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Color Origins in Langatate Crystals

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ABSTRACT

Langatate La5Ga5.5Ta0.5O14 is piezoelectric crystal from langasite family, commonly grown by Czochralski method from Ir crucible. Langatate crystals of different colors (colorless, orange, green) have been studied by optical spectroscopy in UV-Visible (200 – 800 nm) and IR (7000 – 1000 cm⁻¹) ranges. Furthermore, the effects of irradiation by ultraviolet laser source (λ=266 nm) and post-growth annealing in N₂+O₂ atmosphere have been investigated. The yellow-orange is mainly due to an absorption centered in the ultraviolet that extends into the blue of the visible spectrum (250-500 nm). The IR optical absorption spectra of Langatate crystals exhibit an absorption band at 5370 cm⁻¹. It seems linked to a point defect responsible for color. The intensity of the absorption band at 3430 cm⁻¹ increases after annealing in oxygen containing atmosphere. We have discussed phenomenon that can occur simultaneously in langatate crystals and produce very similar colors which are related to structural defects. First, metal ion impurities (as Iron, Titanium...), whose presence is previously confirmed by femtosecond laser ablation coupled with ICP-MS spectroscopy, can contribute to langatate color. Second, ultraviolet absorption leads us to think about charge transfer phenomenon such as O²⁻ → Fe³⁺ and/or Fe³⁺-Fe³⁺ pair transitions. Third, the irradiation by ultraviolet (λ=266 nm) laser source locally color the langatate sample by the creation of color centers. Origins of color centers, particularly those related to oxygen vacancies, (V₀⁻, 2e')⁺, are discussed. And, finally, point defects changes the band gap of langatate, leading to extend the absorption to visible light regions.

Keywords: annealing, color, color centers, charge transfer, impurities, langasite family crystals, optical spectrometry, point defects, ultraviolet irradiation.

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1. Introduction

Owing to its very interesting piezoelectric properties and thermal stability, langate La$_3$Ga$_5.5$Ta$_{0.5}$O$_{14}$ (LGT) crystal seems to be the best candidate to substitute quartz crystal for frequency output devices [1]. In fact, these applications require homogeneous crystals with reproducible physical properties. For that, before manufacturing acoustic devices (sensors, transducers, actuators or ultra-stable resonators), we control the quality of the crystal, using accurate techniques to reveal structural defects [2] [3]. Indeed, point defects, such as impurities and oxygen vacancies significantly affect the physicochemical properties (optical, electrical, piezoelectric…) and induce the crystal color.

Langate crystals (LGT), commonly grown by the Czochralski method from a ternary melt of La$_2$O$_3$-Ga$_2$O$_3$ and Ta$_2$O$_5$ using an iridium crucible, differ from colorless to yellow-orange or pale-green [4] [5] depending on growth and post-growth treatment conditions. LGT crystals without subsequent annealing differs from colorless to yellow-orange crystals, depending on growth atmosphere. LGT grown under Ar [6] [7] or N$_2$ [8] are colorless, while those grown under Ar + O$_2$ are orange [6] [9] [10] [11] and yellow [12] [7]. After vacuum annealing, the LGT samples become colorless and orange colored after air annealing [13] [4] [5]. The authors [7] [14] have suggested that the oxygen present in the growth and annealing atmospheres is responsible for the crystal color linked by the authors [11] [5] [4] to the presence of oxygen vacancies in the structure. According to Buzanov et al. [6], LGT crystals color depends not only on oxygen vacancies but also on cation vacancies and charge exchange in cations. In the opinion of the authors [15], LGT crystals contain iridium ions, which induce its color on the basis that LGT crystals grown from a Pt crucible in air were colorless [6]. However, Taishi et al. [14] obtained orange crystals by Bridgman growth in Pt crucible.

In this study, basing on chemical and spectroscopic analyses, we discuss point defects arising during growth and post-growth treatments and inducing LGT color. For our study, we have selected differently colored langate (LGT) samples from various suppliers.

2. Materials and Methods

2.1. LGT Samples

We have selected langate (LGT) crystals from different suppliers. Langasite La$_3$Ga$_5$SiO$_{14}$ (LGS) has been added for comparison. Details of growth, annealing conditions and color are tabulated in table 1. From the ingots, we cut wafers appropriate to each characterization technique. Moreover, some measurements have been achieved on ingot ‘CI6’ before annealing ‘ba’ and after annealing ‘aa’. The Figure 1 presents photos of samples cut from the ingots.

*Figure 1: Photos of langate (LGT) and langasite (LGS) samples cut from ingots: 1) CI6 ba, 2) CI6 aa, 3) Fn, 4) CK, 5) Fo, 6) M19, 7) LGS.*
Table 1: Growth and post-growth treatment conditions of LGT samples and LGS from different suppliers.

<table>
<thead>
<tr>
<th>Sample reference</th>
<th>Growth axis</th>
<th>Growth atmosphere</th>
<th>Annealing</th>
<th>Color</th>
</tr>
</thead>
<tbody>
<tr>
<td>CI6 ba</td>
<td>X</td>
<td>N₂</td>
<td>Before annealing</td>
<td>colorless</td>
</tr>
<tr>
<td>CI6 aa</td>
<td>X</td>
<td>N₂ + &lt;0.1% O₂</td>
<td>N₂ + &lt;0.1% O₂, 1400°C, 24 h</td>
<td>heterogeneous light orange</td>
</tr>
<tr>
<td>Fn</td>
<td>Z</td>
<td>Ar+1% O₂</td>
<td>Air, 1250 °C, several days</td>
<td>bright orange</td>
</tr>
<tr>
<td>CK</td>
<td>Z</td>
<td>N₂</td>
<td>unannealed</td>
<td>colorless</td>
</tr>
<tr>
<td>Fo</td>
<td>Z</td>
<td>unknown</td>
<td>unknown</td>
<td>green</td>
</tr>
<tr>
<td>M19</td>
<td>Z</td>
<td>N₂ +0.5% O₂</td>
<td>unannealed</td>
<td>light orange</td>
</tr>
<tr>
<td>LGS</td>
<td>Z</td>
<td>Ar+1% O₂</td>
<td>Air, 1250 °C, several days</td>
<td>bright orange</td>
</tr>
</tbody>
</table>

2.2. UV-Visible and IR Spectroscopy

Point defects significantly affect the optical properties and they can be revealed by optical spectroscopy. The optical spectra of LGT and LGS samples with plane and parallel polished surfaces (thickness 4 to 5 mm) were recorded in the wavelength range from 200 to 800 nm on a Lambda 900 UV-Visible spectrometer and in the range 7000 to 15000 cm⁻¹, on a PerkinElmer Frontier FT-IR spectrometer. Moreover, UV-Visible and IR spectra of LGT and LGS samples were recorded after irradiation with 266 nm picosecond laser from Teem Photonics and after annealing in oxygen containing atmosphere for LGT ‘CI6’ sample.

3. Results

3.1. UV-Visible Spectroscopy

The characteristic transmission spectra in UV-Visible ranged from 200 to 800 nm of LGT crystals from the different suppliers are shown in Figure 2.

Figure 2: UV-Vis transmission spectra of LGT and LGS samples.

The yellow-orange of LGT crystal is caused mainly by an absorption centered in the ultraviolet that extends into the blue of the visible spectrum (250-500 nm). The LGT CI6 aa spectrum exhibits a pronounced intrinsic absorption edge around 256 nm. For LGT CK, a shift of the intrinsic absorption edge to a higher wavelength of 270 nm is observed. The absorption band at 280 nm in the spectra of CK and CI6 was recorded by authors of [5] [16] and attributed to “La vacancies” V₃La⁺⁺. The weak and wide absorption band at around 330-340 nm observed in the transmission spectra of CK was attributed to oxygen vacancies V₃O⁻⁻.

An absorption band at λ ~490 nm is observed in LGS and LGT Fo and M19 spectra was previously attributed to F centers (V3O⁻⁻, 2é) and to orange coloration [16]. The defect responsible for the absorption band at 450 nm can be at the origin of the green color of LGT Fo. A spread and a shift to higher wavelength λ ~350-380 nm of the intrinsic absorption edge are observed in the transmission spectra of LGS sample, LGT Fn and Fo. Comparing LGS and LGT Fo spectra, before and after ultraviolet irradiation, we note that the absorption band at ~490 nm splits into two more intense absorption bands at ~480 and ~510 nm, which appear also, with lower intensity, in the spectrum of irradiated LGT CK. For LGT CI6, we note that annealing in oxygen containing atmosphere decreases the intensity of the absorption band at 280 nm.

3.2. IR Spectroscopy

The characteristic transmission spectra in IR range from 7000 to 15000 cm⁻¹ of LGT crystals from different suppliers are shown in Figure 3. IR absorption spectra of colored LGT (Fo, Fn, M19) and LGS samples exhibit an absorption peak at 5370 cm⁻¹, whose intensity increases with the color.
This sharp absorption peak seems linked to a defect responsible to the crystal coloration. The variation of the absorption band intensity is related to the variation of the defect density as shown in Figure 3. The intensity of the absorption band at 3430 cm\(^{-1}\) increases after annealing in oxygen containing atmosphere [17], Figure 3.

Comparing IR absorption spectra of LGT Fn and LGS, Figure 5, we note that LGT transmission edge is at higher wavelength and LGS spectrum exhibits an additional absorption band at ~2650 cm\(^{-1}\). For LGS, we note that ultraviolet irradiation increases the intensity of the absorption band at 5400 cm\(^{-1}\), Figure 5.

4. Discussion

4.1. Metal Ions Impurities

The yellow-orange color of LGT crystal can be due to light absorption by transition metals (Fe, Ti, Cu) and rare-earth elements (Ce, Pr, Nd) impurities under the effect of the electrical field of neighboring oxygen ions. According to chemical analyses performed by Femtosecond laser ablation/ICP-MS coupling (Fs LA-ICP-MS) on LGT ‘CI6 aa’ and LGT ‘Fn’[3], raw materials impurities, in particular metal transition ions such as Fe and Ti and lanthanide such as Ce and Pr, have contaminated grown LGT crystals and can contribute to their color. This is also confirmed by electron spin resonance (ESR) spectra of these LGT samples, which show a spin transition at g between 4.2 and 4.3, characteristic of Fe\(^{3+}\) in tetrahedral environment and a spin transition at g between 2.4-2.1 which is characteristic of Fe\(^{2+}\) in octahedral environment [3]. Moreover, Fs LA-ICP-MS chemical analyses [3] show that the concentrations of transition metal ions impurities increase with the color intensity.
of LGT crystals. LGT crystals are commonly grown by Czochralski method using an Iridium crucible [5] [6] [10] [15]. Kimura et al. [15] reported that LGT crystals contain iridium ions, which are responsible to the crystal color. However, F’s LA-ICP-MS chemical analyses [3] show that iridium concentration in LGT crystals is below 1 ppm. The as-grown LGT Cl6ba in N₂ atmosphere has been colorless and it became orange after annealing in oxygen containing atmosphere Figure 1. I.A. Kaurova et al. [13] found also that the color of LGT crystal grown by Czochralski technique under a 99% Ar + 1% O₂ atmosphere changed after air annealing from light yellow to orange. Annealing affects a color by modifying the valence state [18] of dispersed metal ions. For example, Fe^{2+} can be oxidized into Fe^{3+} by annealing LGT crystals in oxidizing conditions and then LGT color will change.

4.2. Charge Transfer

The optical transmission spectrum of bright orange LGT Fn sample, Figure 2, shows an absorption centered in the near ultraviolet and extended into the blue end of the visible spectrum. It can be due to an interaction between transition metal ions and its neighbors in the LGT structure. The ultraviolet absorption cannot simply be due to metal ions transition.

4.2.1. Oxygen-cation Charge Transfer

The yellow-orange color of LGT crystals can result from light absorbed through the transfer of electrons from the oxygen ions to the metal transition ions impurities such as O^{2-} ⇒ Fe^{3+}. E. Fritsch et al. [19] reported that oxygen-metal charge transfer absorptions produce yellow to orange and brown colors independently of the nature of the host mineral. For example, the yellow color of corundum Al₂O₃ [20] and quartz SiO₂ ‘citrine’ [21] is caused by light absorption through O^{2-} ⇒ Fe^{3+} charge transfer. Annealing in Oxygen containing atmosphere changes LGT Cl6ba from colorless to orange colored, Figure 1. First, heat treatment can induce or increase charge transfers. Second, annealing in oxygen containing atmosphere can oxidize transition metal ions impurities. As the positive charge of the central metal ion increases, the absorption band of the Oxygen ⇒ metal ion charge transfer moves from the ultraviolet toward the visible region [19] and then crystal changes from colorless to orange colored.

4.2.2. Cation-Cation Charge Transfer

The orange color of LGT crystal can be due to intervalence charge transfer between metal ions in the appropriate valence states. During annealing, metal ions impurities such as Fe^{2+} can be partially oxidized and, through diffusion, pairs of Fe^{2+} and Fe^{3+} ions can be formed, which interact through Fe^{2+}-Fe^{3+} intervalence charge transfer by light absorption in the blue end of the spectrum to yield the orange coloration. The yellow-orange color of LGT crystal can be due to light absorption by ion pair transitions such as Fe^{3+}-Fe^{3+}. These absorptions arise from a transition that occurs simultaneously in both ions. They are situated between 300 and 500 nm and generate yellow to red colors. Such transitions are very strongly oriented along the Fe^{3+}-O-Fe^{3+} bonds. As a result, they induce pleochroism [19]. Fe^{3+}-Fe^{3+} pair transitions are mentioned by J. Ferguson et al [22] to elucidate the color of some yellow sapphires α-Al₂O₃ and by S. M Mattson et al [23] to explain the hue of some red crystalline boron silicate called ‘dravites’.

4.3. Color Centers

4.3.1. Electron Color Centers

Electron color centers contribute to LGT crystal color. They can result from hypervalent impurities, and oxygen anions vacancies. According to Femtosecond laser/ICP-MS coupling chemical analysis [3], LGT crystals contain impurities that can occupy interstitial sites or substitute cations of the crystal lattice having or not the same degree of oxidation. If the impurity substitutes a cation with lower valence state such as Titanium cation Ti^{4+} substitutes Gallium cation Ga^{3+}, TiO₂, in order to keep electroneutrality, an available electron will be trapped nearby. This is an electron color center which can contribute to LGT crystal color. It was stated in [24] that the color of LGS crystals is caused by the presence of oxygen vacancies and their complexes in the form of F centers (V₀, 2 e⁻). The authors [4] [5] [11] examined the relationship between the coloration of LGT crystals and their composition by X-ray and neutron diffraction. They observed that LGT crystal color depends on the growth atmosphere and the post growth treatment, and it is
related to the concentration of oxygen vacancies complexes \( V_{O_2}^-(2e^-) \).

H. Kimura et al [15] and O.A Busanov et al [16] observed that the color of LGS and LGT crystals depends on the oxygen partial pressure in growth atmosphere. The color of LGT crystals decreases progressively in intensity when the Oxygen concentration in an Argon-Oxygen growth atmosphere is reduced [16]. Different researchers [4] [5] [9] [11] [13] studied the relationship between the color and the composition of LGT crystals using X-ray (XRD) and neutron diffraction (ND). They conclude that the color of such crystals depends on their oxygen vacancy content and that colored crystals have less oxygen vacancies. We can conclude that Oxygen content in growth atmosphere decreases oxygen vacancies and leads to colored crystals according to quasi-chemical reaction (1). I.A Kaurova et al. [4] stated that LGT crystals were colorless for \( V_{O_2}^- \) and exhibited a color for \( V_{O_2}^- \):

\[
\frac{1}{2} O_2 + V_{O_2}^- + 2e^- \leftrightarrow O_2^x
\]  

(1)

Annealing in oxygen containing atmosphere has changed LGT CI6ba from colorless to orange colored, Figure 1. Kaurova et al. [4] reported that air annealing increases the Oxygen content of the colorless sample and the crystal turned orange; vacuum annealing increases the oxygen vacancy concentration of the yellow sample, especially at annealing temperatures higher than 1200°C and the crystal becomes colorless [4]. G.M. Kuz'micheva et al. [5] reported that in oxygen-free growth atmosphere of langasite family, a dissociation of the melt and evaporation of Gallium suboxide are observed, which results in deficiency of Oxygen and Gallium. For that, LGT CI3 and CI6 have been grown in N\(_2\) atmosphere from Gallium rich charge composition to compensate the loss of gallium.

The site occupancy of the “octahedral” sites, equally occupied by the Ga\(^{3+}\) and Ta\(^{5+}\) cations in the stoichiometric composition La\(_3\)Ga\(_{5.5}\)Ta\(_{0.5}\)O\(_{14}\) can be written as Ga:Ta = 1-x : x, where 0 < x < 1. To maintain electroneutrality, cation and anion vacancies will form.

If 0 < x < 0.5 Ga\(^{3+}\) (Gallium rich melt composition), the crystal will be negatively charged and electroneutrality will be ensured by oxygen vacancies according to the quasi chemical reactions (2).

\[
Ga_2O_5 \rightarrow 2Ga_3'' + 2V_0 + 3O_2
\]

(2)

In this case, the composition of LGT crystal can be written as \( La_3(Ga_{1-x}Ta_x)Ga_5O_{14-(0.5-x)} \), where 0 < x < 0.5

### 4.3.2. Hole Color Centers

If an electron is missing from a location where usually an electron pair exists, a "hole" color center is created. If LGT crystal impurities [3] substitute cations of the crystal lattice with higher valence state such as Fe\(_{Ti}^{3+}\), Fe\(_{Ti}^{3+}\) substitutes Tantalum cation Ta\(^{5+}\), in order to maintain the electroneutrality, monovalent cations (Na\(^+\), K\(^+\), H\(^+\)) will be present nearby. This causes forces on the electrons of the oxygen atom near the Aluminium ion to be weakened and an extreme heat can remove one of the weaker bonded electrons of the oxygen atoms that can be trapped by the monovalent cation. This leaves a hole and different energy levels become available to the new unpaired electron on the oxygen ion.

### 4.4. Interband Transitions

The color of LGT can arise through its electronic structure (band theory). In this case, transitions between bands rather than between energy levels of single atoms are responsible for the color. In fact, crystal defects such as impurities, atom vacancies and interstitial atoms contribute to the band gap narrowing by introducing new energy levels. To predict how the defects altered the electronic properties of the host materials, Chan-Yeup Chung et al. [25] performed DFT calculations of band gap energy of perfect and defect-containing LGT crystals with the hybrid functional HSE06. The calculated band gap of defect-free LGT was 5.279 eV and the calculated band gap corresponding to each defective system were 4.849 eV, 4.399 eV, 4.027 eV, 4.050 eV, 4.348 eV, and 3.678 eV for \( Ga_{Ta}^{3+},V_{Ga},V_{Ta}^{3+},V_{Ga}(1a,oct),V_{Ga}(3f,tet) \), and \( V_{Ga(2d,tet)} \), respectively.

We can qualitatively determine the band gap energy of LGT crystal from the UV-Visible transmission spectrum. However, this is only the case if the interband transition is the dominant mechanism of absorption. The fundamental absorption edge of LGT crystal is 242 nm [27]. A spread and a shift to higher wavelengths of the intrinsic absorption edges of LGT crystals under study are shown in Figure 2.
5. Conclusions

La₃Ga₅.₅Ta₀.₅O₁₄ crystals (LGT) differ from colorless to yellow-orange or pale-green depending on the feedstock, the growth and the post-growth treatment parameters and on trapped defects during the synthesis. The yellow-orange color is caused mainly by an absorption centered in the ultraviolet that extends into the blue of the visible spectrum (250-500 nm) and absorbs violet and blue. After ultraviolet irradiation (λ=266 nm), we note that the absorption band at ~490 nm splits into two absorption bands at ~480 and ~510 nm. We have discussed phenomenon that can occur simultaneously in langatate crystals and produce very similar colors. First, according to fs-laser ablation ICP-MS coupling chemical analyses, LGT crystals contain metal transition impurities coming from raw materials, such as Iron and Titanium, which absorb visible light and contribute to the color. Second, the ultraviolet absorption leads us to think about charge transfer from oxygen ions to the metal transitions ions such as O²⁻ → Fe³⁺ and ion pair transitions such as Fe³⁺-Fe³⁺. Third, we have observed that ultraviolet irradiation induces the orange color by the creation of color centers. We have discussed different forms of color centers that can occur in the LGT structure such as (V₀⁺, 2e⁻)³ and we have presented its origins. Oxygen vacancies, V₀⁺, are linked to the oxygen content in growth and annealing atmospheres. Finally, point defects reduce the band gap energy to the visible light range and LGT color can be caused by transitions between bands.

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