Importance of Long-Range Channel Sr Displacements for the Narrow Emission in Sr[Li$_2$Al$_2$O$_2$N$_2$]:Eu$^{2+}$ Phosphor

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The recently discovered Sr[Li$_2$Al$_2$O$_2$N$_2$]:Eu$^{2+}$ red phosphor, candidate for the next generation of eco-efficient white light-emitting diodes, exhibits excellent emission spectral position and exceptionally small linewidth. It belongs to the UCr$_4$C$_4$-structure family of phosphors containing many potential candidates for commercial phosphors, whose small linewidth, tentatively ascribed to the high-symmetry cuboid environment of the doping site, has drawn the attention of researchers in the last five years. Density functional theory, ΔSCF method, and configuration coordinate models (CCM) are used to provide a complete characterization of this material. Using a multi-dimensional CCM, an accurate description of the coupling of the vibronic structure with the electronic transition is obtained, including the partial Huang–Rhys factors and frequency of the dominant modes. It is shown that, in addition to the first-coordination shell cuboid deformation mode, low-frequency phonon modes involving chains of strontium atoms along the tetragonal axis shape the emission linewidth in Sr[Li$_2$Al$_2$O$_2$N$_2$]:Eu$^{2+}$. This finding sheds new light on the emission properties of UCr$_4$C$_4$-structure phosphors, possessing similar Ca/Sr/Ba channel. The approach provides a robust theoretical framework to systematically study the emission spectra of such Eu-doped phosphors, and predict candidates with expected similar or even sharper linewidth.

1. Introduction

To mitigate CO$_2$ emission, the use of efficient white-light sources for general lighting is becoming increasingly important worldwide. In 2019, in its solid state lighting plan that relies heavily on light-emitting diodes (LEDs) technology, the US department of energy estimated that the energy savings could account for about a 5% reduction in the total primary energy budget of the USA.[1] One way to realize such white light is to coat a primary blue or near ultraviolet LED with one or multiple phosphor material(s) that converts a fraction of the initial high-frequency photons to lower frequency, the resulting light appearing white. There generally exist two ways to obtain white light via LEDs with phosphors: i) a blue LED is used to excite a single yellow phosphor or mixed green and red phosphors; and ii) a near ultraviolet LED chip is used to excite the red, green, and blue phosphors.[2,3]

Among the various phosphor compounds, finding a highly efficient red emitting phosphor is crucial for a further increase in the luminous efficacy while keeping a high color-rendering index. In order to sufficiently cover the red spectral region while minimizing the efficacy loss caused by the eye poor long-wave sensitivity, red phosphors with narrow emission band showing a peak in the optimal red region are needed.[4,5] In 2019, Hoerder et al.[6] reported the experimental discovery of a new narrow-band red-emission phosphor Sr[Li$_2$Al$_2$O$_2$N$_2$]:Eu$^{2+}$ (SALON) that fulfills such requirements with an emission peak located at 614 nm, a full width at half maximum (FWHM) of only 48 nm at room temperature and a low thermal quenching of 4% at operating temperature (420 K). The fact that the emission peak is located at a shorter wavelength than its cousin Sr[LiAl$_3$N$_4$]:Eu$^{2+}$ (SLA)[7] enables a gain of 16% in luminous efficacy compared to SLA, while keeping an excellent color-rendering index and thus enabling a potential leap in the energy efficiency of white-emitting phosphor-converted LEDs. SALON belongs to the UCr$_4$C$_4$-structure family of phosphors, from which most interesting narrow-emission candidates have emerged recently.[8,9]

In our work, we first provide a theoretical characterization of the geometric, electronic and vibrational properties of SALON from first principles, using density-functional theory (DFT). We study the narrow Eu$_{5d}$→Eu$_{4f}$ emission spectrum from a theoretical perspective. For this purpose, we use the ΔSCF method,[10–13] that allows one to compute relevant transition energies and atomic configuration displacement induced by the 5d→4f electronic transition. In combination with ΔSCF calculations,
we use a multi-dimensional configuration-coordinate model with the help of a generating function approach to determine the luminescence intensity spectrum. Indeed, this approach allows us to extract the spectral decomposition of the phonon modes participating to the 5d–4f transition by projecting the atomic displacements induced by the 5d–4f transition onto the phonon eigenvectors of the host structure. For this purpose, we use density-functional perturbation theory (DFPT)[14,15] to compute the phonons modes of SALON. This approach has been used by Alkauskas et al.[16] to study the vibronic structure of diamond nitrogen-vacancy (NV) center and, recently, by Linderälv et al.[17] to study the lineshape of Ce-doped YAG. To our knowledge, the latter work is the only prior existing fully first-principles vibronic analysis of the 5d–4f emission of lanthanide-based phosphors. We note however that some previous vibronic studies, mixing experimental and theoretical analysis, were already carried out.[18–21]

We compare our result with a 1D configuration-coordinate model (1D CCM).

It was suggested by Hoerder et al.[6] that the narrow emission band of SALON originates from the Eu$^{2+}$ optical center with high local-symmetry environment composed of a nearly cubic polyhedron of four oxygen and four nitrogen atoms, resulting in an isotropic structural relaxation upon emission. This would reduce the number of different energetic states involved in the emission process, leading to a narrow emission band.[6] Here, we show that this picture is incomplete.

Our computations indicate that the structural change of the environment of the europium atom when going from excited to ground state is composed of two patterns: i) a nearly isotropic expansion of the first coordination shell of the Eu atom composed of O and N, as suggested by Hoerder et al.;[6] ii) a long-range collective displacement of the Sr channel containing Eu atom away from Eu. This second pattern couples with low-frequency acoustic and $B_n$ phonon modes in a small energy window, and contributes to the small linewidth. It is also responsible for the slow convergence of the results with the supercell size. We propose a geometric series model on the Sr displacements to estimate the residual error.

Overall, we explain the narrow emission spectrum by the combination of i) a small atomic displacement upon emission due to the rigid host structure and ii) an important participation of low-frequency phonon modes in the electronic transition. We finally find that the shape of our theoretical spectrum agrees well with the experimental one.

The paper is structured as follows. Section 2 is devoted to theoretical background: we explain how transition energies and atomic displacements induced by the 5d-4f transition can be used within a configuration-coordinate model framework. The generating function approach that allows one to predict luminescence spectra is also introduced. In Section 3, we present the results of our work. We start with the DFT results on structural and electronic properties of SALON and present its luminescent properties. We describe which phonon modes dominate the electronic properties of SALON and present its luminescent properties. Deviations from linear interpolation can be considered when studying non-radiative decay channels, but have been shown to have a small impact on the luminescence spectrum shape.[22]

In the 1D-CCM, the 4f and 5d Born–Oppenheimer potential energies are displayed as a function of the normal coordinate $Q$. The corresponding ionic configuration is $\Lambda_g^s$ with configuration coordinate $Q_{fg}$. When a photon is absorbed, the system is quickly promoted to an unrelaxed excited 5d state $\Lambda_g^e$ without atomic motion. Due to this change of electronic state, atomic relaxation towards a new equilibrium configuration $Q_e$ takes place with phonon emission. The energy lost in the process is the Franck–Condon shift $E_{FC,e}$. Photon emission occurs then through 5d-4f deexcitation and the system reaches an unrelaxed ground state $\Lambda_g$. The cycle is completed by a last relaxation with phonon emission and energy loss $E_{FC,g}$ until the initial state $\Lambda_g^s$ is reached.

The total energy loss is the Stokes shift $\Delta S$ given by the sum of the two Franck–Condon shifts $\Delta S = E_{FC,e} + E_{FC,g}$. The zero-phonon line refers to a purely electronic transition with energy $E_{ZPL}$. Ignoring the small difference of zero-point energies between the two states, $E_{ZPL}$ is the difference between the two energy surface minima.

We first describe how to consider the luminescent properties with a 1D-CCM as depicted in Figure 1. In this model, all the complexity of the nuclei vibrations is reduced to one effective vibrational mode characterized by a generalized configuration coordinate $Q$ that interpolates linearly between ground- and excited-state atomic coordinates. Deviations from linear interpolation can be considered when studying non-radiative decay channels, but have been shown to have a small impact on the luminescence spectrum shape.[22]

In the 1D-CCM, the 4f and 5d Born–Oppenheimer potential energies are displayed as a function of the normal coordinate $Q$. The corresponding ionic configuration is

$$\frac{1}{\Delta Q} R(Q) = \frac{Q}{\Delta Q} (R_e - R_g) + R_g$$

where $\Delta Q$ is the normal coordinate change from ground-state to excited-state geometries. It writes

$$(\Delta Q)^2 = \sum_{\alpha} m_{\alpha} (R_{\alpha,e} - R_{\alpha,g})^2$$

Figure 1. Configuration coordinate diagram, see text for details.
where $i$ labels Cartesian axes, $\alpha$ atoms, $m_{\alpha}$ atomic masses, $R_{\alpha}$ and $R_{\alpha}$ are respectively atomic positions in 5d excited and 4f ground states. Within such coordinate system, the harmonic ground-state and excited-state energies are respectively

$$E(Q) = \frac{1}{2} \Omega_g^2 Q^2$$  \hspace{1cm} (3)$$

$$E'(Q) = \frac{1}{2} \Omega_x^2 (Q - \Delta Q)^2 + E_{PL}$$  \hspace{1cm} (4)$$

The associated effective vibrational frequencies in the ground (g) or excited (e) states are

$$\Omega_{[g,e]}^2 = \frac{2E_{FC,[g,e]}}{\Delta Q^2}$$  \hspace{1cm} (5)$$

We can use Equations (3) and (4) to compute the Huang–Rhys factors that indicate the average numbers of phonons emitted during the relaxation process:

$$S_{[g,e]} = \frac{E_{FC,[g,e]}}{\hbar \Omega_{[g,e]}} = \frac{\Omega_{[g,e]}^2 \Delta Q^2}{2\hbar}$$  \hspace{1cm} (6)$$

Using a semi-classical treatment, one can estimate the FWHM of the spectrum by considering the expectation value of $Q$ and by computing the density of transition as a function of the emitted energy. At 0 K, the FWHM, $W(0)$, is written as

$$W(0) = \sqrt{8 \ln 2} \frac{S_g}{\sqrt{S_e}} \hbar \Omega_g$$  \hspace{1cm} (7)$$

This formula is valid for large values of $S$, such that the otherwise Pekarian line shape of the emission spectrum is well approximated by a Gaussian envelope. As will be seen later, this condition is not well fulfilled for SALON phosphor ($S = 3$), but we will use it nonetheless as a reference for later comparison.

Focusing now on the multi-dimensional configuration coordinate model (multi-D CCM), the nuclear motions are a superposition of 3N normal modes of vibration $\nu$ represented by normal coordinates $Q_\nu$ and frequency $\omega_\nu$ with $N$ the number of atoms in the supercell. The harmonic approximation is retained: phonon modes are independent of each other and the vibrations in the solids are composed of 3N harmonic oscillators. The vibrational state $\chi_\nu$ is expressed as a product of 3N harmonic oscillator eigenfunctions $\chi_{\nu m}$ with $n_\nu$ the vibrational state of the $\nu$th harmonic oscillator and $n$ the set of 3N vibrational states $\{n_1, n_2, \ldots, n_N\}$.

Making then the Franck–Condon approximation, which states that the transition dipole moment between excited and ground state depends weakly on nuclear coordinates, we write the normalized luminescence intensity at 0 K for a given photon energy $h\omega$ as $I(h\omega) = C \omega A(h\omega)$, with $C$ a normalization constant that is chosen such that the area under the curve is unity, and the emission spectral function is

$$A(h\omega) = \sum_{\nu} \left| \langle \chi_{\nu m} | \chi_{\nu n} \rangle \right|^2 \delta(E_{PL} + E_{g,n} - h\omega)$$  \hspace{1cm} (8)$$

with $E_{g,n} = \sum n_\nu \hbar \omega_\nu$, the energy of the state $\chi_{\nu n}$.\hspace{0.5cm}[16]$\!

Note that, in the 1D-CCM and assuming the same harmonic curvature in the ground and excited states, $\Omega_g = \Omega_e$, such that $S_g = S_e = S$. Equation (8) reduces to

$$A(h\omega) = \sum_{\nu} e^{-S_\nu} \frac{S_\nu}{n_\nu} \delta(E_{PL} + n\hbar\Omega - h\omega)$$  \hspace{1cm} (9)$$

In the multi-D CCM approach, the harmonic approximation allows one to write the 3N-dimensional Franck–Condon overlap as a product of 3N overlap integrals $\langle \chi_{\nu m} | \chi_{\nu m} \rangle = \prod_{\nu} \langle \chi_{\nu m} | \chi_{\nu m} \rangle$. Indeed at 0 K all initial vibrational states are in their lowest state $n = 0$. We finally suppose that the Born–Oppenheimer curvatures are the same in the 4f and 5d state which holds for SALON. This allows one to further simplify Equation (8) since

$$\left| \langle \chi_{g,n} | \chi_{e,0} \rangle \right|^2 = e^{-S_\nu} \frac{S_\nu}{n_\nu} \delta(S_\nu, n_\nu)$$  \hspace{1cm} (10)$$

Quantities that enter Equations (8) and (10) are phonon frequencies $\omega_\nu$ and partial Huang–Rhys factors $S_\nu$. Following the work of Alkauskas et al.,\hspace{0.5cm}[16] we define the weight by which each mode $\nu$ contributes to the atomic position changes when a electronic transition occurs as $p_\nu = (\Delta Q_j/\Delta Q^2)$, where

$$\Delta Q_j = \sum_{\nu,i} m_{\nu,i}^2 (R_{\nu,ai} - R_{\nu,ai}) \Delta r_{\nu,ai}$$  \hspace{1cm} (11)$$

$$\Delta Q^2 = \sum_j (\Delta Q_j)^2$$  \hspace{1cm} (12)$$

and $\Delta r_{\nu,ai}$ is the vector that represents the displacement of atom $a$ in phonon mode $\nu$. This vector is normalized such that $\sum_j \Delta r_{\nu,ai} \Delta r_{\nu,ai} = \delta_{\nu,j}$. In other words, the contribution of a mode to the electronic transition is given by the projection of the phonon eigenvector associated with this mode on the atomic distortion induced by this electronic transition, weighted by the atom mass. It is possible to define an effective frequency as

$$\Omega_{\nu}^2 = \sum_{\nu} p_\nu \omega_\nu^2$$  \hspace{1cm} (13)$$

where $\omega_\nu$ is the frequency of the mode $\nu$. The partial Huang–Rhys factor associated with the mode $\nu$ indicates the number of phonons of mode $\nu$ involved in the 4f–5d transition. Assuming harmonicity, we have

$$S_\nu = \frac{1}{2} \frac{\omega_\nu^2 \Delta Q_j^2}{\hbar \omega_\nu} = \omega_\nu \Delta Q_j^2$$  \hspace{1cm} (14)$$

By introducing a Huang–Rhys spectral decomposition,\hspace{0.5cm}[24]

$$S(h\omega) = \sum_{\nu} S_\nu \delta(h\omega - \hbar\omega_\nu)$$  \hspace{1cm} (15)$$
a generating function approach yields the spectral function $A(\hbar \omega)$

$$A(E_{ZPL} - \hbar \omega) = \frac{1}{2\pi \hbar} \int G(t)e^{i\omega t}dt$$

(16)

where the generating function $G(t)$ is

$$G(t) = e^{(0)-S(0)}$$

(17)

$$S(t) = \int_0^\infty S(\hbar \omega)e^{i\omega t}d(\hbar \omega)$$

(18)

The $S(0) = \sum \nu S_{\nu}$ is the total Huang–Rhys factor and the parameter $\gamma$ represents the homogeneous Lorentzian broadening of each transition. One can also model inhomogeneous broadening due to ensemble averaging with a convolution of $A(\hbar \omega)$ with a Gaussian. Within this multi-dimensional approach, calculating the emission spectrum requires to compute in DFT: i) the zero-phonon line energy $E_{ZPL}$, ii) the atomic configuration change induced by 4f-5d transition ($R_{Crs}$, $R_{Sr}$), iii) the phonon eigenvectors $\Delta_{\nu \alpha}$ and eigenfrequencies $\alpha_{\nu}$. The first two are accessible through the ASCF method while DFPT can deliver the third one. Within the 1D-CCM, computing all phonon modes with DFPT is not needed, only one relevant effective phonon frequency is deduced from the harmonic approximation, the knowledge of the Franck–Condon shift and the normal coordinate change $\Delta Q$.

3. Results and Discussion

3.1. Structural Properties

Based on single-crystal X-ray diffraction data, Hoerder et al. showed that SALON crystallizes in a tetragonal phase (space group $P4_2/m$) with the unit-cell parameter $a = b = 7.959 \text{ Å}$ and $c = 3.184 \text{ Å}$, with 2 formula units per cell and 18 atoms per primitive cell. The structure is a variant of the UC$_4$C$_4$ structure type, well known in the field of phosphor materials to provide a highly condensed and rigid framework structure, with strontium on the corresponding uranium site, aluminium and lithium on the corresponding chromium site, and nitrogen and oxygen on the corresponding carbide site. Two tetrahedra types form a network hosting the strontium cations in one of the resulting channels shown in Figure 2. Strontium atoms are coordinated with four nitrogen and four oxygen atoms which gives a highly symmetrical cube-like coordination.

The structural parameters of undoped SALON computed within DFT are listed in Table 1. We observe globally a slight overestimation of the structural parameters by about 0.5–0.7% which is common for GGA-PBE functionals. Since our computation neglects all vibrational and anharmonic effects, while the experimental structure is measured at room temperature, this error might be underestimated because of zero-point and thermal expansion effects. When a Sr atom is replaced by an Eu atom in the doped configuration, with decrease of symmetry to point group $2/m (C_{2h})$, we observe that the Eu–O and Eu–N bond lengths are quite close to the Sr–O and Sr–N bond lengths, which is reasonable since the experimental atomic radii of Eu and Sr in their VIII coordination state are very close; 1.39 and 1.40 Å, respectively. In contrast, we also observe that the Eu–Sr length is more than 2% bigger than the Sr–Sr length in the undoped structure.

To understand the relaxation process accompanying the change in electronic density induced by the 5d–4f transition, a comparison is performed between the 4f-state geometry ($\Lambda_{\nu}$, $\Lambda_{\nu}^*$) and the 5d-state geometry ($\Lambda^*$, $\Lambda$), see Figure 3. On the left and right of Figure 3a, we present the highest occupied Kohn-Sham orbitals (square moduli of the wavefunctions) associated to the $4f^{7}(S_{7/2})$ ground and $4f^{6}(F_{7/2})5d^{1}$ excited states. We

Figure 2. 2x2x2 supercell of undoped SALON. Tetrahedra T1 are composed of aluminium coordinated with three nitrogen atoms and one oxygen atom forming a $[\text{AlN}_3]^8^-$ unit. Tetrahedra T2 are composed of lithium coordinated with one nitrogen atom and three oxygen atoms forming a $[\text{LiO}_3N]^8^-$ unit. These tetrahedra compose a network of vierer rings arranged in three types of channels of which two are empty (C1 and C2) and the third hosts strontium cations that are coordinated with four oxygen atoms and four nitrogen atoms in a nearly cubic environment. The unit cell is represented with dashed lines.

Figure 3. Computed (left) and experimental (right) geometries of undoped SALON. a) AlN$_3$ unit in T1, SrN$_3$ unit in T2, and AlO$_3$ unit in C3. b) SrO$_3$N$_3$ unit in T1, LiO$_3$N$_3$ unit in T2, and AlN$_3$ unit in C3.
This work could also argue that the excited $\text{Eu}^{2+}$ configuration, leading to stronger Eu–O and Eu–N bonds, due to both Coulomb and covalent strengthening. This uncommon situation of a bond length shortening upon $f \rightarrow d$ excitation has been already observed experimentally and studied theoretically in various other lanthanide compounds.[28,29,33]

Second, we observe a long-range relaxation of the strontium atoms belonging to the strontium channel containing the europium atom (see green shaded area of Figure 3a). When going from 5d to 4f state, those strontium atoms are pushed away from europium along c-axis. Comparing first the displacement magnitude of the two patterns, we observe that the displacements of the first distant Sr is slightly smaller than the O/N displacement associated to the first pattern (see Figure S10, Supporting Information for a visualization of the displacement norm as a function of the distance from europium). However, what matters in the luminescent properties is the normal coordinate change ($\Delta Q^2$), which explains why this second pattern is actually the dominant one, the Sr being heavier. It appears that about half of the total normal coordinate change comes from Sr displacements, see Figure 3c.

It is surprising to observe that the displacement of the first distant Sr is slightly smaller than the second distant Sr. We attribute this effect to the large spatial extent of the 5d$_z$ orbital. Beyond the second distant Sr, the Sr displacements follow an exponential decay, as explained later.

We performed a convergence study of ($\Delta Q^2$) as a function of the supercell size (see Figure 3c and Figure S1, Supporting Information). Increasing the number of atoms in the $x$–$y$ plane ($\sqrt{2} \times \sqrt{2} \times N$ supercells) does not affect the result while increasing the Eu–Eu distance along c-axis does. Indeed, for too small $1 \times 1 \times N$ supercells, there is a spurious cancellation of Sr displacements caused by periodically repeated Eu. The c-component of the supercell size (or equivalently the Eu-Eu distance) required to obtain a strictly converged ($\Delta Q^2$) is very large, and instead, we model this cancellation effect by expressing the Sr displacements $D[i]$ in [Å] unit as a sum of two counter-acting decreasing geometric series,

$$D[i] = C(A^{i-2} - A^{i(N_{\text{Sr}} - 1 - i)})$$

(19)

where $i$ labels the Sr atoms, $N_{\text{Sr}}$ denotes the number of Sr atoms between two adjacent Eu, which is 11 in the case of the $1 \times 1 \times 12$ supercell used, and where $C$ and $A$ are the fitting coefficients. With $A = 0.58$ [Å] and $C = 0.037$ [Å], the DFT displacements are well reproduced with this simple model (see red curve of Figure 3b). In this figure, plain and light green curves refer respectively to the first and second terms of Equation (19). Note that we excluded the first Sr because of the above-mentioned reasons. If we now consider a perfectly isolated Eu, the Sr displacements would be expressed as a simple decreasing geometric series with the same fitting coefficients $A$ and $C$ (plain green curve). We

| Structural parameters of undoped and doped SALON, either in ground 4f or excited 5d state for a $1 \times 1 \times 12$ supercell. |
|-------------------------------------------------------------|
| **Experiment[6]**                          | **Undoped** | **Doped 4f** | **Doped 5d** |
| $a$ [Å]                                      | 7.959       | 8.013 (+0.67%) | 8.012 (+0.01%) |
| $b$ [Å]                                      | 7.959       | 8.013 (+0.67%) | 8.012 (+0.01%) |
| $c$ [Å]                                      | 38.208      | 38.438 (+0.60%) | 38.442 (+0.01%) |
| X–Sr length [Å]                              | 3.184       | 3.131 (-1.66%) | 3.206 (+2.39%) |
| X–O length [Å]                               | 2.659       | 2.667 (+0.30%) | 2.678 (+0.41%) |
| X–N length [Å]                               | 2.760       | 2.782 (+0.79%) | 2.738 (+1.68%) |
| O–X–O angle [°]                              | 73.572      | 73.819 (+0.34%) | 73.719 (-1.13%) |
| N–X–N angle [°]                              | 70.446      | 70.301 (-0.21%) | 70.356 (+0.08%) |
| Polyhedral volume [Å$^3$]                     | 30.411      | 30.870 (+1.49%) | 31.070 (+0.76%) |

The $X$ symbol refers to strontium when undoped and europium when doped. Polyhedral volume refers to the nearly cubic environment of Sr/Eu composed of 4 oxygen and 4 nitrogen. The percentages in parentheses are calculated with respect to the previous column.
compute the error made as the difference between the red and
green displacements. In term of normal coordinate change,
this gives a difference of 0.028 [amu⋅Å²], a 4.8% error with
respect to the Sr contribution to the total \((ΔQ)^2\). Indeed, these
additional Sr displacements lead to small additional displace-
ments of their atomic neighbours that we simply model with
this same percentage error. In total, we estimate the strictly
converged value (i.e., without finite supercell size effect) to be
\((ΔQ)^2\) ≈ 1.33 [amu⋅Å²], 4.7% higher than the computed DFT
value. The impact on the luminescent properties of a specific
channel structure along one crystallographic direction is a
rarely observed effect. We still note some works on the lumi-
nescence of specific compounds with this 1D feature. To
our knowledge however, it is the first time that such long-range
channel relaxation is theoretically highlighted and analysed
in phosphor materials. As it will be seen in Section 3.3, this
greatly impacts the computed luminescent properties and it
might be a common denominator that explains the particular
luminescent properties of this type of phosphor materials.

3.2. Electronic Properties

The electronic band structure and the partial density of states
near the Fermi level of undoped SALON are presented in
Figure 4a, obtained with the relaxed cell parameters. For the
electronic structure over the entire energy range, see Figure S11,
Supporting Information.

The computed indirect band gap of undoped SALON is
3.44 eV, underestimating the experimental gap of 4.4 eV,[8] a
quite standard underestimation, often observed with local or
(semi-)local exchange-correlation functionals, like the pres-
ently used GGA-PBE. Figure 4b shows the band structures of
the four characteristic states of the 1D coordinate model (see
Figure 1) for a 1×1×5 supercell size. We find that in contrast
to the emission spectrum, the electronic band structure con-
verges faster with the supercell size. In the ground state, the
seven 4f states are correctly placed inside the band gap. In the
excited state (\(Δ_{\nu}^2\) or \(Δ_{\varepsilon}^2\)), only one 4f state is left in the band gap.
This change is a typical feature of the DFT+U approach. Then,
we observe one first 5d state, presenting mainly a 5d\textsubscript{xy} character, actually a superposition of \(l = 2, m = -2, 0 \) and 2 angular momentum components, belonging to the \( a_g \) irreducible representation of the above-mentioned 2/m point group. It shows a small dispersion along the c-axis coming from a residual interaction between periodically repeated europium atoms. See Figure S2, Supporting Information for related computed properties. The next higher band lying in the band gap is identified as mainly 5d\textsubscript{z}. When in the excited state, the 4f-like hole state can also be analyzed in terms of its angular momentum components, showing a superposition of \( l = 3, m = -3, -1, 1 \) and 3 components, so belonging to the \( b_g \) irreducible representation of the above-mentioned 2/m point group.

### 3.3. Luminescent Properties

We compute the luminescence properties of doped SALON and report the DFT parameters required for the 1D model in Table 2. A comparison with the similar compound Sr[LiAl\textsubscript{3}N\textsubscript{4}]:Eu\textsuperscript{2+} (SLA) is provided using data from ref. [13], where a similar computational methodology was used. The experimental \( E_{ZPL} \) is difficult to estimate precisely as no vibronic structure is present in the 15K-spectrum.\(^6\) We place it in the range 2.03–2.08 eV. Our computed zero-phonon line energy \( E_{ZPL} = 2.032 \) eV falls within this range. The convergence of \( E_{ZPL} \) as a function of the supercell size can be found in Figure S3, Supporting Information and shows that 0.1% convergence is achieved with respect to the 1\times1\times8 supercell.

Then, comparing with the 15 phosphors studied in ref. [13], we notice that SALON exhibits a small Stokes shift \( \Delta S \) and a small \( \Delta Q \). This indicates that the geometry of the system is not much affected by the 5d–4f transition. We observe also that the effective vibrational energies in the ground (25.15 meV) and excited state (23.86 meV) are close to each other, meaning that the curvature of the Born–Oppenheimer potential in the ground and excited state are quite similar, in contrast to SLA (see Table 2).

We also note that the Huang–Rhys factor and phonon frequency are consistent with the Stokes shift. As explained in reference\(^{34} \) \( \Delta S = 0.182 \) eV should lie between \((2S_g - 1)\hbar\Omega_g = 0.167 \) eV and \( 2S_e\hbar\Omega_e = 0.192 \) eV, which is the case. Finally, the semi-classical FWHM = 119 meV obtained from Equation (7) is quite close to the experimental FWHM = 135 meV at low temperature.

We now present the results from the multi-dimensional approach described in Section 2. The discrete partial Huang–Rhys factors \( S_v \) are shown in Figure 5a along with the spectral decomposition \( S(\hbar\Omega) \) (Gaussian smearing of 4 meV). The phonon density of states is displayed in the background of this figure. See Supporting Information for related computed properties (phonon band structure, phonon frequencies at \( \Gamma \), Born effective charges, and dielectric tensor).

This reveals the nature of the electron-phonon coupling associated to the 5d–4f transition: At low energy (<0.022 eV), phonon modes associated to Sr atoms dominates. This threshold is fixed such that, at 0.022 eV, Sr atoms contribute to half of the phonon density of states. It is also where a dip is observed in the spectral function. We highlight in this low-energy region the four modes presenting the highest partial Huang–Rhys factors \( S_v \). We present them in the phonon dispersion curve along \( \Gamma-Z \) in Figure 5b and we show their eigendisplacements in Figure 5c where the displacements have been scaled by a factor 10 for clarity. The mode that couples the

| Compound | \( E_{ZPL} \) | \( E_{IC, g} \) | \( E_{IC, e} \) | \( \Delta S \) |
|---------|-------------|------------|-------------|----------|
| SALON:Eu | 2.032 | 0.096 | 0.086 | 0.182 |
| SLA:Eu 1 | 2.038 | 0.076 | 0.057 | 0.133 |
| SLA:Eu 2 | 2.049 | 0.060 | 0.111 | 0.171 |
| SALON:Eu | 1.127 | 25.15 | 23.86 | 3.82 | 3.62 |
| SLA:Eu 1 | 0.756 | 33.40 | 29.0 | 2.63 | 1.71 |
| SLA:Eu 2 | 1.222 | 18.30 | 25.0 | 3.27 | 4.46 |

\[ 2.032 \text{ eV} \] falls within the range 2.03–2.08 eV.
most (i) is an acoustic mode directed along \( z \), of high wavelength. On this same phonon branch, the acoustic mode (iv) with half-wavelength also couples strongly. Mode (ii) presents a \( B_u \) representation where strontium atoms of different channels are out of phase. Mode (iii) corresponds to the same mode but with half-wavelength. Indeed, those long-wavelength collective displacements of the Sr atoms along the \( c \)-axis couple strongly with the above-described 5d–4f relaxation of the Sr channel containing Eu.

Above 0.022 eV, the spectral decomposition is made of a multitude of modes containing Al, O and N displacements. We still notice the large peak around 0.035 eV composed of modes that couple strongly with the relaxation of the first coordination shell of Eu (expansion of the nearly cubic environment of Eu). This coupling to a large number of modes instead of one single effective “breathing” mode coincides with the results of Linderälv et al.\(^\text{[5]}\) where a similar feature was found with YAG:Ce\(^{3+}\) phosphor. In Figure 5d, we quantify the importance of Sr-dominated phonon modes by computing their contribution to the total Huang–Rhys factor and find out that the Sr-dominated modes contributes 42% to the narrow emission in Sr[Li\(_2\)Al\(_2\)O\(_2\)N\(_2\)]:Eu\(^{2+}\). For completeness, the convergence of this spectral decomposition as a function of the supercell size is presented in Figures S4 and S5, Supporting Information. It indicates that the result is well converged for high phonon energies while the exact shape of the low energy part of \( S(\hbar \omega) \), composed of modes involving Sr atoms, is not totally converged yet. By increasing the supercell size further, one would simply displace the points highlighted in Figure 5b towards \( \Gamma \) on their corresponding branches. This concern reflects first the difficulty to obtain a strict convergence of the structural relaxation of the Sr channel, as described in Section 3.1. Second, due to the coupling to high-wavelength acoustic phonon modes, obtaining a strict convergence of the low-energy shape requires also to obtain a denser sampling of those phonons. Particular embedded methods to artificially increase the supercell size can be used to tackle this problem.\(^{[16,17]}\) We reserve such improvements for future work. However, we note that the integrated spectral function (i.e., \( S = \sum S_i \)) presents a quicker convergence. In addition, this lack of strict convergence at low energy has only a small impact on the smeared emission intensity \( I(\hbar \omega) \) due to the counter-balancing effect of a decreasing total Huang–Rhys factor and
an increasing effective frequency (see Figure S5, Supporting Information).

The computed effective frequency defined in Equation (13), 25.5 meV, agrees well with the effective frequency of the 1D-model, 25.15 meV, from Equation (5). This make sense since, by defining \( E_{\text{FCG}} = \frac{1}{2} \sum \langle \Delta Q \rangle^2 \omega^2 \), one can show that Equation (5) is equivalent to Equation (13). The computed difference between those two values gives an estimation of the error made by using phonon modes of the undoped structure. Indeed, the first value is computed with the phonon of the undoped structure while the second is computed, within the 1D-model, with the doped structure. We believe that this effect is related to the difference in bond lengths (Sr-O/N/Sr versus Eu-O/N/Sr). This small difference reveals that the phonon modes that contributes to the electronic transitions are already present in the undoped material without the addition of the Eu doping agent. This result is in contrast to the result of Alkauskas et al.\(^{[16]}\) in which the luminescent lineshape of diamond nitrogen-vacancy center could not be correctly described with the phonon modes of the undoped material.

The total Huang–Rhys factor \( S_{\text{multi-D}} = \sum \nu S_{\nu} \) is 3.05, smaller than the 1D \( S_{\text{1D}} = 3.82 \). This difference can be explained by developing Equations (6) and (14). One shows, with the same \( E_{\text{FCG}} \) definition, that \( S_{\text{ID}} \propto \left( \sum \nu S_{\nu} \right)^{1/2} \) is always larger than \( S_{\text{multi-D}} = \sum \nu S_{\nu} = \sum \nu \delta_{\nu} \omega \). \( S_{\text{1D}} \) deviates largely from \( S_{\text{multi-D}} \) when the spread of the spectral function \( S(h\omega) \) is large, which is the case here. We will see that using the Huang–Rhys factor as computed in the more precise multi-dimensional approach leads to a luminescence intensity \( L(h\omega) \) that better matches the experiment.

Following the generating function approach described in Section 2, the emission spectral function \( A(h\omega) \) was calculated based on the spectral decomposition \( S(h\omega) \) (Equation (16)). The resulting curve is then convoluted with a Gaussian of fixed FWHM \( w \). The normalized luminescence intensity is finally obtained considering the cubic frequency dependence \( L(h\omega) = C\omega^3 A(h\omega) \).

On Figure S6, Supporting Information, we present the luminescence intensity as computed in this work at 0 K following 1D model, multi-D model without the Sr-dominated modes (only considering orange area of Figure 5d) and the full multi-D model with all modes. No shift of the zero-phonon line was considered and a small smearing was applied.

In order to compare these results with the experimental spectrum (15 K) and to deal with the indeterminacy of the experimental \( E_{\text{ZPL}} \), we proceed in the following way. For each case, we adjust the shift of the curve (modification of \( E_{\text{ZPL}} \)) and the Gaussian and Lorentzian broadening parameters (\( w \) and \( \gamma \)) to minimize the error defined as the integral of the absolute difference between the simulated and experimental curves (in energy scale), divided by two. Since all spectra are normalized such that \( \int L(h\omega) d(h\omega) = 1 \), this ensures an error bounded to [0 - 1]. The resulting curves are presented in Figure 6.

We find that, in the 1D case, using \( w = 92 \) meV, \( \gamma = 12 \) meV and a shift of 26 meV toward higher energies gives the best fit (error of 12%). In the multi-D case without considering the Sr-dominated modes, using \( w = 30 \) meV, \( \gamma = 75 \) meV, and a shift of 14 meV toward lower energies gives the best fit (error of 8%). When using all the phonon modes in the multi-D model, using \( w = 22 \) meV, \( \gamma = 34 \) meV and a shift of 14 meV toward higher energies gives the best fit (error of 6.5%).

Indeed, the broadening parameters \( \gamma \) and \( w \) used refer physically to homogeneous and inhomogeneous broadening. However, in the case of SALON phosphor, no fine vibronic structure is apparent experimentally, making the exact physical determination of these broadening parameters difficult. Here, \( w \) and \( \gamma \) parameters are used as fitting parameters to match experimental emission spectrum shape. No physical conclusion should be drawn from the values of \( \gamma \) and \( w \) that gives the best fit.

We note first that the 1D model does not reproduce well the shape of the experimental spectrum. Indeed, it has been shown that the single mode approximation is well justified for large coupling \( S \gg 1 \)\(^{[38]}\) which is not the case here. We have also seen that \( S_{\text{1D}} \) overestimates the more precise \( S_{\text{multi-D}} \). Moreover, the spectral decomposition \( S(h\omega) \) is highly asymmetric around the effective frequency: an important weight on a short energy range at low energy and relatively broad distribution after 25–30 meV. This fine feature cannot be captured well by assuming only one effective mode as done in the 1D model.

We observe also that considering all the phonon modes, including the Sr-dominated modes, allows one to obtain the best match with the experiment. In particular, the long tail of the spectrum is well reproduced only with the full computation. This underlies the importance of the long-range Sr displacements in shaping the emission spectrum of SrLi2Al2O2N2:Eu\(^{2+}\). Surprisingly, the spectrum obtained without those Sr modes provides already a good estimate of the experiment. This is however fortuitous: indeed, by only considering the high-energy part of \( S(h\omega) \), the total Huang–Rhys factor is decreased but the effective frequency is increased. On the emission spectrum, these two effects are counter balancing.
each other (see Equation (7)) which explains the small difference between orange and black curve of Figure 6.

To conclude this section, we make the link between the small FWHM of the emission spectrum of SALON with its particular structure. A small FWHM is achieved by reducing at the same time the normal coordinate change $\Delta Q$ and the effective frequency $\hbar \Omega_g$ (see Equation (7)). In the case of SALON, the small $\Delta Q$ comes from the high rigidity of the host structure. The small effective frequency, on the other hand, comes from the large coupling of the Sr channel relaxation with low-frequency phonon modes involving these channels. We believe that these features are not specific to SALON but are also valid for other UCr$_4$C$_7$ type phosphors that present a rigid host structure and this peculiar Ca/Sr/Ba channel.

4. Conclusions

In the present work, we characterize the newly discovered narrow-band emission SrLi$_2$Al$_2$O$_3$:Eu$^{2+}$ red phosphor from first principles. Experimental structural parameters are reproduced within 1% error for undoped SALON. We analyze the structural change of the system upon emission. Two relaxation patterns are distinguished: a nearly isotropic expansion of the cuboid environment of Eu$^{2+}$ composed of 4 oxygen and 4 nitrogen atoms, with small distortion, and a long-range relaxation of the strontium atoms belonging to the strontium channels along the c-axis. At the level of the electronic structure, the $\Delta$SCF method is able to correctly place the 4f and 5d states in the band gap. The lowest 5d state is identified as presenting a 5d$_{x^2-y^2}$ character and shows a large spatial delocalization along the Sr channel. Concerning the luminescent properties, we show that SALON presents a small effective displacement between excited and ground state. By projecting the atomic displacements induced by 4f–5d transition onto the phonon eigenvectors, we identify the phonon modes that couples the most with the transition. This spectral decomposition is dominated by a coupling to low-frequency acoustic and $B_u$ modes involving long-wavelength displacement of these Sr channels, which lowers the effective vibrational frequency. The high-frequency part of this decomposition originates from the coupling to a large number of modes involving O, N and Al. This contradicts the initial guess of Hoeder et al.\cite{Hoeder2021} that argued that the isotropic structural relaxation, coming from the high symmetry environment of Eu$^{2+}$, reduces the number of different vibrational states involved in the emission process. Overall, we computed a total Huang–Rhyys parameter of $S = 3.05$. This small coupling, in combination with a moderate effective frequency, yields a small bandwidth.

Finally, we compute the luminescence intensity spectrum based on this phonon decomposition and compare it with the spectrum obtained with the simpler 1D approach. With respect to experiment, we conclude that the multi-D approach is twice as accurate as the 1D approach. In particular, the tail of the spectrum is very well described within the multi-D approach. These results are important in two aspects. First, they directly provide a theoretical understanding of the observed experimental spectrum. In particular, we highlight the importance of the Sr channel oriented along c-axis. The spectrum of other UCr$_4$C$_7$-type phosphors that present similar Ca/Sr/Ba channel might be similarly affected. Second, the success of the methodology used provides confidence in its potential applicability in the theoretical study of other important phosphors and their luminescent intensity spectra. It could be for instance interesting to apply this multi-dimensional approach to a low-temperature phosphor emission spectrum that still presents an apparent vibronic fine structure, allowing a further validation of the computational methodology.

5. Computational Method

5.1. DFT Parameters

Calculations are performed using density-functional theory (DFT) using ABINIT\cite{abinit} Core electrons are treated with the PAW method, with JTH atomic data.\cite{JTH} The generalized gradient approximation (GGA-PBE) is used to treat exchange-correlation effects\cite{Perdew1996} and a Hubbard U term ($U=7$ eV) is added on 4f states of europium, in line with the value used in a study of fifteen Eu-doped materials.\cite{Pascal2015}

Calculations on europium-doped SALON are conducted with a supercell technique. Most results shown in this manuscript are computed with a $1 \times 1 \times 12$ supercell composed of 216 atoms. Since there are 24 Sr atoms in this supercell, this corresponds to a theoretical doping concentration of 4.16% while the experimental one is 0.7%. Still, since our methodology does not account for concentration-dependent effects such as self-absorption or energy transfer, our only concern is the spurious interaction between europium periodic replica, that should be numerically controlled. To this aim, we performed a careful convergence study of our main results with respect to the supercell size, that can be found in Supporting Information. Because of the anisotropy of the material (tetragonal structure), we checked both an increase of the size in the $x$–$y$ plane and an increase along $z$ axis. It appears that the critical parameter to converge is the long-range structural relaxation of the strontium atoms along $z$-axis, explaining the particular elongated shape of our supercell.

For all supercell calculations, the structures are relaxed below a maximal residual force of $10^{-5}$ Hartree/Bohr. The cut-off kinetic energy is 30 Ha and a $2 \times 2 \times 2$ wavevector grid is used. Phonons are calculated for the undoped structures at the $\Gamma$ point.

Although spin–orbit coupling can present a noticeable effect on the excited state electronic configuration, it is not considered here due to its small effect on the emission spectra. This work focuses on the lowest excited state of the 4f$^5$5d$^1$ multiplet ($^7F_J$, $J = 0$ and $6$) splitting and does not try to account for the other states.

5.2. Simulating the Eu 5d Excited State

The treatment of the Eu 4f$^5$5d$^1$ excited state and the resulting Eu 5d electron and 4f hole interaction might rely on the Bethe–Salpeter equation\cite{Bethe1930} but this would be too computationally demanding for such supercell system. For this reason, we rely on the $\Delta$SCF method whereby the eigenfunction associated to the highest Eu 4f-like state is forced to be unoccupied while the
next energy state (close to a 5d₂ state) is constrained to be occupied. Transition energies are then computed as difference of two total energies. The electron–hole interaction is here mimicked through the promotion of the Eu 4f electron to the Eu 5d state. By doing that, the seven 4f levels split into one unoccupied band that stays within the gap, and six occupied bands that shift downwards, hybridizing inside the valence band.[13]

We suppose that the 4f–5d transition only affects the local geometry of the crystal and not the macroscopic state of the crystal, meaning that the lattice parameters of the crystal will not change. Thus, atomic relaxation in the excited state is done at fixed lattice parameters obtained in the ground state.

Additional details on the ΔSCF method are given in Supporting Information.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest
The authors declare no conflict of interest.

Data Availability Statement
Research data are not shared.

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