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The crystal structure and luminescence properties of the first lithium oxonitridolithosilicate $\text{Li}_3\text{SiNO}_2\text{:Eu}^{2+\ddagger}$

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The compound $\text{Li}_3\text{SiNO}_2\text{:Eu}^{2+}$ was synthesized in high temperature solid-state reactions in weld shut tantalum ampules and the crystal structure of Li_3SiNO_2 has been determined by single-crystal X-ray diffraction. It crystallizes in the monoclinic space group $C2/c$ (no. 15) with the lattice parameters $a = 1049.01(3)$, $b = 1103.42(3)$, $c = 511.86(2)$ pm, $\beta = 116.14(1)^\circ$, and a volume of $V = 0.53187(2)$ nm³. This compound is built up from two different layers, which are arranged alternately along the crystallographic a -axis. The results from single-crystal diffraction were confirmed by the Rietveld analysis of bulk samples. Moreover, Li_3SiNO_2 could be successfully doped with the activator ion Eu^{2+} and the luminescence spectroscopy of single-crystals revealed broad band emission at $\lambda_{\text{max}} = 601$ nm (fwhm = 90 nm).

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Introduction

Although the field of lithium oxonitridosilicates has been known for quite a long time and the first compound $\text{Li}_4\text{Si}_{0.8}\text{N}_{2.4}(\text{Li}_2\text{O})_{1.6}$ was published by Juza, Weber, and Meyer-Simon in 1953,¹ only a few more representatives are known today. A few years later (1981), the well-known LiSiON was synthesized by Laurent, Guyader, and Roullet and analyzed by time-of-flight neutron diffraction.² Ma *et al.* were able to successfully dope the compound LiSiON with the activator ion Eu^{2+} in 2012, which resulted in emission with a maximum of $\lambda_{\text{max}} = 426\text{--}478$ nm (fwhm = 142–193 nm) depending on the activator ion concentration.³ Based on X-ray phase analysis and classical quantitative analysis, compounds such as $\text{Li}_4\text{SiN}_2\text{O}$, Li_3SiNO_2 or $\text{Li}_7\text{SiN}_3\text{O}$ were predicted by Kraśnicka and Podsiadlo at the same time.^{4,5} The structure of the last compound, $\text{Li}_7\text{SiN}_3\text{O}$, was then elucidated by Casas-Cabanas and Palacín in 2014 using Rietveld refinement.⁶ All lithium oxonitridosilicates, whose structures are known to date, contain lithium ions exclusively in tetrahedral coordination. This results in struc-

tures that consist only of the combined network of tetrahedrally coordinated silicon and lithium ions. In the following, lithium oxonitridolithosilicate compounds shall be referred to as structures, which are built up by the aforementioned (Li/Si)–(O/N) tetrahedral network but additionally with lithium as a completing cation which occupies the cavities. Although no luminescent lithium oxonitridolithosilicate is known yet, there are already some established phosphors in the field of alkali lithosilicates.^{7–10} The best common examples are $\text{Na}[\text{Li}_3\text{SiO}_4]\text{:Eu}^{2+}$ (NL $\text{SO}:\text{Eu}^{2+}$) ($\lambda_{\text{max}} = 469$ nm; fwhm = 32 nm), $\text{K}[\text{Li}_3\text{SiO}_4]\text{:Eu}^{2+}$ (KL $\text{SO}:\text{Eu}^{2+}$) ($\lambda_{\text{max}} = 604$ nm; fwhm = 150 nm), $\text{NaK}_7[\text{Li}_3\text{SiO}_4]_8\text{:Eu}^{2+}$ (NKLSO: Eu^{2+}) ($\lambda_{\text{max}} = 515$ nm and 598; fwhm = 49 nm and 138 nm),^{11–14} $\text{RbNa}_3[\text{Li}_3\text{SiO}_4]_4\text{:Eu}^{2+}$ ($\lambda_{\text{max}} = 471$ nm; fwhm = 22.4 nm),^{8,15,16} $\text{RbNa}[\text{Li}_3\text{SiO}_4]_2\text{:Eu}^{2+}$ ($\lambda_{\text{max}} = 523$ nm; fwhm = 41 nm),¹⁷ $\text{RbLi}[\text{Li}_3\text{SiO}_4]_2\text{:Eu}^{2+}$ (RLSO: Eu^{2+}) ($\lambda_{\text{max}} = 530$ nm; fwhm = 42 nm),¹⁸ $\text{RbNa}_2\text{K}[\text{Li}_3\text{SiO}_4]_4\text{:Eu}^{2+}$ ($\lambda_{\text{max}} = 480$ nm; fwhm = 26 nm),¹⁹ $\text{CsNa}_2\text{K}[\text{Li}_3\text{SiO}_4]_4\text{:Eu}^{2+}$ ($\lambda_{\text{max}} = 485$ nm; fwhm = 26 nm),¹⁹ $\text{LiK}_7[\text{Li}_3\text{SiO}_4]_8\text{:Eu}^{2+}$ (LKLSO: Eu^{2+}) ($\lambda_{\text{max}} = 511$ nm; fwhm = 45 nm),²⁰ $\text{Na}_2\text{K}_2[\text{Li}_3\text{SiO}_4]_4\text{:Eu}^{2+}$ ($\lambda_{\text{max}} = 486$ nm; fwhm = 20.7 nm),^{21,22} $\text{RbKLi}_2[\text{Li}_3\text{SiO}_4]_4\text{:Eu}^{2+}$ ($\lambda_{\text{max}} = 474$ nm and 532; fwhm = 24.8 nm and 43.5 nm),^{7,23} and $\text{Cs}_{4-x-y-z}\text{Rb}_x\text{Na}_y\text{Li}_z[\text{Li}_3\text{SiO}_4]_4\text{:Eu}^{2+}$ (CRNLSO: Eu^{2+}) ($\lambda_{\text{max}} = 473$ nm and 531 nm; fwhm = 25.2 nm and 58 nm).²⁴ These compounds exhibit interesting luminescence properties and some of them are characterized by their narrow single band emission in the blue and green spectral range.

In this article, we present the structure of Li_3SiNO_2 based on single-crystal X-ray data refinement. Additionally, the title phase was doped with the activator ion Eu^{2+} and, in comparison with the well-known alkali lithosilicates, nitrogen was suc-

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cessfully incorporated into the crystal structure resulting in a shift of the emission band to higher wavelengths compared to the aforementioned oxosilicates.^{25–27} The here presented compound can be seen as the first example in the field of lithium oxonitridolithosilicates and additionally it exhibits single band emission in the orange spectral region, which is a novelty even for the substance class of oxonitridolithosilicates.

Results and discussion

Crystal structure

The compound Li_3SiNO_2 crystallizes in the monoclinic space group $C2/c$ with the lattice parameters $a = 1049.01(3)$, $b = 1103.42(3)$, $c = 511.86(2)$ pm, $\beta = 116.14(1)^\circ$, and a volume of $V = 0.53187(2)$ nm³. Details on the crystal-structure refinement are given in Table 1. Atomic coordinates, displacement parameters, and interatomic distances are listed in Tables 2–4.

The three-dimensional framework in the structure of Li_3SiNO_2 is built up from two different layers, which are stacked alternately along the crystallographic a -axis. As shown in Fig. 1, the first layer, named layer A, consists of condensed SiO_2N_2 and LiO_3N tetrahedra, where the vertex-sharing SiO_2N_2

tetrahedra form unbranched *zweier* single chains²⁸ along the crystallographic c -axis *via* bridging $\text{N}^{2/3-}$ anions. The structural feature of such unbranched *zweier* single chains was found for the first time in nature in the mineral diopside $\text{CaMgSi}_2\text{O}_6$, which belongs to the mineral group of pyroxenes and, according to the silicate classification by Liebau, it is an inosilicate.^{28–30} SiO_2N_2 tetrahedra can also be found in compounds such as $\text{Ca}_3\text{SiO}_4\text{N}_2$,³¹ where these tetrahedra are also connected to each other *via* N^{3-} anions. The Si–O distances, which vary from 164.53(4) to 167.92(4) pm, and the Si–N distances with values from 171.46(5) to 172.31(5) pm are in good agreement with the sum of the ionic radii.^{23,24,32,33} The occupation of the N1 site with O instead of N leads to a significant under-occupation, indicating the presence of N at this position. These unbranched *zweier* single chains are then connected to each other *via* LiO_3N tetrahedra, which form $\text{Li}_2\text{O}_4\text{N}_2$ dimer units, when only layer A is considered.³⁴ Hereby two LiO_3N tetrahedra share a common edge *via* the O^{2-} anions. Although the dimer units are not connected to each other, they connect the unbranched *zweier* single chains in such a way that they are corner-sharing *via* the N^{3-} anions and edge-sharing *via* two O^{2-} anions. In the LiO_3N tetrahedra, the Li–O distances range from 188.9(2) to 241.8(2) pm and the Li–N distance is 195.0(2) pm. Since one Li–O distance is quite elongated, the corresponding coordination polyhedra can also be described as elongated trigonal pyramids exhibiting a 3 + 1 coordination, where the Li^+ cations are slightly displaced from the centers (see Fig. 1).^{12,33,35}

The connectivity within layer A in the structure of Li_3SiNO_2 is related to that in the compounds $\text{Na}_2\text{Mn}_2\text{S}_3$ ³⁶ and $\text{Na}_6\text{In}_2\text{S}_6$,³⁷ where the SiO_2N_2 tetrahedra correspond to the MnS_4 tetrahedra (Mn1 site) and NaS_4 tetrahedra, respectively, and the LiO_3N tetrahedra correspond to the MnS_4 tetrahedra (Mn2 site) and InS_4 tetrahedra, respectively (see Fig. 2).

The second layer, named layer B, is built up from three different coordination polyhedra consisting of LiO_4 tetrahedra, LiO_2N_4 octahedra, and LiO_4N_2 octahedra, whereby within the latter dynamic positional disorder occurs. In this layer, there are five crystallographically distinguishable lithium sites. The Li1 and Li2 sites form LiO_4 tetrahedra with Li1–O and Li2–O distances ranging from 188.0(2) to 196.9(2) pm and 196.7(2) to 213.8(2) pm, respectively, which are in good agreement with the sum of the ionic radii.^{20,23} The Li3 site is coordinated by four N^{3-} and two O^{2-} anions and constitutes a distorted octahedral arrangement, where the Li3–O distances are 205.0(2) pm and the Li3–N distances are between 218.9(2) and 248.94(5) pm. Since two of the Li3–N distances are slightly longer and are located opposite to each other, this can also be described as an elongated octahedron. For Li4/4A, dynamic positional disorder occurs in the LiO_4N_2 octahedra, where the blurred electron density can be described by two partially occupied Li^+ positions (Li4 and Li4A). The two sites are coordinated by four O^{2-} anions and two N^{3-} anions, where the Li4–O distances range from 200.58(4) to 292.0(3) pm and the Li4–N distance is 222.3(2) pm. The Li4A–O distances range from 204.7(2) to 254.4(8) pm and the Li4A–N distance is 255.8(8) pm, implying a highly distorted octahedron. Due to the spatial

Table 1 Crystal data and structure refinement of Li_3SiNO_2

Empirical formula	Li_3SiNO_2
Molar mass/g mol ^{−1}	94.92
Crystal system	Monoclinic
Space group	$C2/c$
Powder data	
Powder diffractometer	STOE Stadi P
Radiation	Mo-K α_1 ($\lambda = 70.93$ pm)
a /pm	1054.82(8)
b /pm	1102.96(7)
c /pm	512.56(4)
β /°	116.66(1)
V /nm ³	0.53293(6)
Single-crystal data	
Single-crystal diffractometer	Bruker D8 Quest Kappa
Radiation	Mo-K α ($\lambda = 71.073$ pm)
a /pm	1049.01(3)
b /pm	1103.42(3)
c /pm	511.86(2)
β /°	116.14(1)
V /nm ³	0.53187(2)
Formula units per cell Z	8
Calculated density/g cm ^{−3}	2.371
Crystal size/mm ³	0.200 × 0.070 × 0.070
Temperature/K	297(2)
Detector distance/mm	40
Exposure time	0.5° per frame; 40 s per frame
Absorption coefficient/mm ^{−1}	0.604
$F(000)/e$	368
θ -Range/°	2.84–40.25
Range in hkl	±19, ±20, ±9
Reflections total/independent	24 474/1622
R_{int}	0.0318
Reflections with $I \geq 2\sigma(I)$	1579
Data/ref. parameters	1622/66
Absorption correction	Multi-scan
Final R_1/wR_2 [$I \geq 2\sigma(I)$]	0.0222/0.0591
Final R_1/wR_2 (all data)	0.0226/0.0594
Goodness of fit on F^2	1.147
Largest diff. peak/hole/e \AA^{-3}	0.516/−0.640



Table 2 Wyckoff positions, atomic coordinates, and equivalent isotropic displacement parameters U_{eq} (\AA^2) of Li_3SiNO_2 (standard deviations in parentheses)

Atom	Wyckoff-position	x	y	z	U_{eq}	Occ.
Si1	8f	0.22086(2)	0.43007(2)	0.25806(3)	0.00621(6)	1
Li1	4e	$\frac{1}{2}$	0.2978(2)	$\frac{1}{4}$	0.0101(2)	1
Li2	4e	$\frac{1}{2}$	0.4746(2)	$\frac{3}{4}$	0.0153(3)	1
Li3	4e	0	0.4338(2)	$\frac{3}{4}$	0.0205(4)	1
Li4	4e	0	0.3190(3)	$\frac{1}{4}$	0.0188(4)	0.76
Li4A	4e	0	0.275(2)	$\frac{1}{4}$	0.019(2)	0.24
Li5	8f	0.2769(2)	0.1976(2)	0.2176(3)	0.0198(3)	1
O1	8f	0.12887(4)	0.31224(3)	0.05472(8)	0.00791(8)	1
O2	8f	0.38906(4)	0.39292(5)	0.3760(2)	0.01470(9)	1
N1	8f	0.17022(5)	0.44522(4)	0.5324(2)	0.00872(8)	1

Table 3 Anisotropic displacement parameters U_{ij} (\AA^2) of Li_3SiNO_2 (standard deviations in parentheses). The displacement parameters of the atoms Li4 and Li4A were refined isotropically

Atom	U_{11}	U_{22}	U_{33}	U_{23}	U_{13}	U_{12}
Si1	0.00646(8)	0.00652(8)	0.00580(7)	-0.00072(3)	0.00283(5)	0.00006(4)
Li1	0.0103(6)	0.0084(5)	0.0133(6)	0	0.0068(5)	0
Li2	0.0161(7)	0.0172(7)	0.0098(6)	0	0.0031(5)	0
Li3	0.0180(8)	0.0129(7)	0.0179(8)	0	-0.0039(7)	0
Li5	0.0255(6)	0.0191(6)	0.0205(6)	0.0106(4)	0.0154(5)	0.0119(5)
O1	0.0082(2)	0.0068(2)	0.0081(2)	-0.00118(9)	0.0031(2)	-0.0003(2)
O2	0.0077(2)	0.0199(2)	0.0150(2)	-0.0062(2)	0.0035(2)	0.0019(2)
N1	0.0129(2)	0.0072(2)	0.0076(2)	0.0002(2)	0.0059(2)	0.0008(2)

Table 4 Interatomic distances (\AA) in Li_3SiNO_2 (standard deviations in parentheses)

Si1–O2	1.6453(4)	Li3–O1	2.050(2) 2×	Li4A–O1	2.047(2) 2×
Si1–O1	1.6792(4)	∅ Li3–O	2.050(2)	Li4A–O2	2.544(8) 2×
∅ Si1–O	1.6668(4)			∅ Li4A–O	2.296(8)
		Li3–N1	2.189(2) 2×	Li4A–N1	2.558(8) 2×
Si1–N1	1.7146(5)	Li3–N1	2.4894(5) 2×	∅ Li4A–N	2.558(8)
Si1–N1	1.7231(5)	∅ Li3–N	2.339(2)		
∅ Si1–N	1.7189(5)				
		Li4–O1	2.0058(4) 2×	Li5–O1	1.889(2)
Li1–O2	1.880(2)	∅ Li4–O	2.0058(4)	Li5–O1	2.036(2)
Li1–O1	1.969(2) 3×			Li5–O2	2.418(2)
∅ Li1–O	1.947(2)	Li4–N1	2.223(2) 2×	∅ Li5–O	2.114(2)
		∅ Li4–N	2.223(2)		
Li2–O2	1.967(2) 2×			Li5–N1	1.950(2)
Li2–O2	2.138(2) 2×			∅ Li5–N	1.950(2)
∅ Li2–O	2.053(2)				

proximity, only one of the two positions (Li4 and Li4A) can be occupied at the same time and due to a strong correlation between the two sites, only an isotropic refinement of the displacement parameters in the structure solution was applied. For comparison, the displacement ellipsoids of the asymmetric unit of Li_3SiNO_2 are shown in the ESI (see Fig. S1†). The asymmetric units are illustrated for the cases where the Li4 site is described by only one fully occupied Li site and by two partially occupied Li sites, respectively. The occupation of the two sites was approximated in such a way that the two isotropic displacement parameters were approximately equal in size and then fixed manually at this value. The crystal structure refinement results in occupations of $\text{Li4} = 0.76$ and $\text{Li4A} = 0.24$ implying that the Li4 site is statistically preferred. To confirm

the occupations of these sites, five crystals from four different syntheses were measured by single-crystal X-ray diffraction at room temperature. All crystal structure refinements showed the same results with minimal deviations. Single-crystal X-ray measurements at low temperatures ($-90\text{ }^\circ\text{C}$) have shown that the occupation of the Li4 and Li4A sites changes by 8% ($\text{Occ. of Li4} = 0.84$ and $\text{Li4A} = 0.16$) and the Li4 site is more preferred. This indicates dynamic positional disorder between the two sites, which is temperature dependent. The results of the single-crystal measurements at low temperature are shown in the ESI,† listed in Tables S2–S5.† One reason for the dynamic positional disorder could be that the volume of the LiO_4N_2 octahedron ($V = 16.73\text{ \AA}^3$) is larger than that of the LiO_2N_4 octahedron ($V = 14.19\text{ \AA}^3$) and thus the Li^+ cation has more



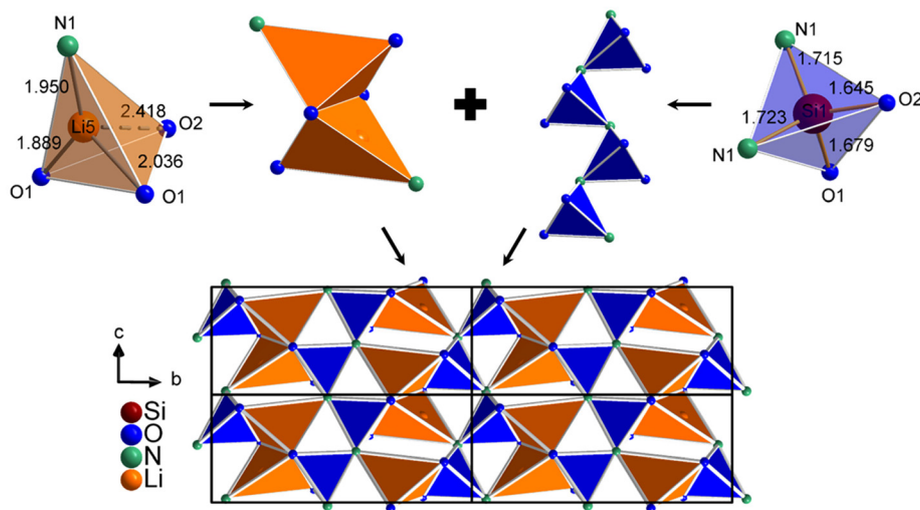


Fig. 1 Layer A of Li_3SiNO_2 . In the upper left, the LiO_3N tetrahedra (orange) with the Li–N and Li–O distances and the dimer units formed from them are shown. In the upper right, the SiO_2N_2 tetrahedron (blue) is illustrated, including the Si–N and Si–O distances, and the unbranched zweier single chains built up from it. In the lower half, layer A, which is formed from LiO_3N and SiO_2N_2 tetrahedra, is represented along $[\bar{1}00]$.

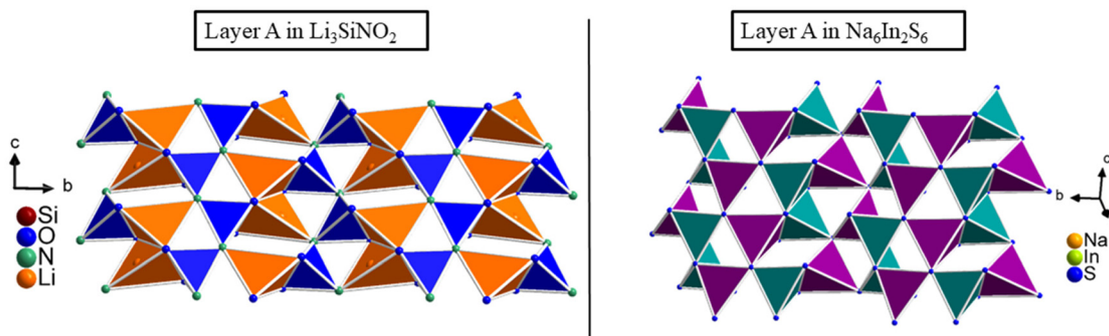


Fig. 2 On the left, layer A of Li_3SiNO_2 is shown compared to layer A of the compound $\text{Na}_6\text{In}_2\text{S}_6$, which is represented on the right. The LiO_3N tetrahedra are displayed in orange, SiO_2N_2 tetrahedra are displayed in blue, the NaS_4 tetrahedra are displayed in violet and the InS_4 tetrahedra are displayed in teal.

space to displace, which becomes higher with increasing temperature (see Fig. 3). The program Vesta 3³⁸ was used to calculate the polyhedron sizes.

The four different coordination polyhedra around the four Li sites, as mentioned before, form an interconnected tetrahedra and octahedra band spanning the condensed layer B (see Fig. 4). The LiO_2N_4 and LiO_4N_2 octahedra built up the octahedra band, named band X, along the crystallographic c -axis, where the LiO_2N_4 octahedra are double edge-sharing to each other by two common N^{3-} anions on each edge. The LiO_4N_2 octahedra, on the other hand, are not connected to each other, but share common edges with the LiO_2N_4 octahedra within the octahedra band X. The LiO_2N_4 and LiO_4N_2 octahedra are connected in such a way that one LiO_4N_2 octahedron shares common edges with three LiO_2N_4 octahedra once *via* two N^{3-} anions and twice *via* one O^{2-} and one N^{3-} anions. Considering the cb -plane, the LiO_4N_2 octahedra are arranged alternately to the right and to the left of the LiO_2N_4 octahedra. A similar con-

nection pattern is formed by the LiO_4 tetrahedra within the tetrahedra band, named band Y. The LiO_4 tetrahedra of the Li2 site are connected to each other along the crystallographic c -axis by double edge-sharing *via* two common O^{2-} anions on each edge. The LiO_4 tetrahedra of the Li1 site, which are not connected to each other within the tetrahedral band Y, always arrange themselves alternately to the left and right of the LiO_4 tetrahedra of the Li2 site from the viewpoint perpendicular to the cb -plane. The LiO_4 tetrahedra of the Li1 and Li2 sites share a common edge *via* two O^{2-} anions. Along the crystallographic b -axis, the bands X and Y are arranged alternately. The bands X and Y are connected in such a way that the LiO_4 tetrahedra of the Li1 site share a common edge with the LiO_2N_4 octahedra and two common edges with the LiO_4N_2 octahedra, each *via* two O^{2-} anions. Additionally, the LiO_4N_2 octahedra and the LiO_4 tetrahedra of the Li2 site are edge-sharing *via* two O^{2-} anions. Furthermore, the c glide plane, derived from the space group $C2/c$, is visible when the mirror plane is placed perpen-



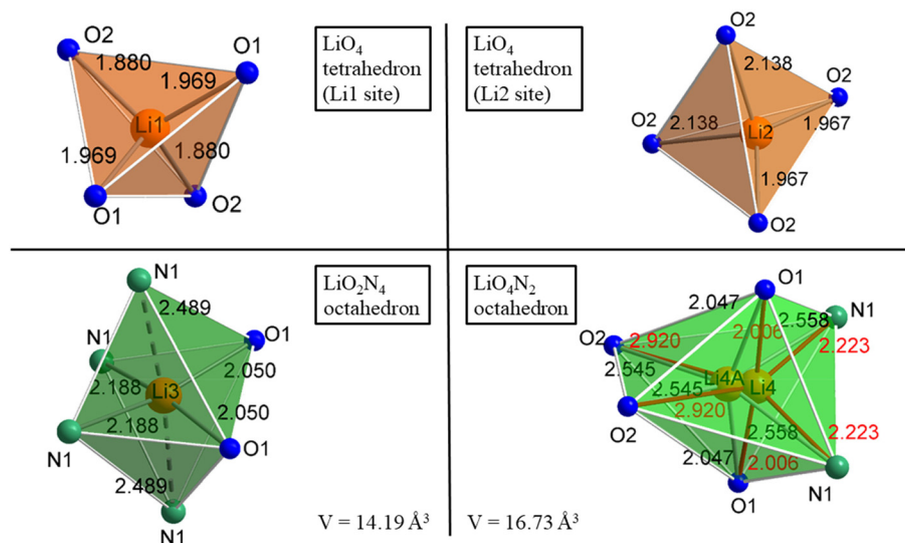


Fig. 3 The different coordination polyhedra of layer B are shown. On the upper half, the LiO_4 tetrahedra of the Li1 and Li2 sites are displayed in orange including the Li–O distances. On the lower left, the LiO_2N_4 octahedron is displayed in green including the Li–O and Li–N distances and the polyhedron size, respectively. On the lower right, the LiO_4N_2 octahedron with the two partially occupied Li^+ positions is represented in bright green including the Li–O and Li–N distances and the polyhedron size. The Li–O and Li–N distances of the Li4 site are shown in red.

dicular to the crystallographic b -axis through the center of the octahedral band X (see Fig. 4). Since the Li^+ cations are not only part of the tetrahedral substructure, we assign this compound to the substance class of lithium oxonitridolithosilicates.

The entire structure of Li_3SiNO_2 is built up from the two layers A and B by stacking them alternately along the crystallographic a -axis with the characteristic pattern BAB'A' (see Fig. 5). Although layers A and B are equivalent in construction to layers A' and B', they are differently orientated due to the most favorable arrangement in terms of energy. Layers A and A' are related *via* a two-fold rotation axis that runs along the crystallographic b -axis through layer B. Layers B and B' are connected *via* a 2_1 screw axis, which is additionally caused by the space group $C2/c$ and runs along the crystallographic b -axis through layers A and A'. The two layers A and B are connected in such a way that the LiO_3N tetrahedra of the Li5 site share a common edge with the LiO_4 tetrahedra of the Li1 site *via* two O^{2-} anions, and are edge-shared to the LiO_4N_2 octahedra and the LiO_2N_4 octahedra *via* an O^{2-} and N^{3-} anion. In addition, the LiO_4 tetrahedra of the Li5 site are also corner-shared to the LiO_4 tetrahedra of the Li2 site *via* an O^{2-} anion. The SiO_2N_2 tetrahedra of the unbranched *zweier* single chains show a similar connection pattern. The SiO_2N_2 tetrahedra share common corners with the LiO_4 tetrahedra of the Li2 sites *via* O^{2-} anions. They share a common edge with the LiO_2N_4 octahedra *via* two N^{3-} anions as well as *via* one N^{3-} and one O^{2-} anion. In addition, it is edge-shared to the LiO_4N_2 octahedra *via* an O^{2-} and N^{3-} anion.

The compound Li_3SiNO_2 could be successfully doped with the activator ion Eu^{2+} and the resulting luminescence properties are discussed in the Luminescence section.

MAPLE, CHARDI, and BLBS

The electrostatic consistency of the crystal structure is proved by MAPLE (Madelung Part of Lattice Energy),^{39–41} BLBS (bond length–bond strength),^{42,43} and CHARDI (Charge Distribution method)^{44,45} calculations. In the latter method, the charge distribution is calculated using the concept of effective coordination numbers in combination with bond strength based on Pauling's concept. The BLBS calculations predict the formal oxidation state of an atom by correlating the bond strengths (bond valences) and bond lengths, where the sum of all bond valences should add up to the oxidation state of the respective atom. The MAPLE and CHARDI calculations were performed for the cases where either the Li4 or the Li4A site is fully occupied. The results of the CHARDI and BLBS calculations are shown in Table 5. The partial MAPLE values and MAPLE sums of Li_3SiNO_2 are listed in Table 6.

The results from the BLBS method show the expected charge for the atoms with the exception of the Li4A atom and are in good accordance with the oxidation states based on crystallographic criteria. The outcome of the BLBS method is supported by the CHARDI calculations, where only the atoms O1 and N1 show larger deviations. The deviations of the N1 and O1 atoms could be explained by the fact that they are spatially very close to the dynamic positional disordered Li4 and Li4A sites and thus interact with them more strongly than the O2 atom. The deviation of the Li4A atom with respect to the BLBS calculation could be due to the fact that the position could describe the blurred electron density very well, but does not correspond to a “typical” Li position in relation to the coordination and the partially longer Li–O/N distances, so the description of a dynamic process with a static model is



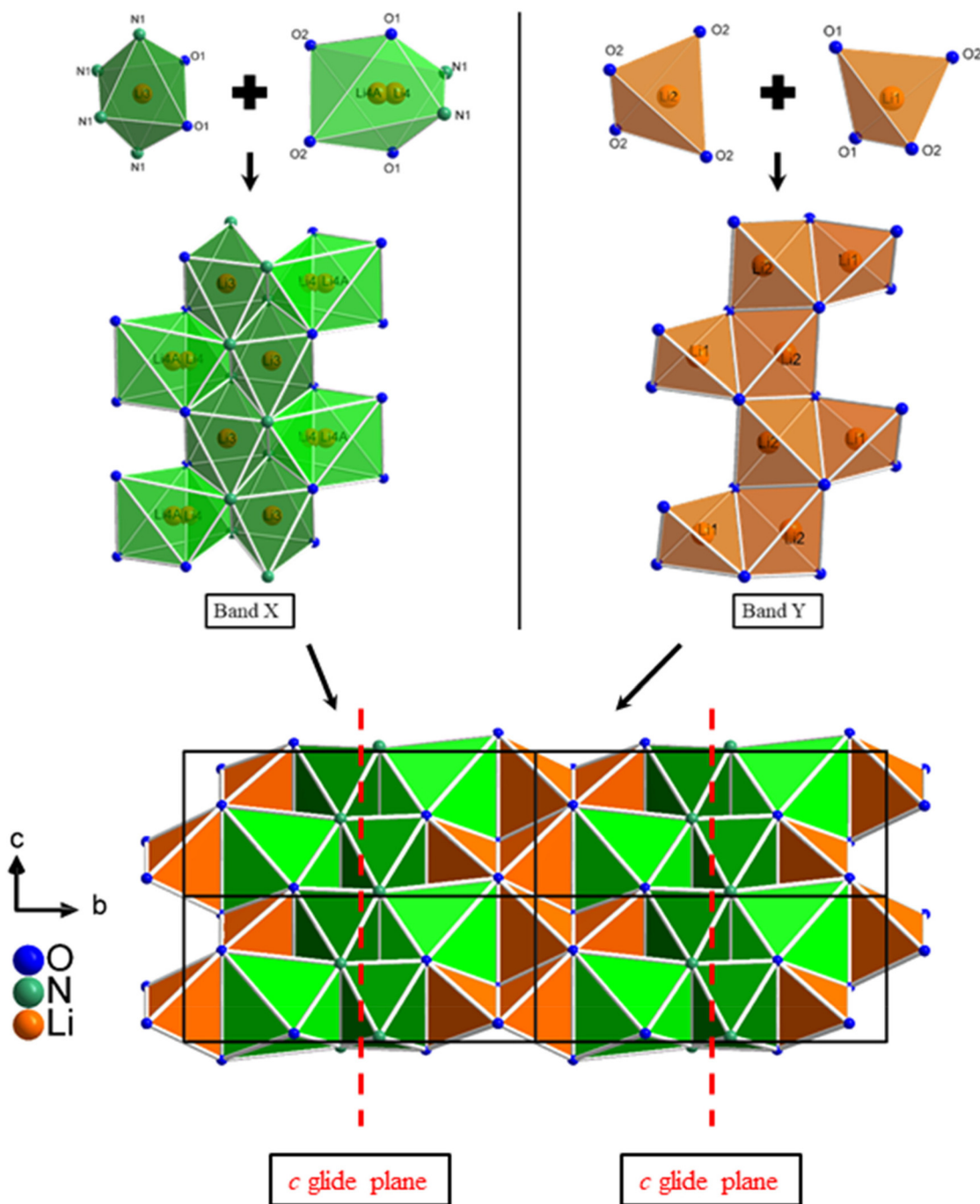


Fig. 4 The highly condensed layer B. In the upper half, the construction of the two different bands (X and Y) are presented. In the lower half, the arrangement and interconnection of the two different bands (X and Y), that build up layer B, are shown. The LiO₄ tetrahedra of the Li1 and Li2 sites are displayed in orange, the LiO₂N₄ octahedra are displayed in green, and the LiO₄N₂ octahedra including the dynamic positional disorder of the Li4 and Li4A sites are represented in bright green.

limited. Furthermore, the partial MAPLE values and the MAPLE sums are compared to the reference values to verify the refined crystal structure. The partial MAPLE values of all individual atoms are in good agreement with the reference values known from the literature and confirm the assignment of the different atoms and the incorporation of N and O into the structure. The results of the calculations are consistent with

those from the SCXRD refinement, and the assignment of the N and O atoms to the corresponding sites seems plausible. Comparing the MAPLE sums of LiSiON and Li₂O with the calculated MAPLE sums of Li₃SiNO₂, the values were found to deviate by only 0.26 and 0.38% depending on the full occupation of the Li4 or Li4A site, which indicates the validity of the crystal structure.



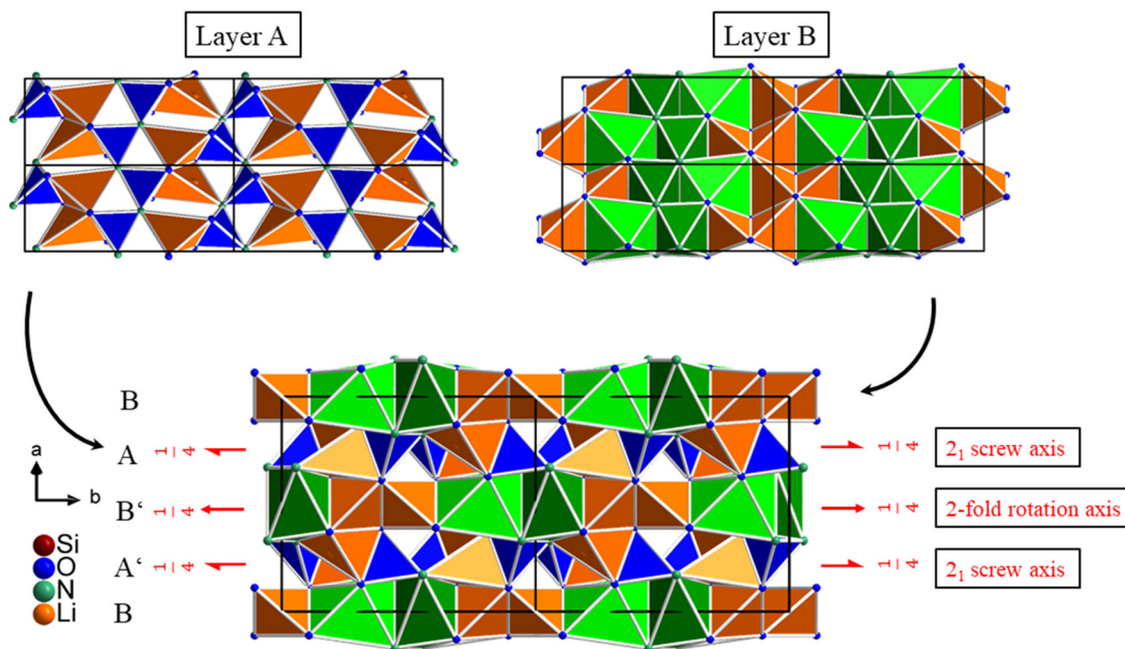


Fig. 5 The entire structure of Li_3SiNO_2 built up from layers A and B. The LiO_4 tetrahedra of the Li1 and Li2 sites and the LiO_3N tetrahedra are displayed in orange, the SiO_2N_2 tetrahedra are displayed in blue, the LiO_2N_4 octahedra are displayed in green and the LiO_4N_2 octahedra including the dynamic positional disorder of the Li4 and Li4A sites are represented in bright green.

Table 5 Charge distribution in Li_3SiNO_2 calculated via CHARDI (ΣQ) and BLBS (ΣV). $\Sigma Q1$ are the values where the Li4 site is fully occupied and $\Sigma Q2$ are the values where the Li4A site is fully occupied

	Si1	Li1	Li2	Li3	Li4	Li4A	Li5	O1	O2	N1
ΣV	4.10	1.10	1.00	1.02	0.89	0.68	1.01	-2.09	-1.85	-3.07
$\Sigma Q1$	4.15	0.92	1.01	0.98	0.93	—	0.93	-2.50	-1.97	-2.52
$\Sigma Q2$	4.21	0.98	0.99	0.99	—	0.83	0.94	-2.59	-2.02	-2.40

Table 6 Partial MAPLE values and MAPLE sums (kJ mol^{-1} of Li_3SiNO_2)^a

Li_3SiNO_2	Li4 fully occupied	Li4A fully occupied	Model
Si1	9200	9261	
Li1	808	780	
Li2	678	635	
Li3	637	696	
Li4	667	—	
Li4A	—	644	
Li5	653	648	
O1	2573	2549	+1
O2	2529	2592	+1
N1	5783	5646	
	$\Sigma = 22\ 133$	$\Sigma = 22\ 106$	$\Sigma = 22\ 191$
	$\Delta = 0.26\%$	$\Delta = 0.38\%$	

^a Typical partial MAPLE values (kJ mol^{-1}): Si^{4+} 9000–10 200; Li^+ 600–860; O^{2-} 2000–2800; N^{3-} 5000–6200.⁴⁷ The MAPLE sums of LiSiON and Li_2O were calculated from the published single-crystal data.

Luminescence

The luminescence properties of $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ were investigated on single-crystals and powder samples, where the latter

was used to determine the excitation spectrum and the thermal quenching behavior.

The powder sample of Li_3SiNO_2 could not be synthesized in the pure phase and contains 3 wt% of LiSi_2N_3 . The results of the Rietveld refinement can be found in the ESI (see Fig. S2 and Table S1†). Li_3SiNO_2 with a nominal activator concentration of 2 mol% Eu^{2+} based on the content of lithium can be excited with near-UV to blue light (e.g. $\lambda_{\text{exc}} = 448$ nm) and the single-crystals exhibit broad band emission at $\lambda_{\text{max}} = 601$ nm with a full width at half maximum (fwhm) of 90 nm (0.32 eV, 2597 cm^{-1}). In comparison, the powder sample ($\lambda_{\text{exc}} = 400$ nm) exhibits broad band emission at $\lambda_{\text{max}} = 606$ nm with a full width at half maximum (fwhm) of 100 nm (0.33 eV, 2695 cm^{-1}). In the CIE-xy color space, this corresponds to the values $x = 0.564(1)$ and $y = 0.433(1)$ for the measured single-crystal emission band and to $x = 0.562(1)$ and $y = 0.436(1)$ for the powder sample. Furthermore, an excitation spectrum of the powder sample was recorded exhibiting a maximum at 376 nm (see Fig. 6a and b). The first lab samples without any systematic engineering regarding the emission properties already yielded quantum efficiencies (QE) above 30%. The compound has a brown to gray body color



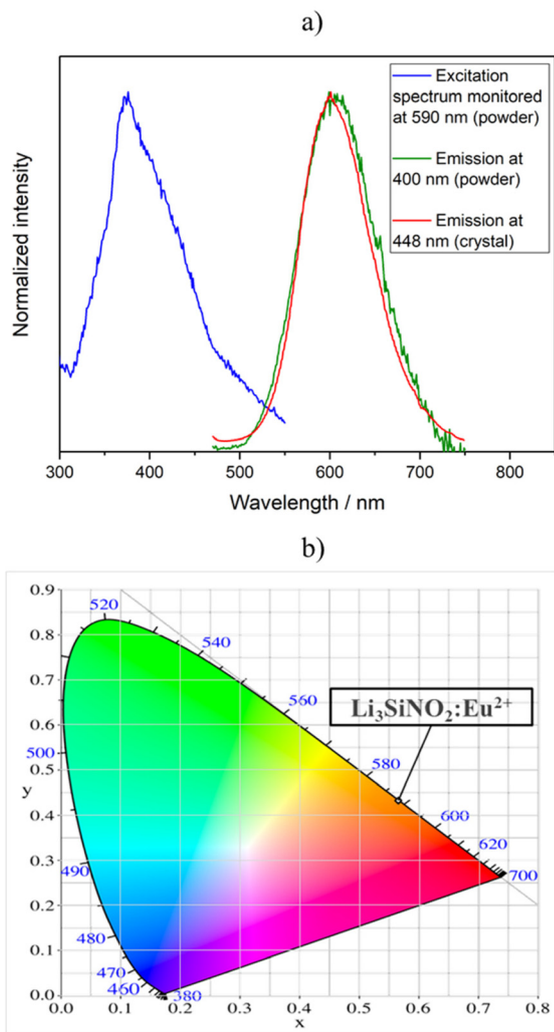


Fig. 6 (a) The red line represents the luminescence spectrum of a $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ crystal recorded at an excitation wavelength of 448 nm. The green line shows the luminescence spectrum of a powder sample of $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ recorded at an excitation wavelength of 400 nm. The blue line represents the excitation spectrum of the powder sample of $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ monitored at $\lambda_{\text{em}} = 590$ nm (based on raw data maximum). (b) It shows the color point of the $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ single-crystals in the CIE-xy color space.⁴⁸

under daylight and is stable under atmospheric conditions for several months.

The thermal quenching behavior (TQ) of $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ was examined on a powder sample (see Fig. 7).

As can be seen in Fig. 7, the relative integral photoluminescence intensity decreases with increasing temperature and drops to 66% of the initial value at a temperature of 225 °C. At a typical operating temperature of 125 °C for a p-LED, the relative integral intensity is still above 85%. Nevertheless, QE and TQ would have to be improved for any industrial use.

Since the compound shows only one emission band, it would indicate that only one position in the structure is suit-

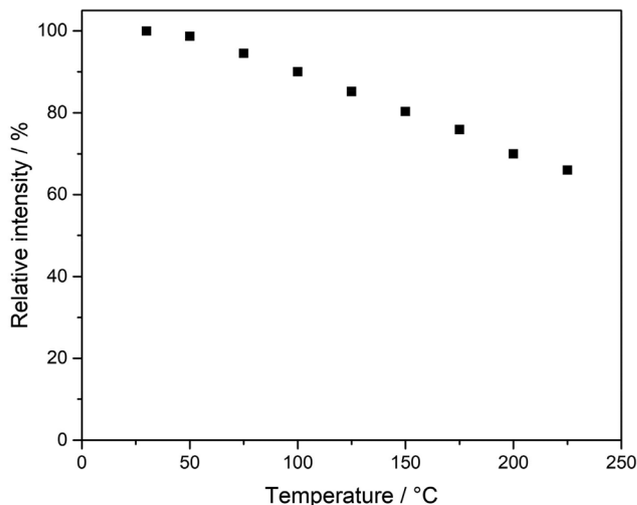


Fig. 7 Representation of the thermal quenching behavior of $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ as the relative integral photoluminescence intensity to that at 25 °C measured in steps of 25 °C up to 225 °C.

able for incorporating the activator ion Eu^{2+} . One possible position would be the Li3 site, which forms an octahedral coordination to two O^{2-} anions with Li–O distances of 205.0(2) pm and to four N^{3-} anions with Li–N distances between 218.9(2) and 248.94(5) pm resulting in a polyhedron size of 14.19 Å³. Although the distances are too short regarding typical Eu–O/N coordination (Eu–O distances range from 240 to 300 pm and Eu–N distances range from 260 to 325 pm),^{49–51} local spatial distortion or enlargement of the octahedron could occur in contrast to the average structure. To maintain charge neutrality, the inclusion of a divalent Eu^{2+} cation may be compensated by Li-vacancies in the surroundings or by local O/N substitution. In comparison with the LiO_2N_4 octahedron of the Li3 site, the LiO_4N_2 octahedron including the dynamic positional disordered Li4 and Li4A sites is slightly larger and would also be a possible position to host the Eu^{2+} cation. The Li4/Li4A site is coordinated octahedrally by four O^{2-} anions with Li–O distances between 200.58(4) and 292.0(3) pm and two N^{3-} anions with Li–N distances between 222.3(2) and 255.8(8) pm resulting in a polyhedron size of 16.73 Å³. Due to the two short Li–O distances, this would also lead to an enlargement or spatial distortion of the octahedron. Compared to compounds such as $\text{Li}_2\text{CaSi}_2\text{N}_4:\text{Eu}^{2+}$,⁵² $\text{Mg}_3\text{GaN}_3:\text{Eu}^{2+}$,⁵³ and $\text{Li-}\alpha\text{-SiAlON}:\text{Eu}^{2+}$,⁵⁴ the polyhedron size and Li–O/N distances of the Li3 and Li4/Li4A sites in $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ are in a similar range and it seems possible that one of them hosts the activator ion Eu^{2+} . In the former compound, there are two crystallographically distinguishable Ca sites, which are octahedrally coordinated by six N^{3-} anions and the compound exhibits an emission maximum at $\lambda_{\text{max}} = 583\text{--}585$ nm. The Ca1 site has Ca1–N distances between 248.9(3) and 250.9(3) pm resulting in a polyhedron size of 16.54 Å³ and the Ca2 site with a Ca2–N distances of 258.6(3) pm has a polyhedron size of 18.95 Å³, whereby the author supposed that both Ca sites are suitable to host the activator ion Eu^{2+} . In $\text{Mg}_3\text{GaN}_3:\text{Eu}^{2+}$, it was assumed



that the activator ion Eu^{2+} is located in octahedral voids, where the possible observed Eu–N distances are between 232.0 and 248.9 pm resulting in an estimated polyhedron sizes between 14.5 and 16.5 Å³ exhibiting an emission maximum at 578 nm with a fwhm of 132 nm. In Eu^{2+} doped Li- α -SiAlON, it was supposed that the activator ion Eu^{2+} occupies a Li site, which is sevenfold coordinated by $\text{N}^{3-}/\text{O}^{2-}$ anions with Li–O/N distances ranging from 205.2 to 276.7 pm. Depending on the Eu^{2+} concentration, the compound Li- α -SiAlON exhibits an emission maximum in the range $\lambda_{\text{max}} = 563\text{--}586$ nm.

It would also be possible that the activator ion Eu^{2+} is situated on the Li2 site. As illustrated in Fig. 8, a lithium channel is formed in the structure of Li_3SiNO_2 along the crystallographic *c*-axis, which consists of an edge-shared LiO_4 tetrahedral band (Li2 site) and corresponds to band Y. This band Y is connected to two unbranched *zweier* single chains of SiO_2N_2 tetrahedra by corner-sharing *via* the O^{2-} anion, respectively. Due to this special arrangement of the unbranched *zweier* single chains of SiO_2N_2 tetrahedra, there would also be two N^{3-} anions in close spatial proximity to the Li2 site, whose Li–N distance would be 314.9 pm. Although this Li–N distance is too large for a Li^+ cation's first coordination sphere, it would be a permissible Eu–N distance for the activator ion Eu^{2+} . Considering these two N^{3-} anions, an octahedral coordination would also result for the Eu^{2+} cation on the Li2 site and is consequently a possible position hosting the Eu^{2+} cation. Due to the additional tetrahedral coordination of the O^{2-} anions of the Li2 site, this would form a distorted octahedron with a polyhedron size of 12.52 Å³ and, due to the short Li–O distances, there would also be an enlargement or spatial distortion of the octahedron. The incorporation of the activator ion Eu^{2+} into lithium channels or domains has already been observed in other compounds known from the literature, such as $\text{RbKLi}_2[\text{Li}_3\text{SiO}_4]_4:\text{Eu}^{2+}$,^{7,23} $\text{Cs}_{4-x-y-z}\text{Rb}_x\text{Na}_y\text{Li}_z[\text{Li}_3\text{SiO}_4]_4:\text{Eu}^{2+}$,²⁴ and $\text{LiK}_7[\text{Li}_3\text{SiO}_4]_8:\text{Eu}^{2+}$.²⁰ However, it should be considered that in these compounds the Li^+ cations form an approximately square planar coordination to four O^{2-} anions instead of a tetrahedral one.

Due to the special arrangement of the various polyhedra, unoccupied distorted octahedral voids are formed in the structure, which could also be a possible position hosting the activator ion Eu^{2+} . A similar behavior was observed in the compound $\text{Mg}_3\text{GaN}_3:\text{Eu}^{2+}$, as mentioned before. The unoccupied distorted octahedral void is coordinated by five O^{2-} anions and one N^{3-} anion, resulting in a MO_5N octahedron with an O1–O2 distance of 465.1, an O2–O2 distance of 511.9, and a N1–O2 distance of 430.6 pm (see Fig. 9). Comparing the anion–anion distances with those of the LiO_2N_4 octahedron (two times a N1–O1 distances of 421.1, and a N1–N1 distance of 497.2 pm) and the dynamic positional disordered LiO_4N_2 octahedron (two times a O2–N1 distances of 501.8, and an O1–O1 distance of 400.9 pm), it is found to be similar in size to the last one and the MO_5N octahedron would result in a polyhedron size of 16.46 Å³. The polyhedron size of the MO_5N octahedron was calculated by placing a dummy atom in the center of the octahedral void without considering any structural relaxation. Considering these anion–anion distances of the MO_5N octahedron, a spatial distortion or enlargement of the octahedral void could occur, when an Eu^{2+} cation is incorporated.

All four positions could act as doping sites for the activator ion Eu^{2+} that explain the observed single band emission. Presumably, however, it is assumable that only one of these sites is preferred, otherwise a split or broad emission profile should be observed. It seems most likely that the activator ion Eu^{2+} is incorporated into the Li2 site building up the lithium channel or in the center of the LiO_4N_2 octahedron including the dynamic positional disordered Li4 and Li4A sites. One possible reason for the latter is that, compared to the octahedrally coordinated Li3 site, it has larger cation–anion distances, which partially correspond to the observed Eu–N and Eu–O distances, therefore resulting in a larger polyhedron size. In the case of the Li2 site, an increased electron density could be observed at this site in the Eu^{2+} doped crystals, which roughly corresponds to the 2 mol% activator ion Eu^{2+} concentration used in the syntheses. In addition, the residual electron density near this site was also found to be aligned towards the

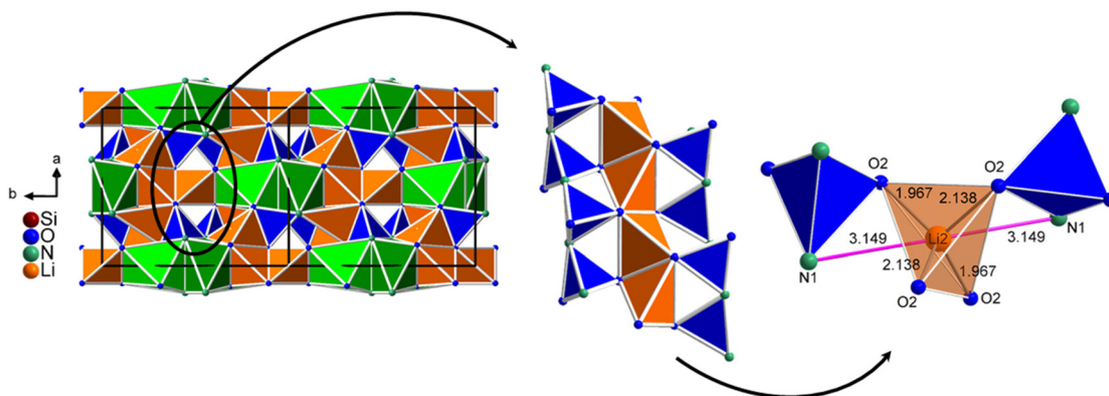


Fig. 8 On the left, the position of one of the lithium channels in the structure of Li_3SiNO_2 is shown. In the middle, the connection of two unbranched *zweier* single chains of SiO_2N_2 tetrahedra (blue) with the LiO_4 tetrahedra of the Li2 site (orange, band Y) is represented. On the right, the Li2 site including the possible octahedral coordination to four O^{2-} and two N^{3-} anions is illustrated.



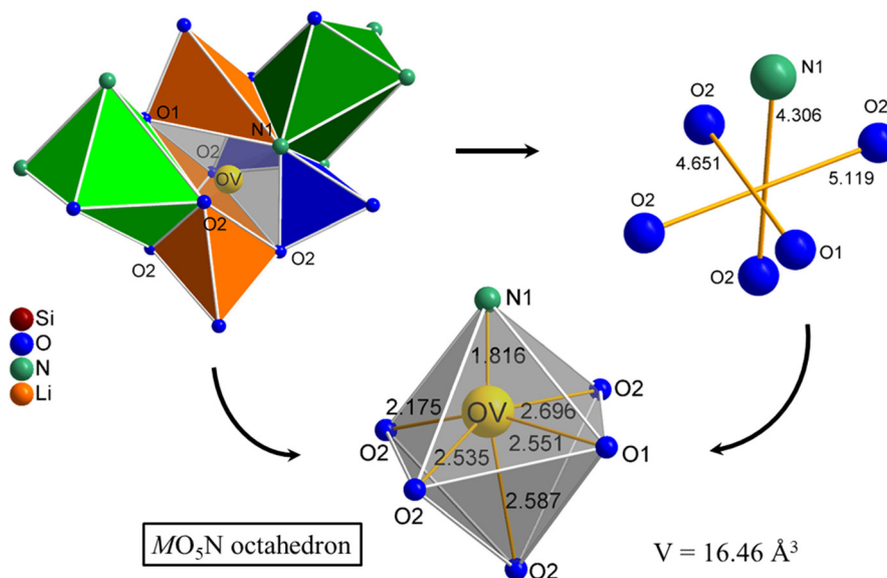


Fig. 9 On the left, the unoccupied distorted octahedral void (OV) in the structure of Li_3SiNO_2 is shown by the labeled O^{2-} anions and the N^{3-} anion, which is constructed by the special arrangement of the various polyhedra. On the right, the corresponding anions of the unoccupied MO_5N octahedron are shown including the specific anion–anion distances. In the middle, the resulting MO_5N octahedron (gray) with hypothetical M–O/N distances including the polyhedron size is illustrated. The LiO_4 and LiO_3N tetrahedra are displayed in orange, the LiO_2N_4 octahedron is displayed in green, the SiO_2N_2 tetrahedra are displayed in blue and the LiO_4N_2 octahedron including the dynamic positional disordered Li4 and Li4A sites is displayed in bright green.

N^{3-} anions and the resulting distance to the N^{3-} anions is around 250 pm, which is approximately in the range of Eu–N distances. Considering the unoccupied distorted octahedral void as a possible position hosting the Eu^{2+} cation, it should be possible to detect low electron density at this position in the single-crystal data. However, no electron density could be observed around this position and consequently the unoccupied distorted octahedral void is presumably not occupied by the activator ion Eu^{2+} . Based on SCXRD, an unambiguous localization of the luminescence center in $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ was not possible.

Compared to other luminescent lithium (oxo)nitride silicates such as $\text{LiSi}_2\text{N}_3:\text{Eu}^{2+}$ (ref. 55) or $\text{LiSiON}:\text{Eu}^{2+,3}$ the Li–O/N distances of $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ are in a similar range. $\text{LiSi}_2\text{N}_3:\text{Eu}^{2+}$

crystallizes in the orthorhombic space group $Cmc2_1$ and has one Li site tetrahedrally coordinated by four N^{3-} anions with Li–N distances between 202(2) and 264(2) pm. The compound exhibits broadband emission with a maximum at $\lambda_{\text{max}} = 572\text{--}584$ nm. The compound $\text{LiSiON}:\text{Eu}^{2+}$, which crystallizes in the orthorhombic space group $Pca2_1$, also has one Li site. The Li^+ cation has a coordination sphere to three O^{2-} and one N^{3-} anions, whose Li–O distances are between 191.8 and 206.8 pm and the Li–N distance is 225.2 pm resulting in an emission maximum at $\lambda_{\text{max}} = 426\text{--}478$ nm (fwhm = 142–193 nm). In the publication of $\text{LiSi}_2\text{N}_3:\text{Eu}^{2+}$ and $\text{LiSiON}:\text{Eu}^{2+}$, no possible positions for the activator ion Eu^{2+} were discussed. In relation to $\text{LiSi}_2\text{N}_3:\text{Eu}^{2+}$ and $\text{LiSiON}:\text{Eu}^{2+}$, a blue shift in luminescence due to the partial substitution of N^{3-} anions by O^{2-} anions

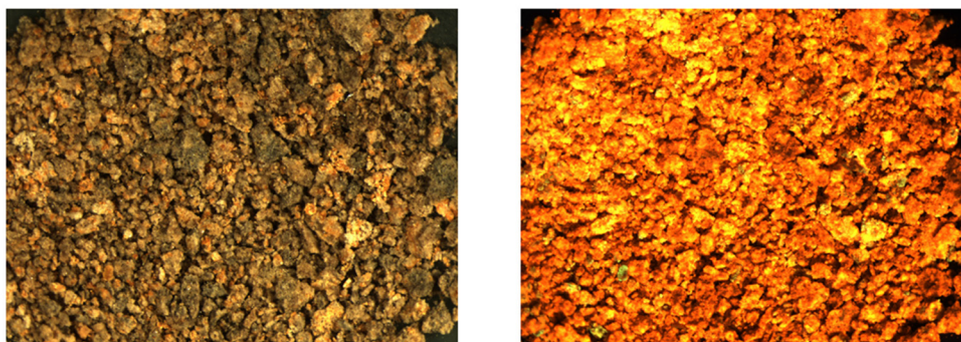


Fig. 10 Left: optical appearance of the 2 mol% Eu^{2+} -doped compound Li_3SiNO_2 under daylight. Right: optical appearance of the 2 mol% Eu^{2+} -doped compound Li_3SiNO_2 under UV-light (365 nm).



cannot be observed for $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$. Since the structures have fundamental differences regarding the coordination of the Li^+ cations, a comparison of these compounds is not possible. Other compounds such as $\text{Sr}[\text{LiAl}_3\text{N}_4]:\text{Eu}^{2+}$ (SLA: Eu^{2+}) ($\lambda_{\text{max}} = 654 \text{ nm}$; $\text{fwhm} = 50 \text{ nm}$)⁵⁶ and $\text{Sr}[\text{Li}_2\text{Al}_2\text{O}_2\text{N}_2]:\text{Eu}^{2+}$ (SALON: Eu^{2+}) ($\lambda_{\text{max}} = 614 \text{ nm}$; $\text{fwhm} = 48 \text{ nm}$),⁵⁷ for example, have comparable structures, where a partial substitution of N^{3-} anions by O^{2-} anions leads to a blue shift in the observed emission bands. The nephelauxetic effect is stronger for the activator ion Eu^{2+} when it is coordinated by N^{3-} anions rather than by O^{2-} anions due to the higher formal charge of N^{3-} anions. Consequently it leads to a lowering of the 5d energy levels resulting in a red-shift of the Eu^{2+} emissions induced from the d to f transitions.⁵⁸

Conclusion

In this study, a new orange-emitting phosphor $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ has been successfully prepared by a solid-state reaction in tantalum ampules. The compound can be excited effectively by near-UV to blue light and the single-crystals of $\text{Li}_3\text{SiNO}_2:\text{Eu}^{2+}$ exhibit an emission maximum at $\lambda_{\text{max}} = 601 \text{ nm}$ with a full width at half maximum (fwhm) of 90 nm (0.32 eV , 2597 cm^{-1}). The crystal structure with its 3D-network can be described as a condensed layered structure consisting of two different layers A and B, which arrange alternately along the crystallographic a -axis. Although layer A has structural similarities to other compounds such as $\text{Na}_2\text{Mn}_2\text{S}_3$ and $\text{Na}_6\text{In}_2\text{S}_6$ and *zwei* single chains can basically also be found in phosphors such as $\text{MSi}_2\text{O}_2\text{N}_2$ ($M = \text{Sr}, \text{Ba}, \text{Eu}$),^{26,59–73} the structural motif and connection pattern in layer B are still unknown. The latter consists of pure Li^+ cations, which build up a dense network of corner- and edge-shared tetrahedra and octahedra with the O^{2-} and N^{3-} anions. Furthermore, we discussed four possible positions in the structure hosting the activator ion Eu^{2+} , which include two different octahedral coordination spheres, one octahedral void spanned by a special arrangement of different polyhedra and a tetrahedrally coordinated Li-channel. In comparison with other compounds known from the literature such as α - and β -SiALON,^{74–76} we see that heavy cations are not necessarily required as counterions in the structure that can be substituted by the activator ion Eu^{2+} resulting in interesting luminescence properties. Although the new phosphor can be synthesized by low-cost educts and shows relatively good stability against moisture and atmospheric conditions, a further optimization of the material's optical properties must be achieved for possible applications as a phosphor material for example in p-LEDs.

Experimental section and theoretical methods

Synthesis

Since it is crucial to maintain control over the amount of oxygen for synthesizing oxonitrides, all preparations were

carried out using an inert gas (Ar 5.0, Messer Austria GmbH) filled glovebox (MBraun, $\text{O}_2 < 1 \text{ ppm}$, $\text{H}_2\text{O} < 1 \text{ ppm}$). The starting materials Si_3N_4 (UBE SN-E10, >99%) and Li_2O (Sigma Aldrich, >97%) with a stoichiometric ratio of 1 : 4 were weighed and mixed with 5 wt% LiF (Sigma Aldrich, >99.99%) as a flux and 2mol% EuF_2 (Alfa Aesar, >99.9%) using an agate mortar. Afterwards, the powder mixture was filled into a tantalum ampule and sealed *via* arc welding under an argon atmosphere. The reaction vessel was then placed inside a silica-glass ampule containing a 400 mbar inert gas atmosphere (Ar 5.0, Messer Austria GmbH). The temperature profile of the synthesis consists of a heating ramp with $4 \text{ }^\circ\text{C min}^{-1}$ up to $960 \text{ }^\circ\text{C}$, a dwelling time of 4 h, and subsequently a cooling ramp of $-0.3 \text{ }^\circ\text{C min}^{-1}$ to $300 \text{ }^\circ\text{C}$. Once cooled down to room temperature, the ampules could be opened and the samples were further examined under atmospheric conditions without detectable decomposition within several months. The product shows intense yellow to orange luminescence under UV-light (see Fig. 10).

Single-crystal X-ray diffraction

Under a polarization microscope, suitable single-crystals of Li_3SiNO_2 were isolated and subsequently investigated with a Bruker D8 Quest diffractometer (Mo- $K\alpha$ radiation, $\lambda = 0.7107 \text{ \AA}$) equipped with a Photon 300 CMOS detector. The programs Saint⁷⁷ and Sadabs⁷⁸ were used for data processing and multi-scan absorption correction. For the structure solution and parameter refinement, ShelXS⁷⁹ using Direct Methods and ShelXL⁸⁰ using the least squares method, both implemented in the program WinGX,⁸¹ were applied. The structural data were standardized employing Structure Tidy⁸² as implemented in Platon (version 170613).⁸³ Further details on the crystal structure investigation of the single-crystal of Li_3SiNO_2 can be obtained from The Cambridge Crystallographic Data Centre *via* <https://www.ccdc.cam.ac.uk/structures/> under the deposition number 2208257.†

Powder X-ray diffraction

Powder X-ray diffraction (PXRD), equipped with a STOE Stadi P powder diffractometer with Ge(111)-monochromatized Mo- $K\alpha_1$ ($\lambda = 70.93 \text{ pm}$) radiation and a Mythen 1K detector operated in Debye–Scherrer geometry across a 2θ range of 2° – 40° , was used for phase analysis. The experimental powder pattern was compared with the calculated powder pattern derived from the single-crystal data of Li_3SiNO_2 . The secondary phase identification was carried out with the ICDD PDF-2 database and the program Topas 4.2⁸⁴ was applied for Rietveld refinements.

Luminescence

The emission spectra of representatives from Li_3SiNO_2 were recorded by a setup of a blue laser diode ($\lambda = 448 \text{ nm}$, THORLABS, Newton, NJ, USA) in combination with a CCD detector (AVA AvaSpec 2048, AVANTES, Apeldoorn, The Netherlands). A tungsten–halogen calibration lamp was used for the previous spectral radiance calibration of the setup. The software AVA AvaSoft (version 7) was used for the data prepa-



ration. The analysis of the luminescence properties of powder samples was carried out using a Fluoromax 4 spectrophotometer (Horiba). The emission spectrum was recorded in the wavelength range between 470 and 750 nm (step size 1 nm) using an excitation wavelength of 400 nm. The same method was used to record the excitation spectra monitored at the corresponding maximum intensity and for the thermal-quenching analysis.

Author contributions

All the authors have accepted responsibility for the entire content of this submitted manuscript and approved submission.

Conflicts of interest

The authors declare no conflict of interest regarding this article.

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