3.0–3.5 eV for self-activated emission. The decrease was attributed to lowering of the lowest component (bottom) of CB as shown in Fig. 4.
It is known that the ZnGa$_2$O$_4$ crystalizes in an ideal normal spinel under ambient pressure, i.e. Zn and Ga ions occupy tetrahedral and octahedral sites, respectively. However, real ZnGa$_2$O$_4$ crystals (by high-temperature sintering) should contain some degree of anti-site coordination, i.e., Zn$^{2+}$ in octahedral sites (Zn$^{\prime}$$_{Ga}$) and Ga$^{3+}$ in tetrahedral site (Ga$^{\prime}$$_Zn$). These defects, as compared to other defects, require much lower formation energy (0.9 eV) according to Pandey et al.\textsuperscript{48}

$$\text{Zn}^{\prime}_Zn + \text{Ga}^{\prime}_Ga \rightarrow \text{Zn'}_{Ga} + \text{Ga}^{\prime}_Zn$$ (4)

The concentration of the two anti-site defects also should be affected by compositional deviation from stoichiometry, or by different applied atmosphere. In a Zn$_x$Ga$_2$O$_{3+x}$ spinel crystal with Zn excess or in higher oxygen partial pressure (e.g. O$_2$), a defect reaction in Eq. (5) can be promoted, where the defect species of Zn$^{\prime}$$_{Ga}$ is populated. On the other hand, in Zn$_x$Ga$_2$O$_{3+x}$ crystals with composition of Zn deficiency or in lower oxygen partial pressure (e.g. N$_2$), a defect reaction in Eq. (6) is promoted and the number of defect species of Ga$^{\prime}$$_Zn$ is increased.

$$3\text{ZnO} + \frac{1}{2} \text{O}_2 \rightarrow 2\text{Zn}^{\prime}_Zn + \text{Zn}_Zn + 4\text{O}_2^0 + 2\text{h}^+$$ (5)

$$3\text{Ga}_2\text{O}_3 \rightarrow 2\text{Ga}^{\prime}_Zn + 4\text{Ga}^{\prime}_Ga + 8\text{O}_2^0 + \frac{1}{2} \text{O}_2 ↑ + 2\text{e}^-$$ (6)

We notice that the decrease of the band-gap was observed in both of the non-e-Air and non-s-O$_2$ samples. Therefore, we consider that the formation of Zn$^{\prime}$$_{Ga}$ may be the reason of the CB lowering. Indeed, we cannot totally exclude the possibility of other defect species (e.g. Zn$^{\prime}$$_Zn$ or V$^{\prime}$$_{Ga}$), although the connections between the observed properties and these defects have not been found.

### 4.2 Redox effects of Cr ions induced by compositional deviation from stoichiometry

The Cr-doped Zn$_x$Ga$_2$O$_{3+x}$ crystals show broad absorption bands peaked at around 560, 420, and 280 nm due to $^4\text{A}_2 \rightarrow ^4\text{T}_2$, $^4\text{A}_2 \rightarrow ^4\text{T}_1$, and $^4\text{A}_2 \rightarrow ^4\text{T}_1(\text{P})$ electronic transitions of Cr$^{3+}$ ions. The Kubelka-Munk function, which is proportional to the absorption coefficient, can be used to quantify the concentration of Cr$^{3+}$ ions. As shown in Fig. 5 (a), when the composition is varied from Zn excess to Zn deficiency, the absorption coefficient of Cr$^{3+}$ decreases, indicating a reduction of Cr$^{3+}$ concentrations in the sample with Zn deficiency. In Fig. 5 (b), the absorption coefficient of Cr$^{3+}$ also decreases from the sample prepared in O$_2$ (Cr-s-O$_2$) to that prepared in N$_2$ (Cr-s-N$_2$).

Cr ions are expected to dominate in the Cr$^{3+}$ valence state in the ZnGa$_2$O$_4$ spinel crystals as luminescent centers. However, abnormal state can be formed, as Peng’s model suggests, in a case when there is a driving force for the stabilization of the abnormal state and the total free energy is reduced. As discussed in Sec. 4.1, in the sample with composition of Zn deficiency or in the one prepared in N$_2$, the defect reaction of Eq. (6) is promoted. Notice that additional negative electrons (e$^-$) are created in this reaction. These electrons can combine with Cr$^{3+}$ ions incorporated in Ga$^{3+}$ sites (Cr$^{x}_Ga$) and reduce them into Cr$^{2+}$ (Cr$^{x}_Ga$), as shown in Eq. (7). The decrease of the Cr$^{3+}$ absorption coefficient in the Cr-d-Air sample may be due to such “self-reduction effect” in the sample with Zn deficiency. For the sample with Zn excess or the one prepared in O$_2$, the defect reaction of Eq. (5) is promoted. Because of the resultant positive holes (h$^+$), an oxidizing reaction in Eq. (8) can keep more Cr ions in the Cr$^{3+}$ state.

$$\text{Cr}^{x}_Ga + e^- \rightarrow \text{Cr}^{x}_Ga$$ (7)

$$\text{Cr}^{x}_Ga + h^+ \rightarrow \text{Cr}^{x}_Ga$$ (8)

In order to support the argument of the self-redox effect, electron spin resonance (ESR) measurements were performed in the Cr-doped Zn$_x$Ga$_2$O$_{3+x}$ samples. The ESR spectra at X band frequency (9.10 GHz) showed a resonance absorption at approximate 0.32 mT at room temperature. This absorption was attributed to nearest-neighbour Cr$^{3+}$ pairs in the spinel Zn$_x$Ga$_2$O$_{3+x}$ crystals.\textsuperscript{50-52} The ESR spectra in the Cr-e-Air and Cr-s-O$_2$ samples showed larger intensity of the resonance absorption, which confirmed higher concentration of Cr$^{3+}$ ions in these two samples. It is note that the Cr$^{2+}$ concentration should increase in the Cr-d-Air and Cr-s-N$_2$ samples when the Cr$^{3+}$ concentration decreases due to the reduction effect. Unfortunately, quantitative evaluation of Cr$^{2+}$ concentration in the Cr-doped Zn$_x$Ga$_2$O$_{3+x}$ samples was unavailable in the absorption, PL, PL spectra and ESR measurements.

![Fig. 5 Kubelka-Munk plot of the Cr-doped Zn$_x$Ga$_2$O$_{3+x}$ crystals. Electronic transitions from ground state ($^4\text{A}_2$) to various excited states ($^4\text{T}_2$, $^4\text{T}_1$, and $^4\text{T}_1(\text{P})$) of Cr$^{3+}$ ion are notated.](image)
4.3 Variation of PL and persistent luminescence in the Cr-doped Zn$_x$Ga$_2$O$_3$ crystals

Under excitation by UV or visible light, the Cr-doped Zn$_x$Ga$_2$O$_3$ samples showed several emission peaks at around 700 nm due to $^2$E $\rightarrow$ $^4$A$_2$ transitions of Cr$^{3+}$. One single peak at 695 nm (N$_2$ line) was attributed to Cr$^{3+}$ in distorted octahedral sites; other peaks at 688 (R line), 679, 708 nm etc. (phonon side bands, PSB) were originated from Cr$^{3+}$ in ideal octahedral sites.

The PLE spectra of the Cr-doped Zn$_x$Ga$_2$O$_3$$_{1-x}$ samples are shown in Fig. 6. The bands at short wavelength (designated as band 0) were attributed to the band-gap absorption. This band shifts from 250 nm (4.96 eV) in the Cr-d-Air and Cr-s-N$_2$ samples to 280 nm (4.42 eV) in the Cr-e-Air and Cr-s-O$_2$ samples, which is consistent with those in the non-doped Zn$_x$Ga$_2$O$_3$ crystals (Fig. 3). Another band at around 340 nm (3.6 eV) (band 2) was attributed to an electronic transition from GS of Cr$^{3+}$, i.e. $^4$A$_2$ level to CB of the host. This band shifts to lower energy in the Cr-e-Air and Cr-s-O$_2$. Other bands at around 280 nm (4.43 eV) (band 1), 420 nm (2.95 eV) (band 3), and 560 nm (2.11 eV) (band 4) due to 3d-3d transitions from $^4$A$_2$ to $^4$T$_1(^4P)$, $^4$T$_1$, and $^4$T$_2$ of Cr$^{3+}$ were also detected in the PLE spectra. The energy of these 3d-3d transitions showing very weak dependence on the preparing condition (composition or atmosphere), are ignorable compared with the variation in $^4$A$_2$ $\rightarrow$ CB and VB $\rightarrow$ CB transition energies (band 0 and band 2).

The PersLE spectra of the Cr-doped Zn$_x$Ga$_2$O$_3$$_{1-x}$ samples in Fig. 7 showed broad bands from 400 nm (3.05 eV) to 270 nm (4.59 eV). These bands have two components: $^4$A$_2$ $\rightarrow$ CB (band 2) and $^4$A$_2$ $\rightarrow$ $^4$T$_1(^4P)$ transition (band 1). Different from the PLE spectra, the $^4$A$_2$ $\rightarrow$ CB transition (band 2) in PersLE spectra shows much higher intensity than the other bands. Also, a red-shift of the $^4$A$_2$ $\rightarrow$ CB transitions (band 2) in the Cr-e-Air and Cr-s-O$_2$ samples was observed as compared with the same transitions in the Cr-d-Air and Cr-s-N$_2$ samples. The bands 0, 3, and 4 present in the PLE spectra could be hardly detected in Fig. 7.

Based on the obtained results, the energy level diagram in Fig. 8 including CB, VB, and various energy levels of Cr$^{3+}$ is constructed. The $^4$A$_2$ $\rightarrow$ CB transition (band 2) provides the energy location of the $^4$A$_2$ level (GS of Cr$^{3+}$) below the bottom of the CB. The ES of Cr$^{3+}$ can then be located using the PLE spectra. For example, in the Cr-d-Air or Cr-s-N$_2$ samples (Fig. 8, left part), the energy difference between $^4$A$_2$ and the CB bottom is approximately 3.7 eV. The $^4$T$_1(^4P)$ level is 4.4 eV above the $^4$A$_2$ level and located inside the CB. On the other hand, the $^4$T$_1$ and $^4$T$_2$ levels are lower than CB. Consequently, electron transfer through the CB to the traps can be realized by excitation of $^4$A$_2$ $\rightarrow$ $^4$T$_1(^4P)$ (band 1) and $^4$A$_2$ $\rightarrow$ CB transitions (band 2), but will fail by excitation of $^4$A$_2$ $\rightarrow$ $^4$T$_1$ (band 3) and $^4$A$_2$ $\rightarrow$ $^4$T$_2$ transitions (band 4). Also, direct excitation into CB ($^4$A$_2$ $\rightarrow$ CB $\rightarrow$ traps) was proved as a more effective route for electron trapping than $^4$A$_2$ $\rightarrow$ $^4$T$_1(^4P)$ $\rightarrow$ CB $\rightarrow$ traps. A fast relaxation of $^4$T$_1(^4P)$ to $^4$T$_1$ may decrease the electron trapping efficiency in the latter case.

Fig. 8 Energy level diagrams of Cr-doped Zn$_x$Ga$_2$O$_3$$_{1-x}$ crystals showing location of Cr$^{3+}$ energy levels relative to CB and VB as well as the energy level of traps.
As shown in Fig. 4, when the molar ratio of Zn to Ga was varied from Zn deficiency to Zn excess, or when the applied atmosphere was changed from N₂ to O₂, the CB bottom will be lowered by 0.3-0.4 eV. Based on the PersLE spectra, the energy difference from ⁴A₂ to the CB bottom decreases by approximately 0.3 eV in the Cr-e-Air or Cr-s-O₂ samples. The energy of the ⁴A₂ level relative to VB top remains unchanged in different samples (Fig. 8).

TL glow curves of the Cr-doped ZₙGa₂O₃₋ₓ samples are shown in Fig. 9. Again, the glow curve in the Cr-d-Air sample is similar to that in the Cr-s-N₂ sample. The glow curves show a band peaked at approximately 340 K with narrow full width at half maximum (FWHM = 70 K). The glow curves broaden in the Cr-e-Air and Cr-s-O₂ samples (FWHM = 180 K) and slightly shifted to lower temperature.

The peak temperature and FWHM of the TL glow curves reflect trap depth (thermal activation energy, \( \Delta \epsilon \)) and the trap depth distribution. As shown in Fig. 8, the change in \( \Delta \epsilon \) (decrease of the average value and broadening of the distribution) can be due to lowering of the CB bottom.

In Fig. 10, we compare the persistent luminescence decay curves of the Cr³⁺-doped ZₙGa₂O₃₋ₓ crystals after excitation by UV light (250-380 nm). The Cr-e-Air and Cr-s-O₂ samples show higher intensity of persistent luminescence in the initial stage (the first 30 s) and decay in a faster rate than the Cr-d-Air and Cr-s-N₂ samples. The persistent luminescence (in radiance) of the Cr-d-Air and Cr-s-N₂ samples 3600 s after removal of the UV light was several times higher than the Cr-e-Air and Cr-s-O₂ samples.

As discussed in Sec. 4.2, the composition and atmosphere showed a redox effect on the valence state of Cr ions. More Cr ions were stabilized in higher valence state, Cr³⁺ state in the Cr-e-Air and Cr-s-O₂ samples. These two samples showed higher intensity of PL and persistent luminescence in the initial stage. However, the detrapping rate in the Cr-e-Air and Cr-s-O₂ is too fast, which may be due to lower and broad distribution of \( \Delta \epsilon \) as shown in Fig. 8.

From the viewpoint of persistent phosphors used at RT, a narrow distribution of \( \Delta \epsilon \) at approximately 0.6 eV (corresponding narrow glow peaks centred at 300-350 K) is expected. Slight Zn deficiency in the composition should be adopted to keep a narrow distribution of \( \Delta \epsilon \). However, we have to consider the self-reduction effect and decrease of Cr³⁺ concentration in the composition of Zn deficiency. In Ref. 19, it was shown that Bi₂O₃ is a good stabilizer for Cr³⁺ state in a Cr-Bi-codoped Zn₀.₉₈Ga₂O₃.₉₈ sample. The co-doped sample showed greatly enhanced persistent luminescence because of a narrow glow peak at 330 K as well as a much higher concentration of stabilized Cr³⁺ ions.

5 Conclusion

In the process of investigating optical properties in non-doped ZₙGa₂O₃₋ₓ (0.98 ≤ x ≤ 1.02) crystals, decrease of band-gap energy due to lowering of CB bottom was observed in the sample with 2 % Zn excess in composition and in the sample prepared in O₂ atmosphere. The similar lowering of CB bottom in these two samples may be due to significant formation of Zn₇Ga anti-sites considering the most probable defect chemical reactions in composition of Zn excess or high partial oxygen pressure. The lowering of CB bottom resulted in decrease and broader distribution of trap depth. Slight Zn deficiency in the composition should be adopted to keep narrow distribution and trap depth for longer persistent luminescence decay. However, the sample with composition of Zn deficiency showed a self-
reduction effect on Cr ions into lower valence states. Effective measure should be taken to stabilize Cr ions in Cr³⁺ state.

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Notes and references

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