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Microstructure, doping and optical properties of $\text{Co}^{2+}:\text{ZnAl}_2\text{O}_4$ transparent ceramics for saturable absorbers: Effect of the ZnF_2 sintering additive

Alexander Belyaev^{a,*}, Liza Basyrova^b, Vadim Sysoev^c, Maxim Lelet^d, Stanislav Balabanov^a, Vladimir Kalganov^e, Vladimir Mikhailovski^e, Mikhail Baranov^b, Evgeniia Stepanidenko^b, Vladimir Vitkin^b, Olga Dymshits^f, and Pavel Loiko^g

^a*G.G. Devyatykh Institute of Chemistry of High-Purity Substances, RAS, 49 Tropinin St., 603951 Nizhny Novgorod, Russia*

^b*ITMO University, Kronverkskiy pr., 49, 197101 Saint-Petersburg, Russia*

^c*Lobachevsky State University of Nizhny Novgorod, 23 Gagarin Ave., 603950 Nizhny Novgorod, Russia*

^d*Research Institute for Chemistry, Lobachevsky State University of Nizhny Novgorod, 23 Gagarin Ave., 603950 Nizhny Novgorod, Russia*

^e*St. Petersburg State University, 7-9 Universitetskaya Emb., 199034 St. Petersburg, Russia*

^f*Vavilov State Optical Institute, 36 Babushkina St., 192171 St. Petersburg, Russia*

^g*Centre de recherche sur les Ions, les Matériaux et la Photonique (CIMAP), UMR 6252 CEA-CNRS-ENSICAEN, Université de Caen Normandie, 6 Boulevard du Maréchal Juin, 14050 Caen Cedex 4, France*

*Corresponding author, e-mail: belyaev@ihps.nnov.ru

ABSTRACT.

Transparent cobalt-doped zinc aluminate spinel (gahnite), $\text{Co}^{2+}:\text{ZnAl}_2\text{O}_4$, ceramics were fabricated by hot pressing of $(\text{Zn},\text{Co})\text{Al}_2\text{O}_4 / \text{ZnF}_2$ nanopowders at 1520 °C for 4 hours. A novel approach was suggested for the preparation of $(\text{Zn},\text{Co})\text{Al}_2\text{O}_4$ precursor: the $(\text{Zn},\text{Co})\text{Al}_2\text{O}_4$ powders were synthesized by mixing an alcohol solution of aluminium isopropoxide with a joint aqueous solution of zinc *formate* (the source of zinc), cobalt nitrate and zinc fluoride (the sintering additive) followed by drying and calcination in air. The study of the thermal behavior of precursors revealed an optimum calcination temperature (700 °C) preventing the loss of the sintering additive (ZnF_2). The effect of the ZnF_2 content (3-10 wt%) on the microstructure, the cobalt doping concentration, absorption and luminescence of ceramics was systematically studied. The ceramics exhibit a close-packed microstructure with a mean grain size of 50-70 μm and high in-line transmission (about 84% at $\sim 2 \mu\text{m}$). The actual concentration of Co^{2+} ions in tetrahedral (T_d) sites of gahnite is lower than the Co^{2+} doping level and it decreases with the ZnF_2 content. We propose a possible mechanism of this variation including both the losses of Co^{2+} via evaporation of the intermediate CoF_2 phase and a partial location of Co^{2+} ions in octahedral (O_h) sites of gahnite due to its partly inverse spinel structure. The developed ceramics are promising for saturable absorbers of erbium lasers.

Keywords: Transparent ceramics, Hot pressing synthesis, Sintering additive, Gahnite, Cobalt ions, Microstructure, Saturable absorbers of lasers.

1. Introduction

Spinel crystals are complex oxides with a cubic symmetry (sp. gr. $Fd\bar{3}m$) having a general chemical formula of AB_2O_4 where A and B are cations with variable valence. Typically, A is a divalent cation, e.g., Mg^{2+} , Fe^{2+} , Zn^{2+} or Mn^{2+} and B is a trivalent one, e.g., Al^{3+} or Fe^{3+} (the so-called 2-3 spinels). A well-known example is $MgAl_2O_4$ [1]. Spinel crystals offer good mechanical and thermal properties and they possess a broad transparency range.

3d transition-metal (TM) ions feature partially filled 3d orbitals which are not shielded. Their spectroscopic properties determined by transitions of 3d electrons are sensitive to the ligand field symmetry and strength. TM ions embedded in tetrahedral (T_d) sites in crystals are attractive because of their broadband linear and nonlinear absorption [2]. The latter arises from the fact that the T_d local field symmetry results in high transition intensities (several orders of magnitude stronger than those for the octahedral (O_h) field). As a result, crystals doped with TM ions entering into the T_d sites are used as saturable absorbers (SAs) of lasers [3-5].

Spinel and, in particular, $MgAl_2O_4$, are attractive for doping with TM ions [6] such as Ni^{2+} [7-10], Co^{2+} [11], Mn^{2+} [12-16], Fe^{2+} [17-19] or Cr^{3+} [20]. For $MgAl_2O_4$, one example is doping with divalent cobalt (Co^{2+}) ions replacing the Mg^{2+} cations in T_d sites and featuring broad absorption at $\sim 1.5 \mu m$ [11]. This absorption spectrally matches the emission of eye-safe erbium lasers. As a result, $Co:MgAl_2O_4$ based crystalline materials have been recognized as SAs of pulsed erbium lasers [21-24] with applications in range-finding, remote sensing (LIDAR), medicine, *etc.*

Zinc aluminate (gahnite) $ZnAl_2O_4$ is another well-known compound in the spinel crystal family [25]. In the mineral form, its color ranges between green, blue, yellow or brown due to the TM ion impurities. Compared to $MgAl_2O_4$, gahnite features better thermal conductivity. E.g., in [26] for ceramic $ZnAl_2O_4$, it was estimated to be ~ 25 W/mK. $ZnAl_2O_4$ exhibits high mechanical strength (Mohs hardness: 8.0), high thermal and chemical stability and wide band gap (~ 6 eV). Unfortunately, bulk single crystals of $ZnAl_2O_4$ cannot be obtained by conventional melt growth methods due to dissociation of zinc aluminate at high temperatures close to the melting point (about 1950 °C, depending on the impurities). As an alternative to single crystals, $Co:ZnAl_2O_4$ -based transparent glass-ceramics (GCs) [27,28] and $Co:ZnAl_2O_4$ ceramics [29] were proposed.

Polycrystalline transparent ceramics are obtained at temperatures significantly lower than the melting point of corresponding bulk crystals. Their spectroscopic properties resemble those of the crystals. As compared to GCs, they do not contain residual glass phase. The development of cobalt-doped spinel ceramics was manifested by $Co:MgAl_2O_4$ ones [30-35]. Recently, transparent $ZnAl_2O_4$ ceramics were obtained by Goldstein *et al.* by hot isostatic pressing at 1550 °C using hydrothermally synthesized powders [36]. Since then, several studies dedicated to undoped gahnite ceramics [37-40] and only few devoted to Co^{2+} -doped ones

[29] appeared. More recently, Ni²⁺-doped ZnAl₂O₄ transparent ceramics [9] were prepared by melting-vaporizing procedure using TeO₂ and Na₂O as melting agents.

One of the limiting factors in the work on obtaining this material is the absence of commercially available powders. The existing methods of synthesis of ZnAl₂O₄ powders have certain difficulties. In particular, in the case of simultaneous precipitation of zinc and aluminium nitrates with ammonia solution [41,42], due to formation of complex compounds of zinc with ammonia that are soluble in water, the resulting powder is enriched with aluminium. The same applies to cobalt, which is also coordinated by ammonia, so that it is impossible to control its exact content in the powder. According to literature data, powders synthesized by hydrothermal method [36], decomposition of sulphate salts [37,38], as well as mixing of powders of aluminium and zinc oxides [39,40] were used to fabricate ceramics. Recently, many works on the synthesis of powders have been published, among which are sol-gel synthesis [43], combustion synthesis [44-48], hydrothermal synthesis [14].

Recently, we showed that synthesis the ZnAl₂O₄ powders by hydrolyzing alcohol solutions of aluminium isopropoxide with an aqueous solution of zinc acetate [49] is promising for producing transparent ceramics. In this work, we optimize the ZnAl₂O₄ powder synthesis by replacing zinc acetate with zinc formate. Zinc formate easily decomposes [50] to form zinc oxide and a mixture of gases, i.e., CO, CO₂, H₂ and H₂O (↑), without formation of carbon residues in the temperature range of 250-300 °C. Other possible option is zinc nitrate. However, nitrate groups contribute to formation of hard particles, which are very difficult to grind in mills of any type. The transition to nitrate forms of metal sources will be possible using automated techniques for obtaining fine precursor powder, e.g., spray drying. Compared to nitrate sources, formate ones make it possible to obtain softer powders. The obtained powders are sintered to produce transparent ceramics by hot pressing in the presence of zinc fluoride (ZnF₂) [51] as a sintering additive. Also in this work, we propose to introduce ZnF₂ directly into the precursor sol, thus excluding an additional step of introduction of the sintering additive at later stages.

In the present paper, we describe the production of Co:ZnAl₂O₄ transparent ceramics by hot pressing from powders with different content of cobalt and sintering additive (ZnF₂). We also report on the effect of the sintering additive on microstructure, cobalt doping and spectroscopic properties of ceramics.

2. Experimental

2.1. Thermal evolution of precursor powders

The precursors for production of Co-doped ZnAl₂O₄ powders were prepared by hydrolysis of an alcohol solution of aluminium isopropoxide (Al(OCH(CH₃)₂)₃), joint aqueous solution of zinc formate (Zn(CH₂O)₂) and cobalt nitrate (Co(NO₃)₂), as well as aqueous solution of zinc fluoride (ZnF₂). The solubility of ZnF₂ in water is about 15 g/l (at 20 °C). The molar ratio (zinc + cobalt) / aluminium was 1:2. The procedure for the preparation of reagents and

hydrolysis is described in detail in [49]. The resulting precursor was dried in a vacuum box (at a pressure of 60 kPa) under heating up to 250 °C with a rate of 2 °C/min. The dried precursor was grinded in a planetary ball mill and calcined in air in the temperature range of 300 to 900 °C for 20 min. Zinc aluminate powders doped with 0.02 at.% or 0.10 at.% cobalt, and 3, 6 or 10 wt% zinc fluoride were obtained at the optimum temperature of 700 °C for 20 min.

The study of thermal behavior of the precursor and the choice of the heat-treatment regime were carried out by the thermal gravimetric analysis (TGA) and the differential thermal analysis (DTA) using a synchronous TGA / DTA analyzer (model DTG-60H, Shimadzu) in air at a heating rate of 5 °C/min. The phase composition of powders was identified by X-ray diffraction (XRD) using the LabX XRD-6100, Shimadzu diffractometer.

Zinc fluoride has high vapor pressure ($p(\text{ZnF}_2) = 7 \text{ Pa}$ at 775 °C [52], 53 Pa at 855 °C [53]). Calcination of the $\text{Co:ZnAl}_2\text{O}_4/\text{ZnF}_2$ precursor at high temperatures may lead to a significant loss of the sintering additive. To avoid this, a thermal gravimetric study of decomposition of the precursor was carried out. The TGA / DTA data of the 0.1 at.% $\text{Co:ZnAl}_2\text{O}_4 / 6 \text{ wt\% ZnF}_2$ dried precursor are shown in Fig. 1. The mass loss (about 7 wt%) at the temperatures up to 200 °C accompanying an endothermic peak (105 °C) corresponds to the removal of water. A small exothermic peak at ~245 °C may correspond to oxidation of formate groups by atmospheric oxygen and oxidation initiated by NO_3^- groups. The significant mass loss of ~25 wt% between 220 °C and 300 °C accompanying an endothermic peak (276 °C) corresponds to decomposition of formate groups. Further mass losses (5 %) starting at above 500 °C correspond to evaporation of the sintering additive of ZnF_2 (melting point: 872 °C) and removal of hydroxyl groups. Thus, significant losses comparable to the content of the sintering additive can be explained by the predominant evaporation of ZnF_2 . An exothermic peak at 850 °C corresponds the crystallization of zinc aluminate. The reason for the appearance of an exothermic peak with a maximum of 585 °C is not entirely understandable. Most likely, it is associated with the formation of intermediate fluorine-containing compounds because an exothermic peak at 585 °C was not observed on the TGA / DTA curve of the precursor without zinc fluoride, according to [49] and the laboratory protocol. To decrease the unwanted loss of the sintering additive, the powders should be calcined at the lowest possible temperature for the shortest possible time. The use of zinc in the formate form allowed us to decrease the temperature and time of powder calcination, as high temperatures are not required for removal of organic residues.

Fig. 2 shows XRD patterns of powders calcined at the temperatures in the range of 300–900 °C for 20 min. Gahnite crystallization begins at 500 °C, as confirmed by the appearance of weak and broad diffraction peaks. The calcination at 900 °C gives a significant increase in the crystal size and crystallinity of the powder, as indicated by notably narrowed diffraction peaks increased in intensity. The mean size of crystallites in the powders calcined at 700 °C and 900 °C was estimated using the Scherrer's equation [54] to be 8 nm and 37 nm, respectively. The crystallization of zinc aluminate at 500 °C and 700 °C occurs by a solid-

phase reaction, and at 900 °C - by dissolving and re-precipitation in the in the $\text{ZnAl}_2\text{O}_4 / \text{ZnF}_2$ eutectic melt. It should be noted that the XRD patterns of calcined powders do not contain reflexes of ZnF_2 . The mass loss of the $\text{Co:ZnAl}_2\text{O}_4 / \text{ZnF}_2$ precursor at above 700 °C observed during the TGA / DTA analysis (Fig. 1) is undesirable, so that the heat treatment of the powders at 700 °C is preferable.

Note that weak reflexes of zinc oxide (ZnO) are present in the XRD pattern of powder calcined at 900 °C. The impurity phase of ZnO may indicate that the crystallization of zinc aluminate has not been completed, but will be completed at hot pressing of ceramics. Note that these powders were not used for the synthesis of ceramics in the present work. If calcination of powders at higher temperatures would be required, the initial batch should be enriched in aluminium if the powder stoichiometry is shifted to zinc.

2.2. Fabrication of ceramics

To fabricate transparent ceramics, synthesized powders were hot pressed in a home-made machine at a temperature of 1520 °C for 4 h. The powders were pressed under 50 MPa in a steel mold (diameter: 15 mm) and placed into a graphite mold. The hot pressing included the following stages: (i) heating up to 950 °C with a constant pressure of 3 MPa, (ii) first dwell for 10 min with a constant pressure, (iii) heating up to 1200 °C with a gradual pressure rise up to 40 MPa, (iv) second dwell for 20 min with a constant pressure, (v) heating up to 1520 °C, (vi) final dwell for 4 h, (vii) free cooling. Sintering of ceramics was carried out in a vacuum atmosphere.

The general mechanism of action the sintering additive for liquid-assisted-sintering (ZnF_2 , in our case) includes (i) its role as a lubricant that promotes sliding of particles during shrinkage, (ii) recrystallization in the melt due to dissolution of small particles and precipitation on large particles and (iii) formation of sintering additive-associated vacancies that facilitate sintering at later stages of the sintering process.

The ceramic disks (thickness: 2 mm, diameter: 15 mm) were polished to laser-grade quality from both sides. They were transparent, Fig. 3, and had a blue coloration due to the cobalt doping. For a fixed Co content in the initial powders, the coloration decreased with increasing the ZnF_2 content (see below).

For spectroscopic studies, half of the ceramic disk was additionally annealed at 1400 °C for 30 min in air to eliminate the possible oxygen vacancies.

2.3. Characterization of ceramics

The X-ray powder diffraction (XRD) patterns of ceramic samples were recorded using a Shimadzu XRD-6000 diffractometer, Cu $K\alpha$ radiation (1.54060 Å) with a Ni filter. The lattice constant a for ceramics was determined by Rietveld refinement.

The microstructure of a clean fracture surface of ceramic disks was studied using a scanning electron microscope (SEM, MERLIN, Carl Zeiss). The size distribution of grains

was analyzed by manual counting using ImageJ software. The composition of ceramics was studied by Energy Dispersive X-ray (EDX) spectroscopy. For this, we used a similar electron microscope (accelerating voltage: 20 kV, beam current: 2 nA) equipped with an INCA X-Act (Oxford Instruments) microanalyzer. To avoid the unwanted charging of the dielectric ceramic surface, it was coated with a thin (thickness: 10 nm) carbon film prior to SEM studies.

All spectroscopic studies were performed at room temperature (20 °C). The transmission spectra were measured with a Shimadzu UV 3600 spectrophotometer (0.3–2 μm). The Raman spectra and the luminescence were detected using a confocal Raman microscope (in-Via, Renishaw) equipped using a Leica $\times 50$ objective (N.A. = 0.75). The excitation wavelength λ_{exc} was 514 nm (Ar⁺ laser line).

The luminescence decay was studied using a time-resolved confocal fluorescence microscope equipped with a $\times 100$ objective (N.A. = 0.95) (MicroTime 100, PicoQuant). As the excitation source, a picosecond (ps) diode laser emitting at 405 nm was used.

3. Results and discussion

3.1. Microstructure

The structure and the phase purity of ceramics were confirmed by X-ray powder diffraction, Fig. 4. The measured XRD patterns correspond to that of undoped ZnAl₂O₄ (according to the ICSD (Inorganic Crystal Structure Database) card #9559) [55]. Co:ZnAl₂O₄ is cubic (sp. gr. O_h⁷ – $Fd\bar{3}m$, #227). The lattice constant a for ceramics was weakly dependent on the ZnF₂ content. For ceramic prepared from powders containing 0.1 at.% Co and 10 wt% ZnF₂, $a = 8.088 \text{ \AA}$.

A similar microstructure was observed for all studied ceramics. The example SEM images of a fracture surface of ceramics prepared from powders containing 0.1 at.% Co and 3 / 10 wt.% ZnF₂ are shown in Fig. 5(a,b). The ceramics exhibit a close-packed structure with clean grain boundaries and a lack of μm -sized pores. The mean grain size was determined by counting about 100 crystallites for each sample. The measured linear dimensions of grains were multiplied by a shape factor of 1.2 (accounting for slight deviations from the spherical shape). Fig. 6(a) shows the grain size distribution for the series of ceramics prepared from powders doped with 0.02 at.% Co and 3–10 wt% ZnF₂. For all samples, a relatively broad range of grain sizes (20–100 μm) was observed revealing a dependence on the position in the ceramic disk: larger grains in the central part and smaller grains at the perimeter of the disk.

The effect of the ZnF₂ content on the mean grain size D_{grain} is shown in Fig. 6(b). For 0.02 at.% Co doping, with increasing the amount of the sintering aid, first, D_{grain} slightly increases (from 55 μm for 3 wt% ZnF₂ to 68 μm for 6 wt% ZnF₂) and then decreases to 60 μm for 10 wt% ZnF₂. A similar tendency is observed for 0.1 at.% Co doping. These values are larger than those observed in an undoped ZnAl₂O₄ ceramics ($D_{\text{grain}} = 40 \mu\text{m}$, using the same shape factor) produced by a similar method with 5 wt% ZnF₂ [51]. The observed trends for

the variation of the grain size may be masked by the relatively broad grain size distribution observed in the studied ceramics; therefore, it is difficult to draw conclusions in this relation.

The composition of ceramics was studied by EDX. The presence of fluorine was detected in all the studied samples. The concentration of F⁻ ions in ceramics slowly increased with the ZnF₂ content in the precursor powders. We assume that the F⁻ ions may be diffusely distributed at the grain boundaries or dissolved in the grains. The EDX spectra of ceramics prepared from powders containing 0.1 at.% Co and 3-10 wt% ZnF₂ are shown in Fig. 7. The element mapping (Al, Zn, O, Co and F) obtained by the means of EDX is shown in Fig. 8.

3.2. Raman spectra

The structure of ceramics was confirmed by Raman spectroscopy, Fig. 9. According to the group-theory analysis, the irreducible representations at the Γ point ($\mathbf{k} = 0$) of the Brillouin zone are as following: $\Gamma = A_{1g} + E_g + T_{1g} + 3T_{2g} + 2A_{2u} + 2E_u + 5T_{1u} + 2T_{2u}$ [56]. Five modes (A_{1g} , E_g and $3T_{2g}$) are Raman-active, five ($5T_{1u}$) are IR-active and other are acoustic or silent. In the Raman spectra of ceramics, all five modes are observed at 195 cm⁻¹ (T_{2g}), 417 cm⁻¹ (E_g), ~510 cm⁻¹ (T_{2g}), 657 cm⁻¹ (T_{2g}) and ~800 cm⁻¹ (A_{1g}). The spectra are weakly dependent on the starting composition. The two most intense modes are due to the motion of Al and O atoms [57].

A careful inspection of the Raman spectra did not reveal characteristic vibrations of possible fluoride phases (CoF₂ or ZnF₂) and ZnO. In particular, for CoF₂, these are 68, 246, 366 and 494 cm⁻¹ [58] and for and for ZnF₂ – 70, 253, 350 and 522 cm⁻¹ [59]. For ZnO, the characteristic Raman bands are at 99, 333 and 438 cm⁻¹ [60].

3.3. Optical absorption

The in-line transmission (T) spectra of polished ceramic disks are shown in Fig. 10(a). First, let us discuss the transmission at 1.9 μm (out of cobalt absorption bands). Depending on the composition of the precursor, it amounts to $T = 83.2\%$ - 84.0% , which is close to the theoretical value set by Fresnel losses, $T_0 = 84.3\%$, assuming a refractive index n of 1.80 [61]. High transmission indicates good optical quality of ceramics. For the increased cobalt concentration in the precursor, the transmission slightly decreases. However, we did not observe a direct relation between transmission and the ZnF₂ content.

The observed spectral bands are due to absorption of Co²⁺ ions in tetrahedral sites in the spinel structure. The Co²⁺ ion has the electronic configuration of [Ar]3d⁷. The ground-state is ⁴A₂(⁴F), so that the spin-allowed transitions occur to quartet states, ⁴F and ⁴P. However, the transition to the lower-lying excited-state ⁴T₂(⁴F) is forbidden in an undistorted T_d ligand field, so it has a small oscillator strength [11] and it is observed as a weak band at ~2.55 μm . The intense and broad (1.06–1.64 μm) absorption band with local peaks at 1248, 1297, 1428, and 1503 nm corresponds to transitions to the ⁴T₁(⁴F) state. The second intense band in the visible (0.50–0.68 μm) with local peak at 547, 587 and 620 nm corresponds to transitions to

the ${}^4T_1({}^4P)$ state. The local peaks in both bands are related to the spin-orbit splitting of the quartet states [30]. For example, for the ${}^4T_1({}^4F)$ state of tetrahedral Co^{2+} ions in gahnite, the previously determined energies of the spin-orbit splitted sub-levels of ${}^4T_1({}^4P)$ at 4.2 K are 6756(Γ_6), 7072(Γ_8), 7776(Γ_7) and 7850(Γ_8) cm^{-1} [25], corresponding to the wavelengths in absorption of 1480, 1414, 1286 and 1273 nm, respectively. For the studied ceramics, the positions of sub-levels may slightly vary as compared to the single-crystal.

Besides the spin-allowed transitions, the spectra contain weak bands assigned to transitions to the higher-lying splitted 2G , 2P and 2H doublet excited-states. In particular, the transitions to the 2E , 2T_1 , 2A_1 and ${}^2T_2({}^2G)$ excited-states (in an increasing energy order) contribute to intense absorption band in the visible. At shorter wavelengths ($<0.50 \mu m$), weak structured absorption is due to transition to the ${}^2T_1({}^2P)$, 2T_2 , 2T_1 , 2E and ${}^2T_1({}^2H)$ excited-states (in an increasing energy order). Here, the assignment is according to the energy levels reported in [25,62]. More details about the crystal-field of tetrahedral Co^{2+} ions in gahnite can be found elsewhere [63]. Weak and broadband absorption at these wavelengths may also originate from Co^{2+} ions in octahedral sites [64] (see Section 4.4).

A detailed view on visible absorption of an example ceramic sample made of powders doped with 0.1 at.% Co and 3 wt% ZnF_2 is given in Fig. 10(b). The bands at 477 nm, ~445 nm and 406 nm are assigned to transitions to the ${}^2T_1({}^2P)$, 2T_2 and ${}^2T_1({}^2H)$ states, respectively. Other transitions to the 2H doublet states are not visible because of their weak intensity. For the as-sintered ceramic, a broad absorption band at ~275 nm is observed overlapping with the UV absorption edge ($\lambda_{UV} = 245$ nm, optical bandgap E_g : ~5.5 eV). After annealing, the Co^{2+} absorption features do not change implying that new Co^{2+} sites with T_d symmetry are not formed, while the intensity of the UV band increases significantly thus shifting the absorption edge to $\lambda_{UV} = 292$ nm. In the previously studied undoped $ZnAl_2O_4$ ceramics sintered with 5 wt% ZnF_2 [49], no such absorption was detected resulting in $E_g = 6.05$ eV. In the present work, sintering of ceramics was carried out in a vacuum atmosphere, which may lead to formation of oxygen vacancies. Oxidative annealing contributes to the drain of vacancies (see Section 4.5) with formation of pores of nm and sub- μm size leading to additional light scattering and a red-shift of the UV absorption edge of the material.

The scheme of energy levels of tetrahedral Co^{2+} ions can be analyzed within the equivalent Tanabe-Sugano diagram [65] for the d^3 complexes in octahedral field, Fig. 11(a). Here, we omit the “g” subscripts because all the states are even (*gerade*). The Co^{2+} absorption in the $ZnAl_2O_4$ matrix corresponds to the region of the diagram where the branches corresponding to the 4P and 2G states intersect. As a result, the doublet states 2E and ${}^2T_1({}^2G)$ are lying below the quartet ${}^4T_1({}^4P)$ one, as shown in Fig. 11(b) according to the crystal-field splitting from [25]. As compared to $Co:MgAl_2O_4$ [66], gahnite shows smaller splitting of the ${}^4T_1({}^4F)$ and ${}^4T_1({}^4P)$ states.

3.4. Cobalt doping

A general chemical formula for spinels is AB_2O_4 (cubic system, sp. gr. $Fd\bar{3}m$). Each unit cell contains 32 anions (O^{2-}) in a face-centered cubic packing. Inside the oxygen lattice, the cations (A^{2+} and B^{3+}) have 32 octahedral (O_h) and 64 tetrahedral (T_d) available interstitial sites per unit cell. In a normal spinel structure, 1/8 of the T_d sites are occupied by divalent cations A^{2+} and 1/2 of the O_h sites are occupied by trivalent cations B^{3+} . Spinel can demonstrate a certain degree of cation disorder. It is described by an inversion parameter δ which is a fraction of B^{3+} cations in the T_d sites (or, equivalently, a fraction of A^{2+} cations in the O_h sites). As a result, the formula for spinels may be written as $^{IV}[A_{1-\delta}B_\delta]^{VI}[B_{2-\delta}A_\delta]O_4$. The spinel is normal for $\delta = 0$ and inverse for $\delta = 1$. For $\delta = 2/3$, cations have a completely random distribution over the T_d and O_h sites [63].

In the case of gahnite, this formula is $^{IV}[Zn_{1-\delta}Al_\delta]^{VI}[Al_{2-\delta}Zn_\delta]O_4$. Here, the superscripts are the coordination numbers (C.N. = IV and VI for the T_d and O_h sites, respectively). The Co^{2+} ions in the spinel structure are expected to replace the Zn^{2+} ones. In this way, some part of Co^{2+} ions may be located in octahedral sites [63,67]. Because a center of symmetry is present for the O_h site, the intensity of transitions for $^{VI}Co^{2+}$ ions will be several orders of magnitude weaker as compared to the $^{IV}Co^{2+}$ ones [25]. Thus, the former species will contribute to absorption with much lower intensity.

It is known that $Co:ZnAl_2O_4$ exhibits a notable inversion, depending on the size of the crystallites and Co^{2+} concentration. With increasing the Co^{2+} doping level reaching the stoichiometric composition of cobalt aluminate ($CoAl_2O_4$), δ increases [67]. It was also shown that the inversion parameter decreases with increasing the crystallite size [64]. However, the presence of a notable fraction of octahedral Co^{2+} ions in bulk gahnite crystals was also claimed [25].

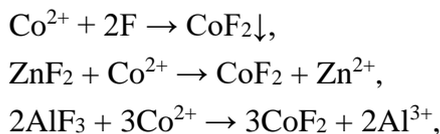
It is known that the lattice constant of Co^{2+} -doped $ZnAl_2O_4$ is an indication of a degree of inversion. For an undoped $ZnAl_2O_4$ ceramics studied for comparison, we have determined $a = 8.087 \text{ \AA}$ which is a typical value for undoped synthetic gahnite [68]. The assumption that Co^{2+} ions are entering into the spinel structure solely in T_d sites should lead to a decrease of a . Indeed, for a C.N. = IV, the ionic radii of Co^{2+} and Zn^{2+} are 0.58 \AA and 0.60 \AA , respectively [69]. In [67], it was suggested that the main reason for increasing a is the intersite-cation exchange (quantified by increasing δ parameter), namely, Co^{2+} for Al^{3+} substitution on the O_h sites. Indeed, the corresponding ionic radii of Co^{2+} and Al^{3+} for C.N. = VI are 0.745 \AA and 0.535 \AA [69].

Now, let us analyze the broad absorption band due to the $^4A_2(^4F) \rightarrow ^4T_1(^4F)$ transition of $^{IV}Co^{2+}$ ions, Fig. 12(a). For a fixed concentration of Co^{2+} in the precursor, the absorption coefficient (α_{abs}) decreases linearly with increasing the ZnF_2 content, e.g., as shown in Fig. 12(b) for the reference wavelength of $1.54 \mu m$. Moreover, even extrapolation of the $\alpha_{abs}(ZnF_2)$ dependence to the limit of absent sintering additive gives a lower value of the absorption coefficient than the expected one, $\alpha_{abs0} = \sigma_{GSA} \cdot N_{Co0}$. Here, σ_{GSA} is the ground-state absorption (GSA) cross-section of Co^{2+} ions in T_d sites, N_{Co0} is the Co^{2+} concentration assum-

ing that all cobalt ions enter into the ceramic grains. For the initial concentrations of 0.02 at.% and 0.1 at.% Co, $N_{\text{Co}0} = 0.303 \times 10^{19} \text{ cm}^{-3}$ and $1.515 \times 10^{19} \text{ cm}^{-3}$, respectively (as calculated assuming the density of ceramics $\rho = 4.61 \text{ g/cm}^3$). Previously, for transparent GCs based on Co:ZnAl₂O₄ nanocrystals, σ_{GSA} at 1.54 μm was determined to be $2.0 \times 10^{-19} \text{ cm}^2$ [27] and $2.45 \times 10^{-19} \text{ cm}^2$ (used in the present work) [30] being lower than that for the Co:MgAl₂O₄ single-crystal, $3.5 \times 10^{-19} \text{ cm}^2$ [3]. A similar analysis can be performed based on the visible absorption band of Co²⁺ ions. In particular for ZnAl₂O₄, it is complicated by the lack of reliable data on the GSA cross-sections for the $^4\text{A}_2(^4\text{F}) \rightarrow ^4\text{T}_1(^4\text{P})$ transition.

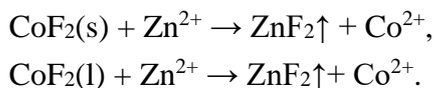
From the measured absorption coefficient α_{abs} at 1.54 μm , we estimated the actual concentration of Co²⁺ ions in T_d sites $N_{\text{Co}}(T_d) = \alpha_{\text{abs}}/\sigma_{\text{GSA}}$ (Table 1). It decreases linearly with the ZnF₂ content and it is about 2 times lower than the $N_{\text{Co}0}$ value. There can exist two reasons for this. One is the presence of a fraction of Co²⁺ ion in O_h sites (as described above) that does not contribute to linear absorption at $\sim 1.54 \mu\text{m}$. A similar mechanism was proposed previously for Co:ZnAl₂O₄-based transparent glass-ceramics to explain their non-saturable losses [28]. However, as all the studied ceramics are synthesized under the same conditions, this fraction is expected to be nearly independent of the ZnF₂ content. Consequently, another reason is a chemical process involving ZnF₂ and occurring during sintering of ceramics.

Let us describe the role of ZnF₂. We suppose that cobalt (II) fluoride (CoF₂) is the key intermediate of the sintering process. CoF₂ can be formed both at the stage of precursor synthesis and during the sintering itself (through an interaction of ZnF₂ and AlF₃ (if it is formed during the evolution of a precursor or during the sintering) with cobalt ions within the green body), according to the following reactions:



respectively.

The formed CoF₂ is less volatile ($\sim 12 \text{ Pa}$ at 950 °C [70]) than ZnF₂ ($\sim 458 \text{ Pa}$ at 950 °C, extrapolated from [53]) and AlF₃ ($\sim 395 \text{ Pa}$ at 950 °C [71]), hence, it is removed later (at higher temperatures). Indeed, the melting points of ZnF₂ and CoF₂ are 872 °C and $\sim 1200 \text{ °C}$, respectively. CoF₂ may be removed by evaporation and / or sublimation. The AlF₃ vapor pressure is lower than that of ZnF₂ but of the same order; therefore, sintered ceramics can be slightly enriched with zinc (see Fig. 2, 900 °C), as AlF₃ can also evaporate. The following equations describe the process of regeneration of spinel lattice according to the reverse exchange reactions ((l) and (s) stand for liquid and solid phases, respectively):



If the rate of these processes is less than the rate of evaporation / sublimation of CoF₂, the actual Co²⁺ concentration in the ceramics will decrease. Here, CoF₂(l) is a melt. The melting point of CoF₂ is $\sim 1200 \text{ °C}$ and during the hot pressing, cobalt fluoride which has not been

evaporated yet, will be completely liquid. In more details, taking into account the thermodynamic approach, similar processes are considered for the $\text{MgAl}_2\text{O}_4 / \text{LiF}$ system in [72].

Separately, it is worth noting that there is a possibility of presence of diffusely distributed CoF_2 at the grain boundaries as a residue, because fluorine is found in ceramics at the level of the detection limit (Section 4.1). However, this option does not change the model of probable formation of CoF_2 .

Note that we did not use $\text{Co:ZnAl}_2\text{O}_4$ ceramics fabricated without the sintering aid for the analysis in Fig. 12(b) as they were translucent and it was impossible to quantify the Co^{2+} absorption.

To conclude, the reduction in Co^{2+} concentration in T_d sites with respect to the value obtained when extrapolating the $\alpha_{\text{abs}}(\text{ZnF}_2)$ dependence to the limit of no sintering aid is due to the ZnF_2 -assisted evaporation of cobalt in the form of CoF_2 . The residual difference in concentrations (45% for 0.1 at.% Co and 28% for 0.02 at.% Co) is ascribed to the inverse spinel structure (predominantly) and losses of cobalt not related to ZnF_2 . Indeed, cobalt may also reduce to the metallic form with the zero oxidation state (e.g., due to the reaction with CO gas or graphite). This will however lead to formation of pores and translucent character of ceramics which is not observed in our case. It should be also noted that oxidative annealing would lead to the oxidation of metallic cobalt and the entry of Co^{2+} into the spinel lattice, as well as to an increase in the intensity of cobalt absorption bands. However, it is not noted (see Fig. 10(b)). Thus, this mechanism seems to play a minor role.

3.5. Luminescence

The luminescence spectra of as-sintered $\text{Co:ZnAl}_2\text{O}_4$ ceramics are shown in Fig. 13(a). The spectra contain an intense band centered at ~ 645 nm (emission bandwidth: 43 nm) and a weak and broad band centered at ~ 860 nm both ascribed to Co^{2+} ions in T_d sites. The spin-allowed transitions in emission occur from the ${}^4T_1({}^4P)$ state to the ${}^4A_2({}^4F)$ and ${}^4T_2({}^4F)$ ones, respectively. As mentioned above, the doublet states 2E and ${}^2T_1({}^2G)$ in $\text{Co:ZnAl}_2\text{O}_4$ are lying below the ${}^4T_1({}^4P)$ state while their populations are thermalized at room-temperature. Thus, ${}^2E({}^2G)$ is the emitting state.

With the increase of the actual Co^{2+} concentration in T_d sites (cf. Table 1), or, equivalently, with the decrease of the ZnF_2 content in the precursor powders, the luminescence intensity gradually increases while no significant change of the spectral shape is observed. No concentration-quenching is observed up to at least 0.050 at.% Co in T_d sites (actual).

We have monitored the effect of annealing on the emission properties of ceramics, see Fig. 13(b). After annealing, the intensity of luminescence related to Co^{2+} ions in T_d sites notably increases while the emission ascribed to defects is suppressed. This is because annealing efficiently removes oxygen vacancies leading to better crystallinity of the material.

Other spectral features are the sharp emission lines at 676–723 nm and a weak and broad emission centered at ~ 740 nm, Fig. 13(c). Previously, we demonstrated [29] that gahn-

ite ceramics contains chromium (Cr^{3+}) impurity ions that come with ZnO raw material. The sharp emission lines at 676–723 nm are due to Cr^{3+} ions substituting the Al^{3+} cations in O_h sites (the so-called N-lines) [73,74]. The broad emission centered at ~740 nm is assigned to the structure defects [75,76]. In particular, the transitions from the conduction band (CB) or shallow donor defect states (e.g., Al^*_{Zn} – aluminium antisite defects) to deep donor defect states (V^*_{O} – oxygen vacancies) may explain this emission.

The luminescence decay was studied for the as-sintered ceramic samples. When exciting the samples at 405 nm and detecting luminescence in the whole visible spectral range, the measured luminescence decay curves exhibited intense “fast” time component (decay time: $\tau_{\text{fast}} = 3.0 \pm 0.2$ ns, being weakly dependent on composition of the starting powders) assigned to emission of defects, and much weaker “slow” component ($\tau_{\text{slow}} \sim$ hundreds of ns) attributed to emission of Co^{2+} ions. Indeed, by spectral filtering only the emission band at ~645 nm, see Fig. 13(a), we observed predominantly the latter decay. The luminescence decay curves were well fitted with a single-exponential law, Fig. 14. For ceramics obtained from powders doped with 0.1 at.% Co, the determined decay time τ_{slow} decreased in the range 272–222 ns for the ZnF_2 content decreasing from 10 to 3 wt% (or, equivalently, with increasing concentration of Co^{2+} ions in T_d sites). The τ_{slow} value has a meaning of the lifetime of the ${}^2\text{E}({}^2\text{G})$ Co^{2+} state. Our observations agree well with the previous studied of $\text{Co}:\text{ZnAl}_2\text{O}_4$ -based GCs [27,28]: it was determined that the lifetime of the ${}^2\text{E}({}^2\text{G})$ excited-state of Co^{2+} ions in T_d sites decreases from 200 to 25 ns for increasing the concentration of CoO in the initial glass (0.1–2 wt%).

As a final remark, one may point out the cost effectiveness and simplicity of synthesis of Co^{2+} -doped ZnAl_2O_4 nanopowders and transparent ceramics in the present work. This is in line with the recent studies [77,78] focusing on similar materials.

4. Conclusions

We report on fabrication of a series of highly transparent cobalt-doped zinc aluminate spinel (gahnite) ceramics – $\text{Co}^{2+}:\text{ZnAl}_2\text{O}_4$. For this, we employed a modified approach with the following features: (i) an original way of fabrication of the precursor nanopowders by using zinc *formate* as a source of zinc and an aqueous solution of zinc fluoride as a sintering additive. The first detail allowed us to reduce the temperature and time of calcination of powders (as formate easily decomposes at 250–300 °C) without carbonization of the powder (formate does not contain C–C bonds). The second detail is the preparation of the precursor using solutions of metal sources *together* with an aqueous solution of the sintering additive (ZnF_2), which eliminates the need for additional steps for its introduction at later stages. Such principles have never been used before for ZnAl_2O_4 transparent ceramics; (ii) an optimum temperature of powder calcination (700 °C) determined from thermal studies and reducing losses of the sintering additive (ZnF_2), (iii) the particular temperature of hot-pressing (1520 °C) to produce transparent ceramics. With this improved chemical way, we obtained ceram-

ics with excellent transparency close to the theoretical limit and suitable microstructure (relatively small grains, clear grain boundaries and the lack of μm -sized pores).

We determined that the concentration of Co^{2+} ions in T_d sites in the ceramic specimens is lower than the initial concentration in the precursor powders and, furthermore, it decreases with the ZnF_2 content. This is ascribed to two main reasons: (i) accommodation of Co^{2+} ions in octahedral sites (inversion of spinel structure) and the loss of Co^{2+} via evaporation of an intermediate phase of CoF_2 . A linear variation of the tetrahedral Co^{2+} concentration with the ZnF_2 content allowed us to control precisely the absorption at around $\sim 1.5 \mu\text{m}$ (the ${}^4\text{A}_2({}^4\text{F}) \rightarrow {}^4\text{T}_1({}^4\text{F})$ transition of ${}^{\text{IV}}\text{Co}^{2+}$ ions) without deterioration of optical quality of ceramics. The latter is of practical importance for development of ceramic saturable absorbers for eye-safe erbium lasers. Further efforts should be applied on revealing the effect of octahedral Co^{2+} ions on nonlinear properties of $\text{Co}:\text{ZnAl}_2\text{O}_4$ ceramics (e.g., via non-saturable losses).

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List of figure captions

Fig. 1. Thermal evolution of the 0.1 at.% Co:ZnAl₂O₄ / 6 wt% ZnF₂ precursor: thermal gravimetric analysis (TGA) mass loss curve (*black*) and differential thermal analysis (DTA) curve (*blue*) measured under heating in air.

Fig. 2. X-ray diffraction (XRD) patterns of the powders doped with 0.1 at.% Co and 6 wt% ZnF₂ annealed at 300, 500, 700 and 900 °C. *Circles* indicate the diffraction peaks of ZnAl₂O₄, *asterisks* – ZnO.

Fig. 3. Photographs of transparent ceramics sintered at 1520 °C for 4 h from 0.02 at.% Co (*upper row*) and 0.1 at.% Co (*lower row*) doped ZnAl₂O₄ powders with 3, 6 or 10 wt% ZnF₂.

Fig. 4. X-ray powder diffraction (XRD) patterns of ceramic samples prepared from powders containing 0.1 at.% Co and 3 / 10 wt% ZnF₂, *numbers* denote the Miller's indices (*hkl*), *red peaks* – diffraction peaks for undoped ZnAl₂O₄ (ICSD card #9559).

Fig. 5. Scanning Electron Microscope (SEM) images of a fracture surface of ceramics produced from powders containing 0.1 at.% Co and (a) 3 wt% ZnF₂ and (b) 10 wt% ZnF₂.

Fig. 6. (a) Grain size distribution for ceramics produced from powders containing 0.02 at.% Co and 3, 6 or 10 wt% ZnF₂; (b) Mean grain size vs. the ZnF₂ concentration in the precursor powders.

Fig. 7. EDX spectra for ceramics prepared from powders containing 0.1 at.% Co and 3-10 wt% ZnF₂.

Fig. 8. EDX mapping for ceramics prepared from powders containing 0.1 at.% Co and 10 wt% ZnF₂: (a) SEM image of the fracture surface, (c-f) the corresponding element maps: (b) Al, (c) O, (d) Zn, (e) F and (f) Co.

Fig. 9. Raman spectra of Co:ZnAl₂O₄ transparent ceramics (the spectra are shifted for the sake of comparison), $\lambda_{exc} = 514$ nm.

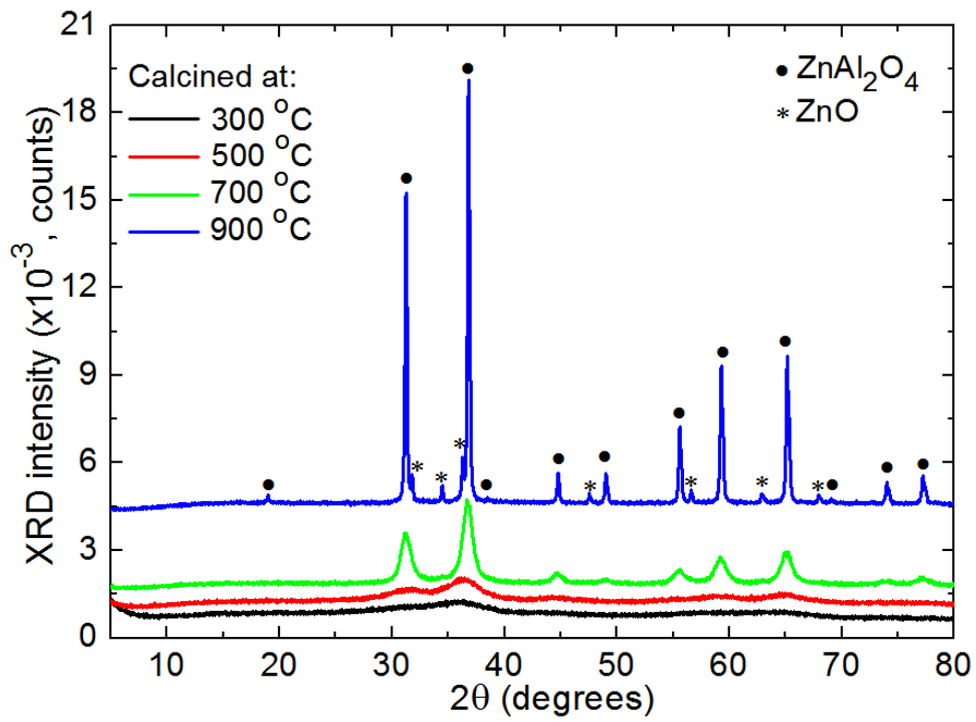
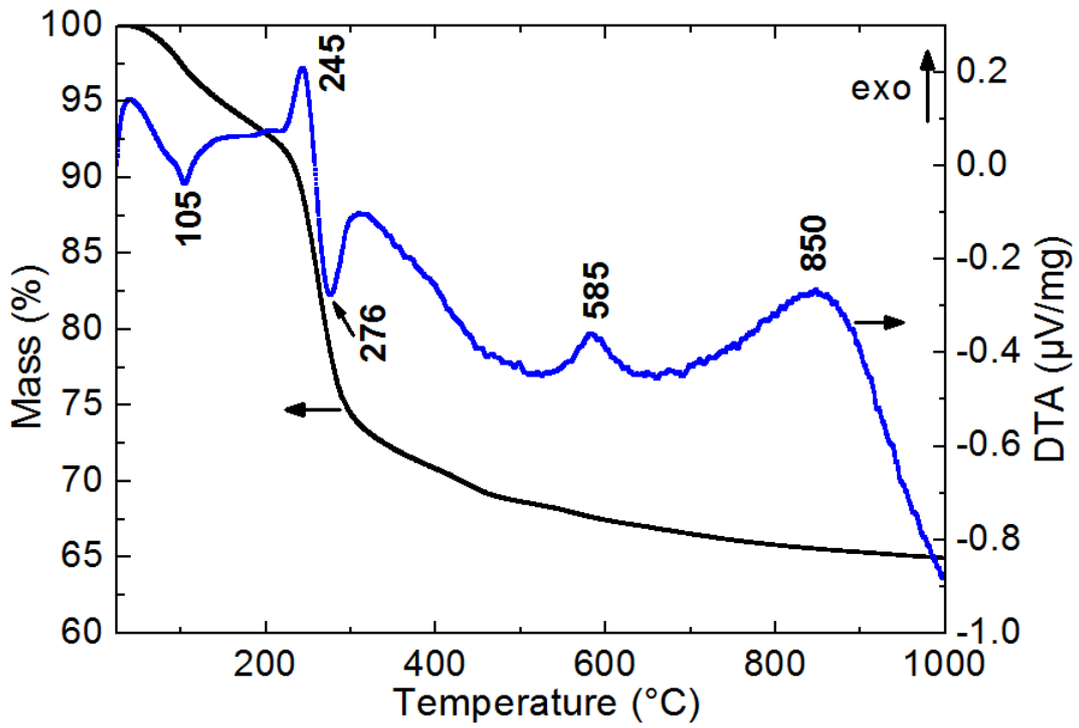
Fig. 10. In-line transmission spectra of laser-grade polished Co:ZnAl₂O₄ ceramic disks (thickness: 2 mm): (a) transmission of as-sintered samples; (b) effect of annealing on transmission in the UV and visible of ceramics prepared from powders doped with 0.1 at.% Co and 3 wt% ZnF₂. *Numbers* in (b) indicate the peak wavelengths.

Fig. 11. (a) Tanabe-Sugano diagram of energy levels of d⁷ tetrahedral complexes (for $C/B = 4.50$), *vertical dashed line* marks the case of Co²⁺:ZnAl₂O₄; (b) crystal-field splitting of Co²⁺ ions in ZnAl₂O₄ and MgAl₂O₄ crystals (after [25,65]). *Green arrows* indicate the observed transitions in emission.

Fig. 12. Analysis of the ⁴A₂(⁴F) → ⁴T₁(⁴F) absorption transition of Co²⁺ ions in T_d sites in Co:ZnAl₂O₄ ceramics: (a) absorption spectra; (b) variation of absorption coefficient α_{abs} at ~1.54 μm with the ZnF₂ concentration in the precursor: *dashed lines* – expected absorption calculated from the GSA cross-section and the nominal Co²⁺ concentration.

Fig. 13. Luminescence of Co:ZnAl₂O₄ transparent ceramics: (a) spectra of non-annealed samples; (b) effect of annealing on the luminescence spectra of ceramics prepared from powders doped with 0.02 / 0.1 at.% Co and 3 wt% ZnF₂; (c) details of impurity and defect emissions for ceramics prepared from powders doped with 0.02 at.% Co and 3 wt% ZnF₂, *numbers* indicate the peak wavelengths. $\lambda_{\text{exc}} = 514$ nm.

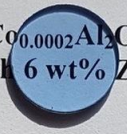
Fig. 14. Luminescence decay curves of Co:ZnAl₂O₄ transparent ceramics prepared from powders doped with 0.1 at.% Co and 3–10 wt% ZnF₂ (semi-log plots): *circles* – experimental data, *solid lines* – their single-exponential fits, τ_{slow} – luminescence lifetime. $\lambda_{\text{exc}} = 405$ nm (ps pulses), $\lambda_{\text{lum}} \sim 650$ nm.



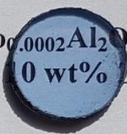
$\text{Zn}_{0.9998}\text{Co}_{0.0002}\text{Al}_2\text{O}_4$ powder
with 3 wt% ZnF_2



$\text{Zn}_{0.9998}\text{Co}_{0.0002}\text{Al}_2\text{O}_4$ powder
with 6 wt% ZnF_2



$\text{Zn}_{0.9998}\text{Co}_{0.0002}\text{Al}_2\text{O}_4$ powder
with 10 wt% ZnF_2



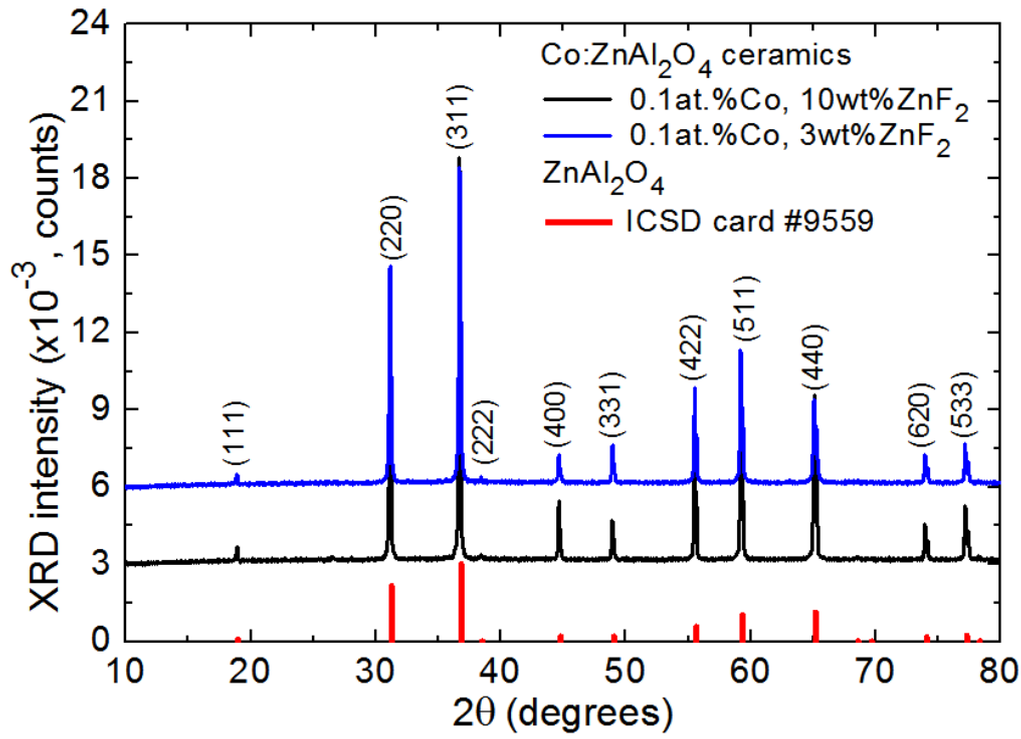
$\text{Zn}_{0.999}\text{Co}_{0.001}\text{Al}_2\text{O}_4$ powder
with 3 wt% ZnF_2

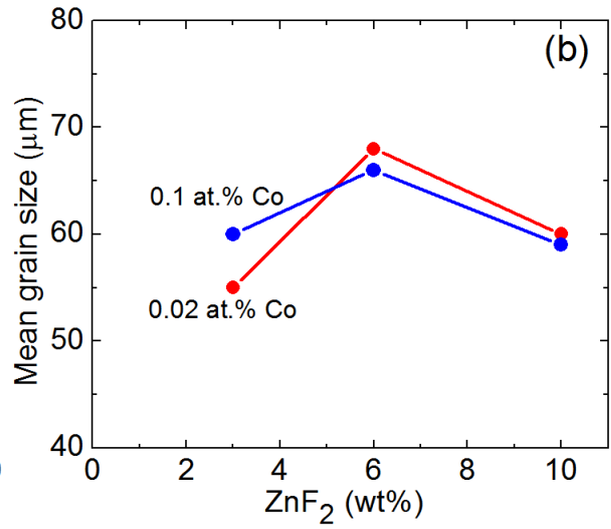
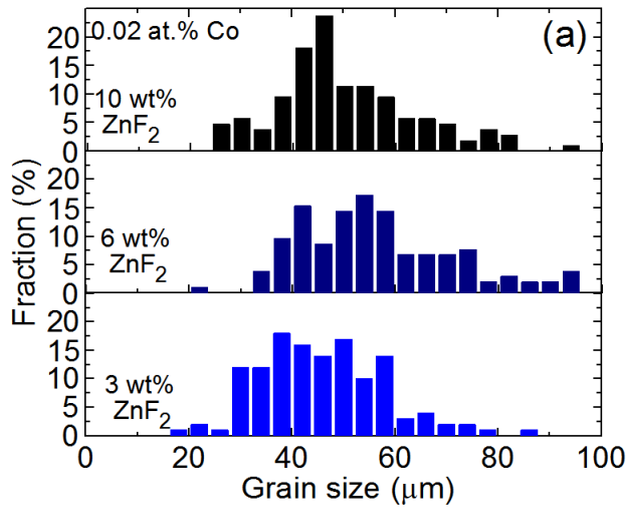
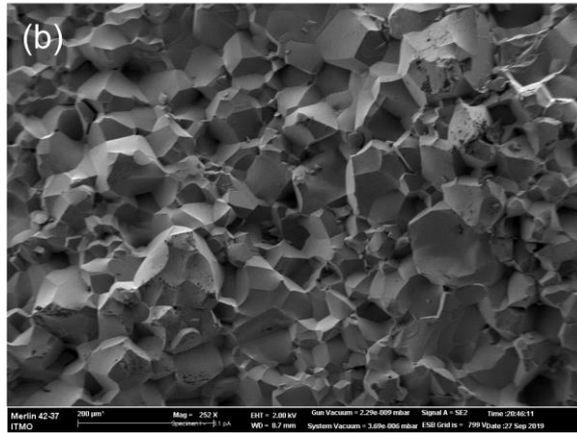
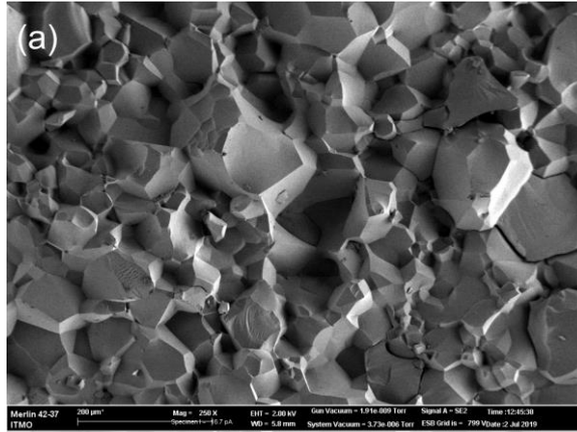


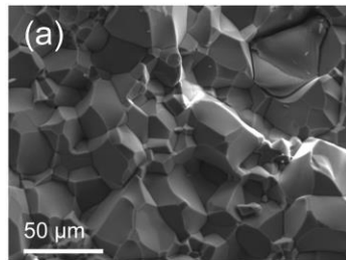
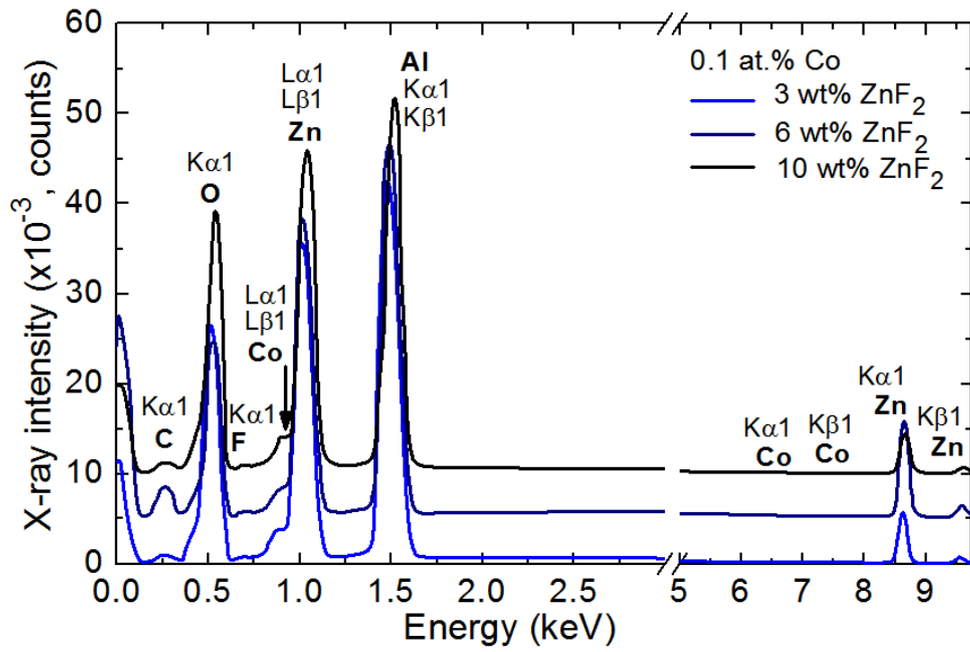
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with 6 wt% ZnF_2



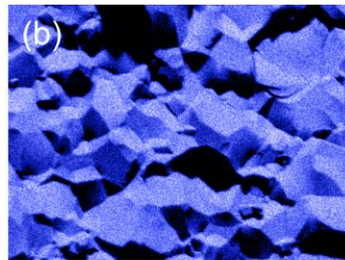
$\text{Zn}_{0.999}\text{Co}_{0.001}\text{Al}_2\text{O}_4$ powder
with 10 wt% ZnF_2



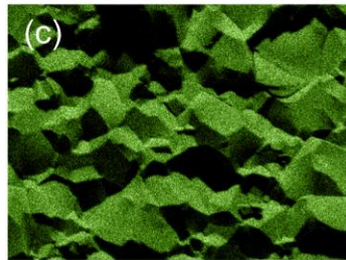




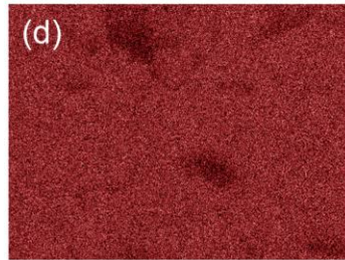
SEM



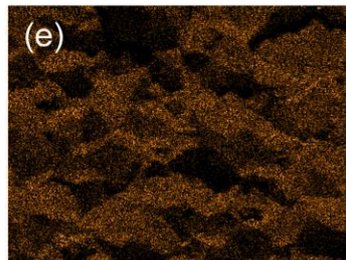
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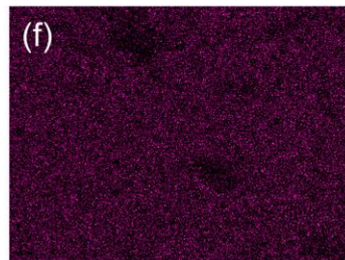
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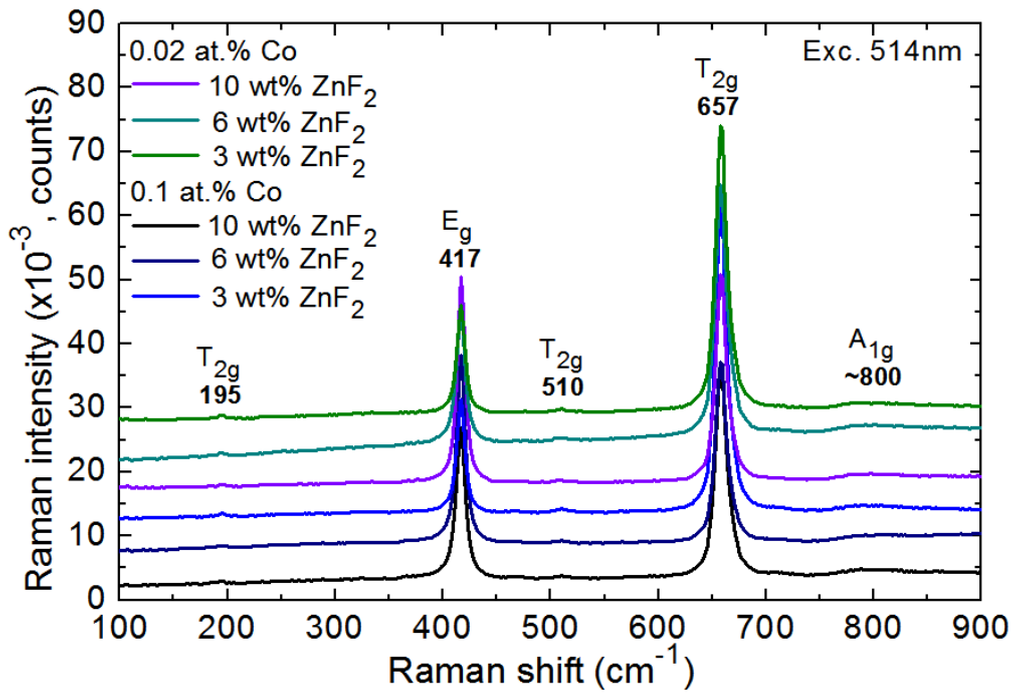
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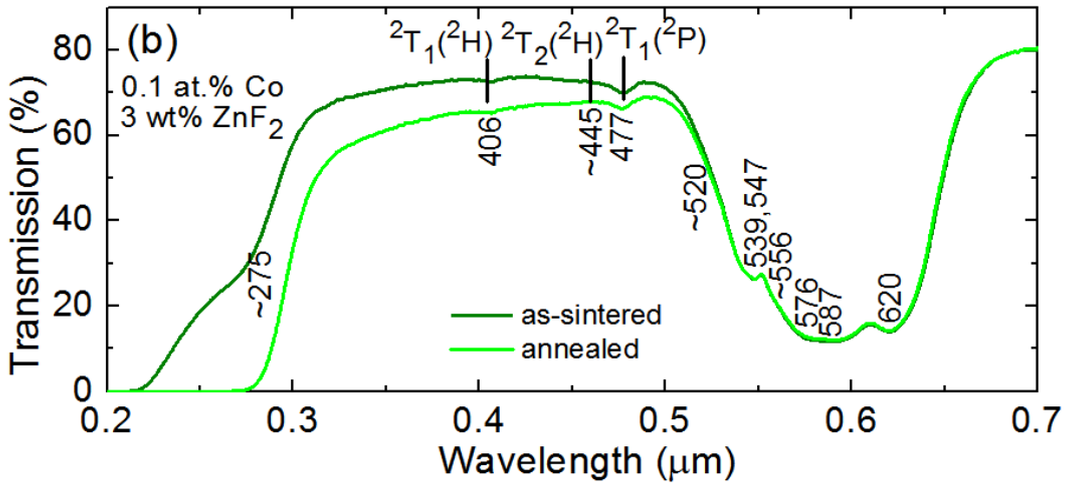
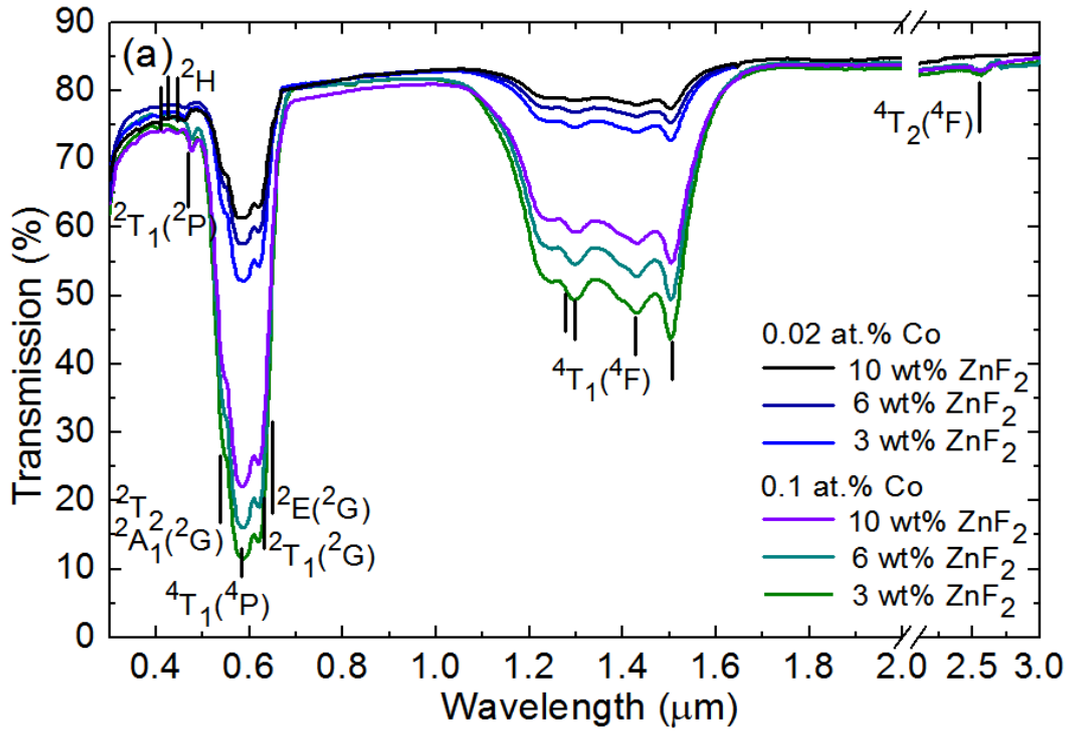


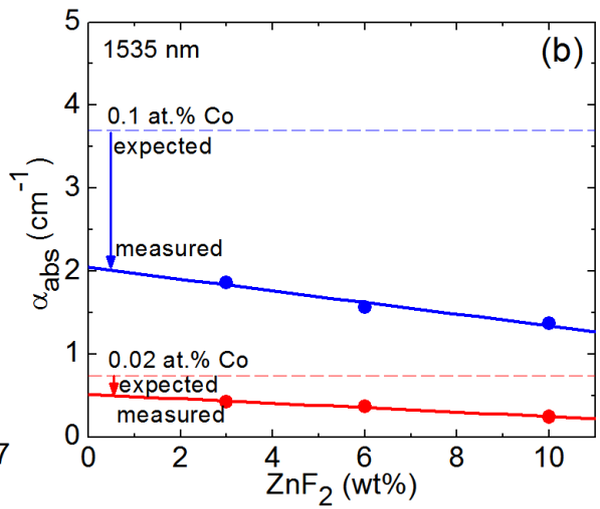
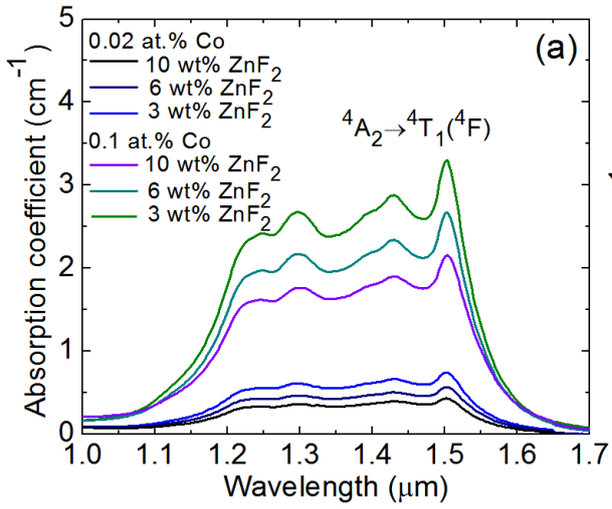
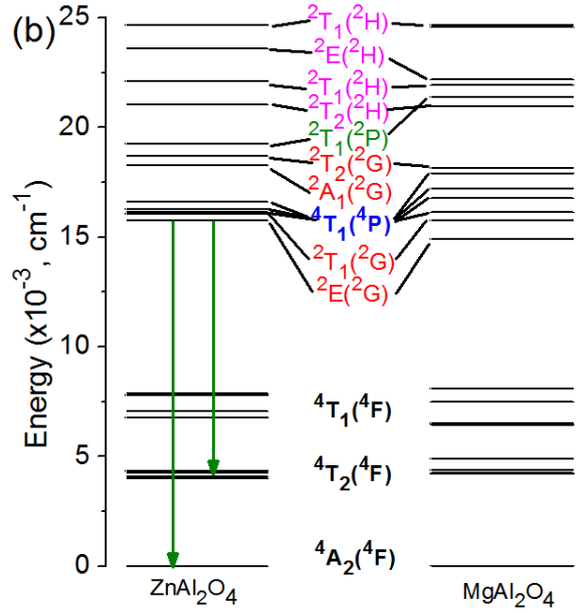
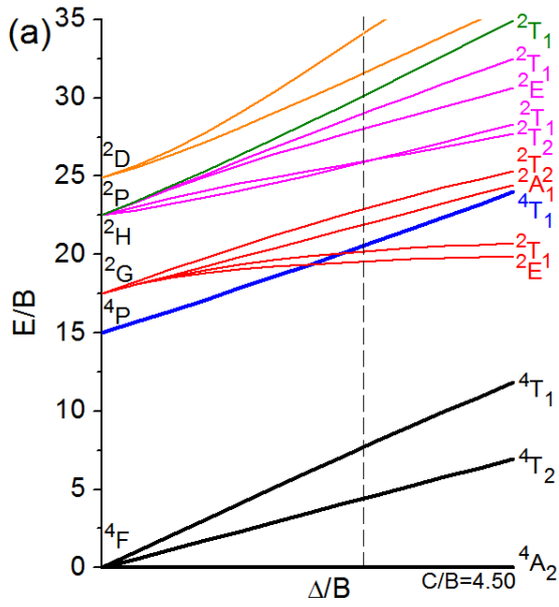
F K α 1,2

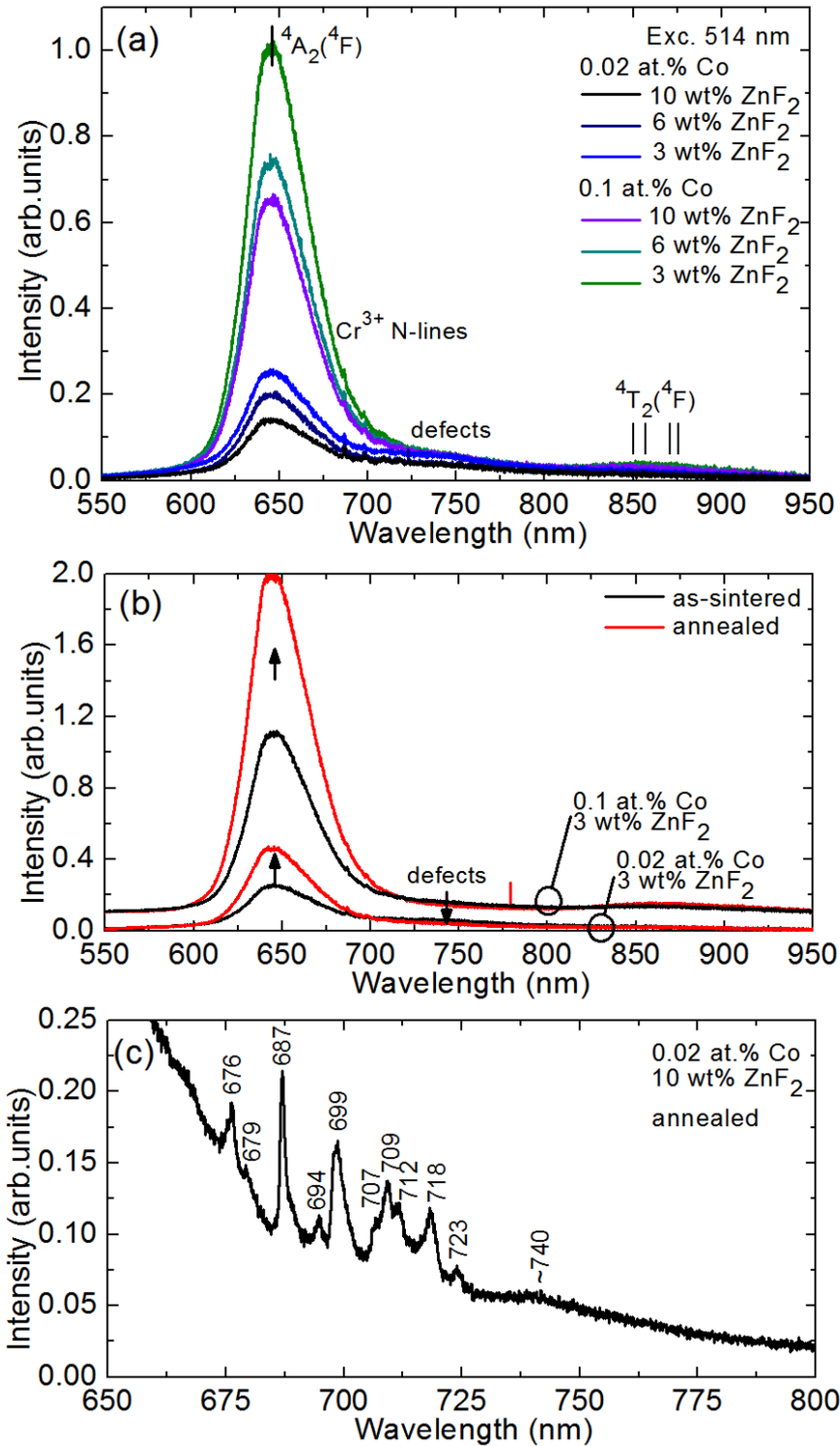


Co K α 1









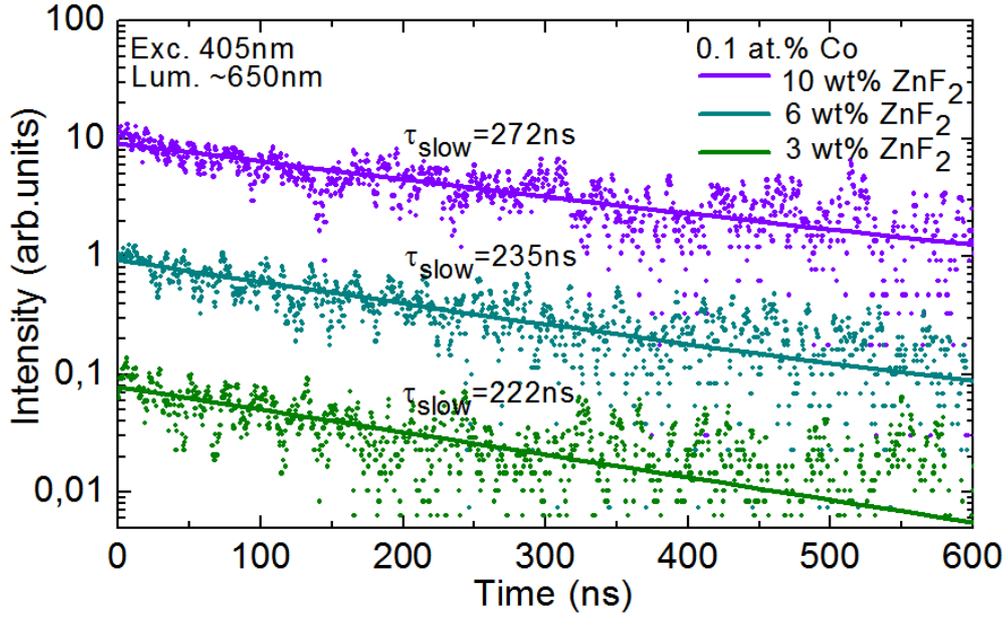


Table 1. Concentration of Co^{2+} ions in tetrahedral (T_d) sites in $\text{Co}:\text{ZnAl}_2\text{O}_4$ initial powders and transparent ceramics (determined from the spectroscopic data).

Material	ZnF_2 , wt%	Content of Co^{2+} in T_d sites, at.% (cm^{-3})	
Initial powders	3–10	0.02 (0.303×10^{19})	0.10 (1.514×10^{19})
Ceramics	3	0.011 (0.17×10^{19})	0.050 (0.76×10^{19})
	6	0.010 (0.15×10^{19})	0.052 (0.64×10^{19})
	10	0.007 (0.10×10^{19})	0.037 (0.56×10^{18})