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Borate polyanions tuning persistent luminescence in Eu and Dy co-doped strontium aluminate[†]

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Although the longest duration of persistent luminescence in Eu and Dy co-doped strontium aluminate (S4A7ED) is obtained by processing with 30 mol% B_2O_3 (S4A7EDB), this composition does not contain the highest amount of the long persistence phase. To elucidate how the structural arrangement of atoms in S4A7EDB induced by synthesis using 30–40 mol% B_2O_3 yields the longest persistence, the crystal environment of B in S4A7ED is evaluated by using micro-Raman and FTIR spectroscopy. Polyatomic vibration modes containing B–O are revealed, in forms of metaborate and other borate polyanions. Based on these results, we propose a structural model for S4A7EDB, in which B introduces non-bridging oxygen, which offers excess negative charge and enlarged adjacent Sr sites, both of which enable Eu and Dy dopant clustering that yields extreme persistent luminescence.

Introduction

Eu^{2+} , Dy^{3+} and B co-doped $Sr_4Al_{14}O_{25}$ (S4A7EDB) is well known to exhibit persistent luminescence. Boron oxide, initially used as a sintering flux agent to facilitate synthesis at lower temperatures and stoichiometric grain growth, was noted to extend persistent luminescence in Eu and Dy co-doped $SrAl_2O_4$ (SAED) and S4A7ED from *ca.* 10 minutes to longer than 10 h.^{1–3} Understanding the exact role of B in forming the crystal structure has motivated investigations into its location and crystal environment.^{4–6} Nag had reported the existence of BO_4 units, which were revealed by NMR and FTIR,⁵ and which was confirmed by spatially resolved analysis of the B–K ionization edge energy loss near-edge structure (ELNES).⁴ Boron incorporation in RE co-doped S4A7 was also associated with a blue shift in emission at 400 nm, which was attributed to lattice shrinkage.⁶ Additional peaks were observed in the photoluminescence (PL) and thermoluminescence (TL) spectra, which were interpreted as additional traps due to the incorporation of B into the SAED crystal structure,⁷ while B modification of trap depth and density has been reported in other persistent afterglow compounds as well.^{8,9} To explain the role of boron,

theoretical studies suggested the clustering of the point defects, Eu^{+2} , Dy^{+3} , B, and oxygen vacancies.¹⁰ In nano-S4A7 produced by a microemulsion method, 15–20 mol% has been reported as the optimum boric acid concentration for the maximum intensity and duration of the persistent luminescence.¹¹

B_2O_3 is known to serve as a network former in oxide glasses, where it forms polyhedral units due to being covalently bonded with charge-screening oxygen.^{12,13} Each oxygen is saturated, *i.e.*, bonded to 2 cations. Network modifiers, on the other hand, contribute non-bridging oxygens to the network. Because they do not completely link the maximum number of polyhedral units—*i.e.*, 2 units per oxygen—their presence lowers the network connectivity, resulting in more open networks. Alkaline oxides and alkali earth oxides are common modifiers, and the presence of their cations provides local electrical charge neutrality.^{14,15} Network intermediates act as glass formers or modifiers, depending on the local network structure and composition.^{12,14} The structure of boron oxide consists of corner-sharing triangular polyhedra. Vitreous boron oxide has planar six-membered boroxyl rings containing trigonal uncharged $[BO_3]$, where \textcircled{O} indicates bridging oxygen, and O indicates non-bridging oxygen.¹⁶ Combinations of basic borate structural units form superstructural units, such as borate rings, diborates, metaborate ring-chains, pyroborate dimers, *etc.*^{13,17,18}

Cation additions can change the oxide network character. For example, B_2O_3 incorporation into alumina glass induces changes in the types and connectivity of borate and aluminate structural units. Boron oxide functions as both a former and modifier in alumino-borate (Al_2O_3 – B_2O_3) networks, depending on their relative concentration ratio.¹⁹

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The change in the structural function of boron oxide is commonly referred to as the boron anomaly. With the addition of modifier oxides, planar trigonal charge-neutral $[\text{BO}_3]$ triangles initially transform to negatively charged, tetrahedral $[\text{BO}_4]^-$ of 3-D character. Upon increasing the amount of modifier added to Al_2O_3 – B_2O_3 , the number of non-bridging oxygens increases, breaking up the oxide network connectivity.^{16,19,20} Above a critical modifier concentration, the coordination number of boron converts back from four to three, in the form of complex triangular moieties—the ones with non-bridging oxygen on meta $[\text{BO}_2\text{O}]^-$, pyro $[\text{BO}_2\text{O}_2]^{2-}$ and finally ortho $[\text{BO}_3]^{3-}$ units.^{13,21} Thus, while boron effectively densifies the alumina polyhedral network at low modifier concentrations, the increase in non-bridging oxygen yields increased free volume in the network at elevated modifier concentrations. The structural connectivity of borate-containing glass can be quantified by the relative amount of 3- and 4-coordinated boron, with the relative ratio being defined as $N4 = (\text{B}4/(\text{B}3 + \text{B}4))$. This metric facilitates the correlation of the structure with optical and thermal properties.

The role of oxides in the complex strontium aluminate SrO – Al_2O_3 – B_2O_3 network depends on the composition. Al^{3+} ions can be either a network former in Al_2O_3 , forming AlO_4 tetrahedra, or a network modifier, by contributing non-bridging oxygen in the formation of AlO_6 octahedra.^{19,22,23} Sr^{2+} ions are network modifiers in SrO , where they break up the network and provide charge compensation in interstitial sites.²⁴ B_2O_3 can provide charge compensation to stabilize Al polyhedra.¹⁹

The crystal structure of $\text{Sr}_4\text{Al}_{14}\text{O}_{25}$ is shown in Fig. 1a. $\text{Sr}_4\text{Al}_{14}\text{O}_{25}$ crystallizes in the orthorhombic space group $Pm\bar{m}$ with unit cell parameters of $a = 24.785$ Å, $b = 8.487$ Å, and $c = 4.886$ Å; $V = 1027.772$ Å³; and $\alpha = \beta = \gamma = 90^\circ$. A network of corner- and edge-sharing alumina polyhedra defines a scaffold,

i.e., chains of octahedra (Fig. 1b), sharing a single edge, defines the scaffold columns, while chains of corner-sharing tetrahedra bridge the columns (Fig. 1c). Sr^{2+} cations provide charge compensation in large interstitial cavities in two crystallographically non-equivalent sites: $\text{Sr}1$ coordinated by 10 oxygen and of Wyckoff symmetry 4j and $\text{Sr}2$ coordinated by 7 oxygen and of Wyckoff symmetry 4i (Fig. 1d).

Previous work in our group revealed the significant and multi-functional role of boron in the structural evolution of S4A7EDB during Pechini processing. Boron was associated with RE incorporation into adjacent Sr^{2+} sites in S4A7,²⁵ as well as with a more uniform distribution of RE dopants in Sr^{2+} sites across the microstructure by nanoCL analysis.³ The boron-rich intergranular phase enabled tuning of the RE dopant concentration in S4A7 grains to reach their solubility limits, to avoid concentration quenching.⁴ In addition, trigonally coordinated boron was observed in the intergranular phase, while it was primarily of tetrahedral coordination in the S4A7 grains.⁴ We also observed that processing with boron oxide not only lowered the temperature needed for the crystal structure of S4A7 to form, but also modified the diffusion kinetics of Eu and Dy in playing a role in extending the afterglow persistence in S4A7EDB.^{3,4,25} Finally, the amount of boron oxide used in Pechini processing of S4A7EDB influenced the duration of persistent luminescence, with the longest in compounds containing 30–40 mol% B_2O_3 . Interestingly, the total amount of the long afterglow phase was not at the maximum when this amount of B_2O_3 is used.²⁶

In previous studies, vibration spectroscopy techniques have proved useful in determining the structure and local environment in polymorphic crystal structures and glassy ones, as well as those of borates. The Raman modes of SrAl_2O_4 (SA) have also been considered in several studies.^{27–29} Meanwhile, crystal

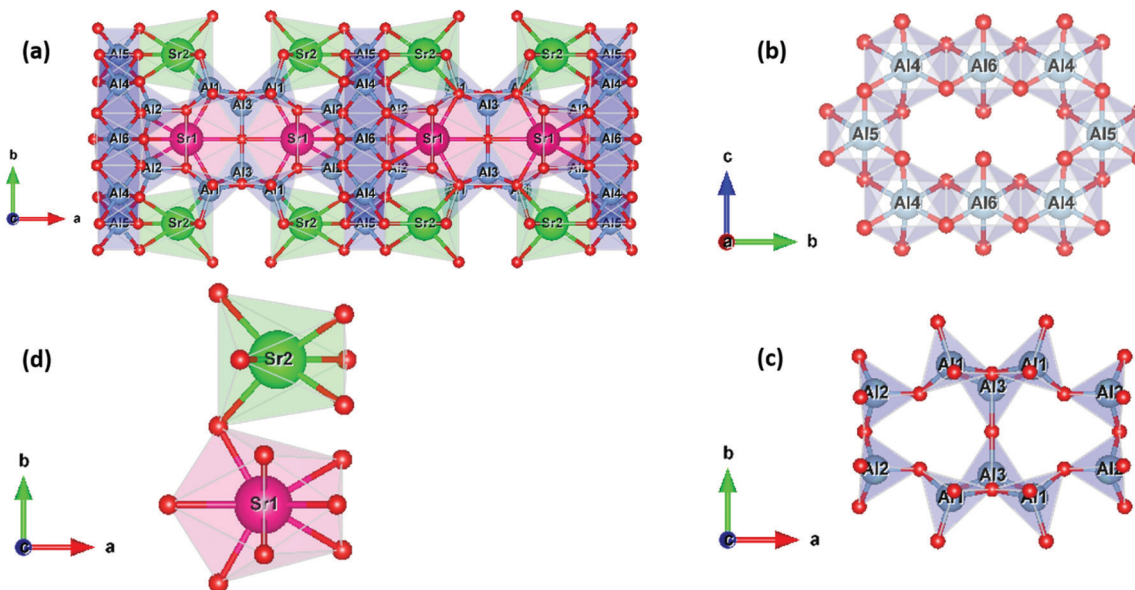


Fig. 1 (a) Components of the $\text{Sr}_4\text{Al}_{14}\text{O}_{25}$ unit cell viewed along the [001]; (b) 6 oxygen-coordinated alumina scaffold; (c) 4 oxygen-coordinated alumina scaffold; (d) the coordination polyhedra of the Sr (1) site (Wyckoff 4j) and Sr (2) site (Wyckoff 4i).



structure changes due to dopant and impurity incorporation have been studied with Raman spectroscopy.³⁰ Raman and infrared spectroscopy are commonly used as complementary techniques for elucidating the local structure in modified glass and crystalline structures.^{31–37} Keiner and co-workers demonstrated the use of micro-Raman and FTIR to correlate spatially resolved information about molecular and network structures.³⁸

In this work, to understand how the B_2O_3 content tunes the structure and luminescence behavior of S4A7ED, we synthesized S4A7EDB by Pechini processing using different amounts of B_2O_3 ; analyzed the phase composition and changes in the atomic arrangements in the crystal structure by XRD, the crystal environment of B by micro-Raman and FTIR spectroscopy; and measured the duration of persistent luminescence by afterglow decay.

Experimental methods

$\text{Sr}_4\text{Al}_{14}\text{O}_{25}$ compound doped with 1 at % Eu, 1 at % Dy and $0 \leq x \leq 60$ mol% B_2O_3 was synthesized by a sol-gel Pechini process as described by Inan Akmehmehmet *et al.*³ The raw materials were aluminum nitrate nonahydrate $[\text{Al}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}]$, strontium nitrate anhydrous $[\text{Sr}(\text{NO}_3)_2]$, europium nitrate hexahydrate $[\text{Eu}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}]$, dysprosium nitrate pentahydrate $[\text{Dy}(\text{NO}_3)_3 \cdot 5\text{H}_2\text{O}]$, boron oxide (B_2O_3), chelating agent citric acid monohydrate ($\text{C}_6\text{H}_8\text{O}_7$), and the gelling agent ethylene glycol ($\text{C}_2\text{H}_6\text{O}_2$). The amorphous powder precursors were calcined at 600 °C for 6 h and then 1150 °C for 10 hours in air. The mixtures were then ground and reduced at 1150 °C for 4 h under a reducing atmosphere (96% Ar, 4% H_2).

The finely ground powder is used in all measurements. The afterglow persistence of S4A7EDxB samples ($0 \leq x \leq 60$ mol% B_2O_3) was collected by using a custom-built optical bench photoluminescence system consisting of a photomultiplier tube (H7421; Hamamatsu Photonics Deutschland GmbH, Herrsching, Germany) coupled with a photon counting unit (C-8855, Hamamatsu Photonics Deutschland GmbH), after illuminating the powder with a 365 nm UV light source for 10 minutes, and then recorded for up to 14 h.

Crystal structure analysis and phase identification were performed by XRD (Bruker AXS, D8 Advance: Cu-K α source ($\lambda = 1.54056 \text{ \AA}$), scanning rate 1 s per step, 0.02° increment, over a 2 θ range of 10° to 90°). Initial phase analysis of the diffraction patterns performed by search/match from the International Centre for Diffraction Data (ICDD) database in the DIFRACT.-SUITE EVA software (Bruker AXS, Germany). Match ICDD files: Orthorhombic $\text{Sr}_4\text{Al}_{14}\text{O}_{25}$ (S4A7) PDF No. 74-1810, monoclinic SrAl_4O_7 (SA2) PDF No. 70-1479, hexagonal $\text{SrAl}_{12}\text{O}_{19}$ (SA6) PDF No. 70-0947, monoclinic SrAl_2O_4 (SA) PDF No. 74-0794, orthorhombic SrB_2O_4 (SB) PDF No. 84-2175 and orthorhombic $\text{Sr}_2\text{B}_2\text{Al}_2\text{O}_8$ (SAB) PDF No. 77-0402. The DiffracPlus TOPAS 4.2 software package (Bruker AXS, Karlsruhe, Germany)³⁹ was used for the structural analysis by the X-ray Rietveld refinement method with the corresponding crystallographic information

file (CIF) of the identified phases collected from the Inorganic Crystal Structure Database (ICSD; FIZ Karlsruhe 2019, version 4.2.0)⁴⁰ and FindIT crystal database software. VESTA⁴¹ was used for viewing crystal structures—reference ones and refined data—, and DIFRACT.SUITE EVA (Bruker AXS, Germany) was used for determining the percent crystallinity.

Fourier transform infrared (FTIR) spectra were collected using a Nicolet IS10 FT-IR spectrometer (Thermo Scientific, USA) in ATR mode in the range of 4000 to 525 cm^{-1} with a resolution of 4 cm^{-1} at 128 scans. Spectral processing and analysis were performed using the program Omnic (Thermo Scientific) program⁴² and the software program Fityk[®] version 1.3.1 for spectral deconvolution.⁴³

Micro-Raman analysis was performed on polished sintered ceramic pellets with using a Raman system (Renishaw inVia) mounted on a confocal microscope: 0.05 mW diode laser, $\lambda_{\text{ex}} = 532 \text{ nm}$, 1 s exposure, 1 cm^{-1} spectral resolution; 856 nm diameter focused laser spot at 50 \times ; system calibrated to 520 nm^{-1} of Si. A representative dataset was ensured *via* measuring at 5 different locations. The manufacturer provided WiRE 3.4 software application was used for data acquisition and analysis by multivariate curve resolution-alternating least squares (MCR-ALS) method using Empty Modelling[™]. The baseline correction was performed with a 5th order polynomial, and spectral deconvolution analysis was performed using Fityk[®].⁴³

Results

The afterglow decay lifetime of S4A7EDxB, where $0 \leq x \leq 60$ mol% B_2O_3 , was recorded for 14 hours under ambient conditions after being illuminated with a UV light source ($\lambda_{\text{ex}} = 365 \text{ nm}$) for 10 minutes, and the decay curves are summarized in Fig. 2a. All S4A7 powders exhibited persistent luminescence, regardless of the B_2O_3 content. The longest persistence luminescence was observed in $x = 30$ mol% B_2O_3 and 40 mol% B_2O_3 compounds. Fig. 2b shows the afterglow decay curves for S4A7EDxB ($30 \leq x \leq 40$ mol% B_2O_3). The two-slope behavior in the decay curves indicated 2 independent detrapping mechanisms, corresponding to τ_1 and τ_2 , respectively, which was obtained from the decay-curve fitting equation:

$$I = \alpha_1 \exp\left(\frac{-t}{\tau_1}\right) + \alpha_2 \exp\left(\frac{-t}{\tau_2}\right)$$

α_1 and α_2 are constants, τ_1 and τ_2 are the decay lifetimes times, as shown in Fig. 2c. The two different lifetimes, τ_1 and τ_2 , indicate the rapid and slow components of radiative relaxation. The S4A7EDxB compound with the $x = 30$ mol% B_2O_3 doped composition had $\tau_1 = 1.14 \text{ h}$, $\tau_2 = 5.8 \text{ h}$, while the $x = 40$ mol% B_2O_3 doped one showed $\tau_1 = 1.21 \text{ h}$ and $\tau_2 = 6.13 \text{ h}$.

To determine the phase composition of S4A7EDxB, where $0 \leq x \leq 60$ mol% B_2O_3 , all samples were analyzed using XRD (Fig. 3). Calibration of the instrument was performed using a corundum standard. The evolution of the main (421) diffraction peak with the B_2O_3 content is shown in Fig. 3b, across the entire composition range studied, with an emphasis on peak

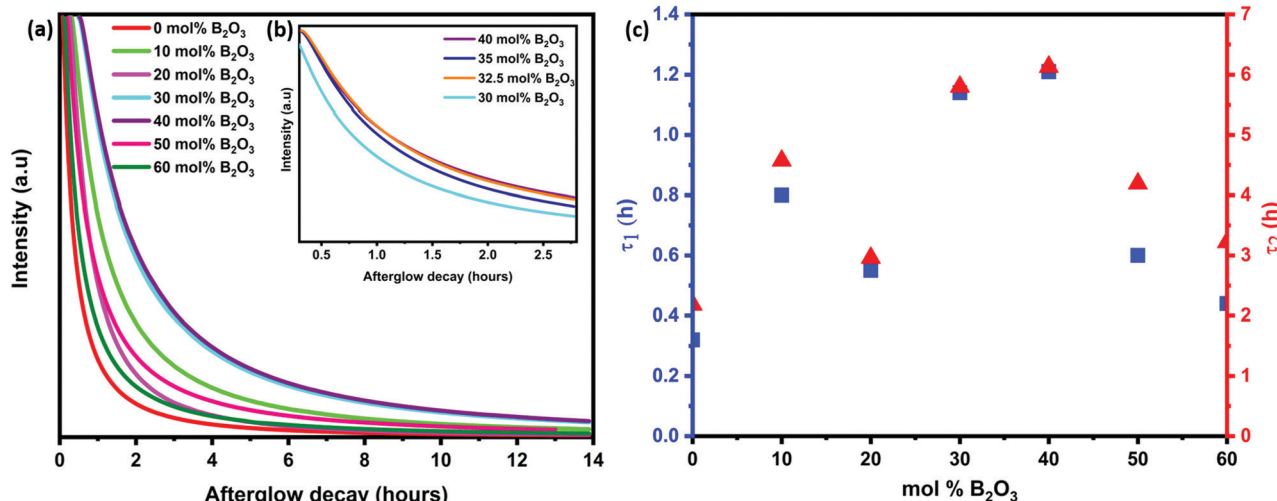


Fig. 2 (a) Afterglow decay curves for S4A7EDxB ($0 \leq x \leq 60 \text{ mol\% B}_2\text{O}_3$); (b) afterglow decay curves for S4A7EDxB ($30 \leq x \leq 40 \text{ mol\% B}_2\text{O}_3$); (c) decay time constant analysis of S4A7EDxB, where $0 \leq x \leq 60 \text{ mol\% B}_2\text{O}_3$ samples: τ_1 for relaxation from shallow traps and τ_2 for relaxation from deep traps.

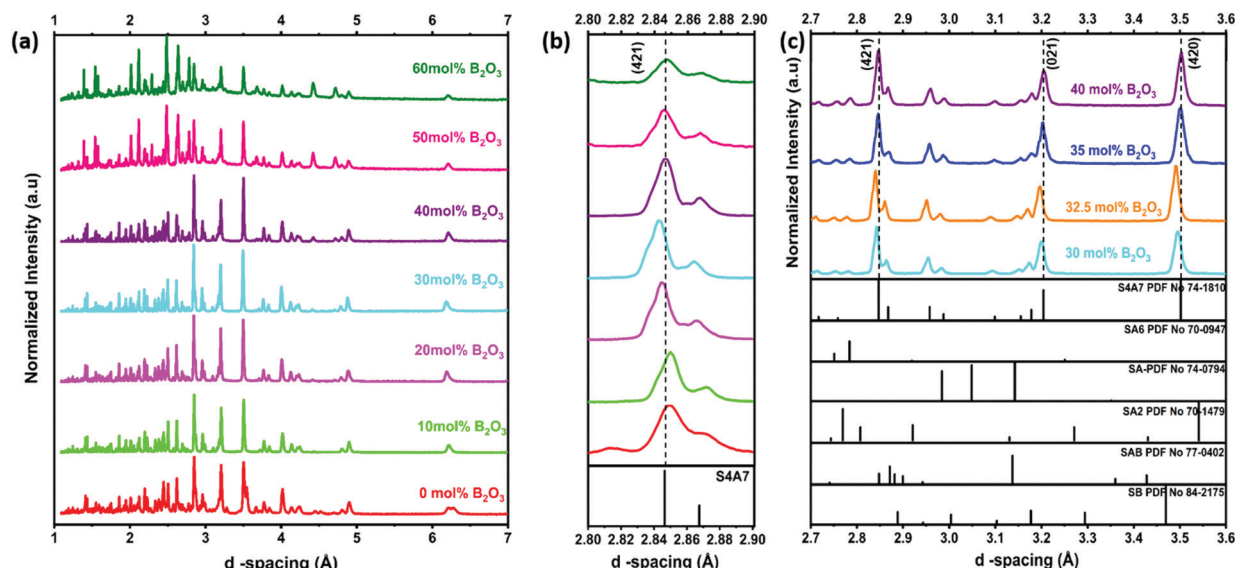


Fig. 3 X-ray diffractograms of S4A7EDxB ($0 \leq x \leq 60 \text{ mol\% B}_2\text{O}_3$) (a) full range; (b) magnified around the (421) diffraction peak; and (c) diffractograms for S4A7EDxB ($30 \leq x \leq 40 \text{ mol\% B}_2\text{O}_3$).

shifts in the $20 \leq x < 40 \text{ mol\% B}_2\text{O}_3$ compositions in Fig. 3c. The main diffraction peaks corresponding to the crystallographic planes (421), (021), and (420) shifted from the reference positions with the addition of B₂O₃ for $0 < x < 40 \text{ mol\%}$. In S4A7EDxB compounds with $20 \leq x < 40 \text{ mol\% B}_2\text{O}_3$, the (421) shifted toward a lower *d*-spacing (Fig. 3b and c).

The presence of an amorphous phase was revealed by quantifying the relative amount of crystalline phase in Fig. 4a and Table S1 (ESI[†]). The results of Rietveld quantitative phase analysis are shown in Fig. 4b and Fig. S1 (ESI[†]) shows the results of the XRD patterns of S4A7EDxB samples ($0 \leq x \leq 60 \text{ mol\% B}_2\text{O}_3$), while Table S2 (ESI[†]) summarizes the results of quantitative analysis of XRD data. The sample prepared without

B₂O₃ was 83.3% crystalline and was multi-phase, being dominated by the S4A7 phase (72.8%), along with SA2 (14.85%), SA6 (6.38%) and SA (5.9%) as the minor strontium aluminate phases. The maximum amount of phases exhibiting room temperature persistent luminescence, S4A7 and SA, was found in the 10 mol% B₂O₃ compound, which was of 93.3% S4A7 phase (and negligible SA content).

S4A7EDxB for $x = 30 \text{ mol\% B}$ consisted of 81.02% S4A7, 5.15% SA, 9.09% SA6 and 4.74% SA2 phases. A similar result was obtained for the compound prepared with 32.5 mol% B₂O₃. Increasing the B₂O₃ content to 35 mol% led to a significant drop in the S4A7 content, which decreased further upon increasing the B₂O₃ content, whereas the amount of SA6 phase



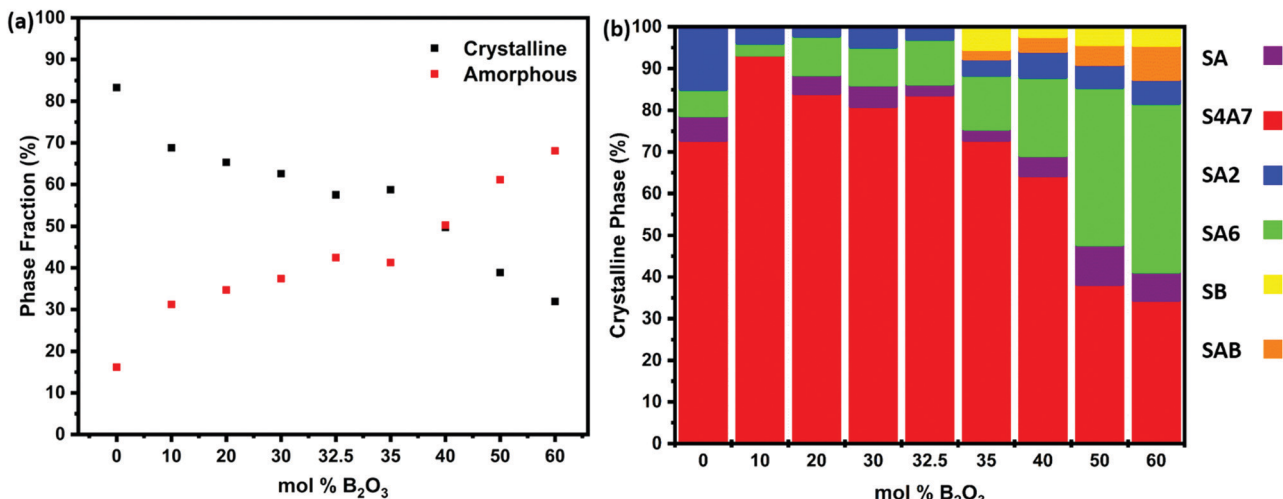


Fig. 4 (a) Percent crystallinity and amorphous phase fraction determined from the XRD data of S4A7EDxB ($0 \leq x \leq 60$ mol% B₂O₃) samples; (b) phase distribution revealed by Rietveld phase analysis of the diffractograms shown in Fig. 3.

increased. The diffractogram of S4A7EDxB for $x = 40$ mol% B showed additional peaks, indicating the presence of SrB₂O₄ (SB) and Sr₂B₂Al₂O₈ (SAB) phases along with S4A7, SA6, SA2 and SA phases. Further addition of B₂O₃ resulted in the dominance of SA6 over all other crystalline strontium aluminate phases. When S4A7EDxB was processed with 60 mol% B₂O₃, the amorphous content reached 68%. In the remaining crystalline material, SA6 dominated the multi-phase distribution, along with substantial amounts of SB and SAB phases.

The FTIR spectra were analyzed to identify the structural changes in strontium aluminate compounds arising from the different amounts of B₂O₃. The FTIR spectra of S4A7EDxB, where $0 \leq x \leq 60$ mol% B₂O₃, are shown for the region 500–4000 cm⁻¹ in Fig. 5a. No peak was detected above 1500 cm⁻¹. The region 500–1000 cm⁻¹ showed vibrations related to the alumina scaffold of the S4A7 crystal structure in Fig. 5b. In this region, the characteristic vibrations of O–Al–O were observed in the phase-pure, undoped S4A7 reference spectra (Fig. S2a, ESI†).

Absorption bands were observed for Al–O features, both anti-symmetric stretching (700–1000 cm⁻¹ region) and anti-symmetric bending (550–650 cm⁻¹). When the B₂O₃ content exceeded 40 mol%, the intensity decreased for some vibrations in the 800–1000 cm⁻¹ range (in the FTIR spectra); meanwhile, the increase in the shoulder indicating enhanced absorption at around 710 cm⁻¹ suggested the bending vibrations of B–O–B linkages in the borate network. No absorption peak was detected in the 1000–1600 cm⁻¹ range for the reference (undoped) and the $x = 0$ mol% B₂O₃ compounds. With the addition of B₂O₃, two distinct absorption profiles were observed in the range of 1180–1200 cm⁻¹ and 1330–1370 cm⁻¹, as shown in Fig. 5c. These two isolated bands became apparent when deconvoluted by assuming Gaussian profiles (Fig. 5d). The absorption band at 1180–1200 cm⁻¹ corresponded to asymmetric B–O stretching vibrations of BO₄ units and the absorption at 1330–1370 cm⁻¹ was assigned to B–O stretching

vibrations of BO₃ groups.^{21,44,45} Using the integrated area beneath these 2 vibration peaks, we determined the relative change in N₄ = (B₄/(B₃ + B₄)) as a function of the B₂O₃ content (Fig. 5e).

The intensity of these bands increased in a linear manner with the B₂O₃ content, until reaching 30 mol% B₂O₃. Across the range $30 \leq x \leq 40$ mol% B₂O₃, the N4 ratio reached a maximum value in the 32.5–35 mol% B₂O₃ compounds, and then declined in the 40 mol% B₂O₃ one.

When the B₂O₃ content exceeded 50 mol% B₂O₃, we observed a sharp increase in peak intensities, concurrent with the amorphous content.

The changes in the vibrational modes of crystalline S4A7 caused by the incorporation of B₂O₃ into the crystal structure were also investigated by using Raman spectroscopy analysis. The phase-pure, orthorhombic S4A7 compound was characterized to serve as a reference fingerprint (Fig. S2b, ESI†). The Raman spectra of S4A7EDxB, where $0 \leq x \leq 60$ mol% B₂O₃, are summarized in Fig. 6a. With an increase in B₂O₃, we observed a weakening in the intensity of the main 318 cm⁻¹ peak, accompanying a shift toward shorter wavelengths (Fig. 6b). Furthermore, the appearance of additional peaks in the range of 700–900 cm⁻¹ was observed in S4A7EDxB ($30 \leq x \leq 60$ mol% B₂O₃), which largely accounted for the boron-related structural groups (Fig. 6c). Fig. 6d shows the evolution in the Raman shift within 700–900 cm⁻¹ for the S4A7EDxB compounds with $30 \leq x \leq 60$ mol% B₂O₃, correlated with the deconvolution analysis of these peaks in Fig. 6e.

The B \bullet O₃ unit has a D_{3h} symmetry point group symmetry and has $\Gamma_{D_{3h}} = A'_1(R) + 2E'(IR, R) + A''_2(IR)$, in which R and IR represent Raman and infrared activity. These vibration modes are symmetric stretching $\nu_1(A'_1)$, out-of-plane bending $\nu_2(A''_2)$, asymmetric stretching $\nu_3(E')$, and in-plane bending $\nu_4(E')$ modes. The replacement of bridging oxygen (O) by a non-bridging one (O⁻) reduces the symmetry to C_{2v} from D_{3h} , while the doubly degenerate (E') asymmetric stretching mode will split into two IR-active components of A₁ and B₂ symmetry.^{46–49}



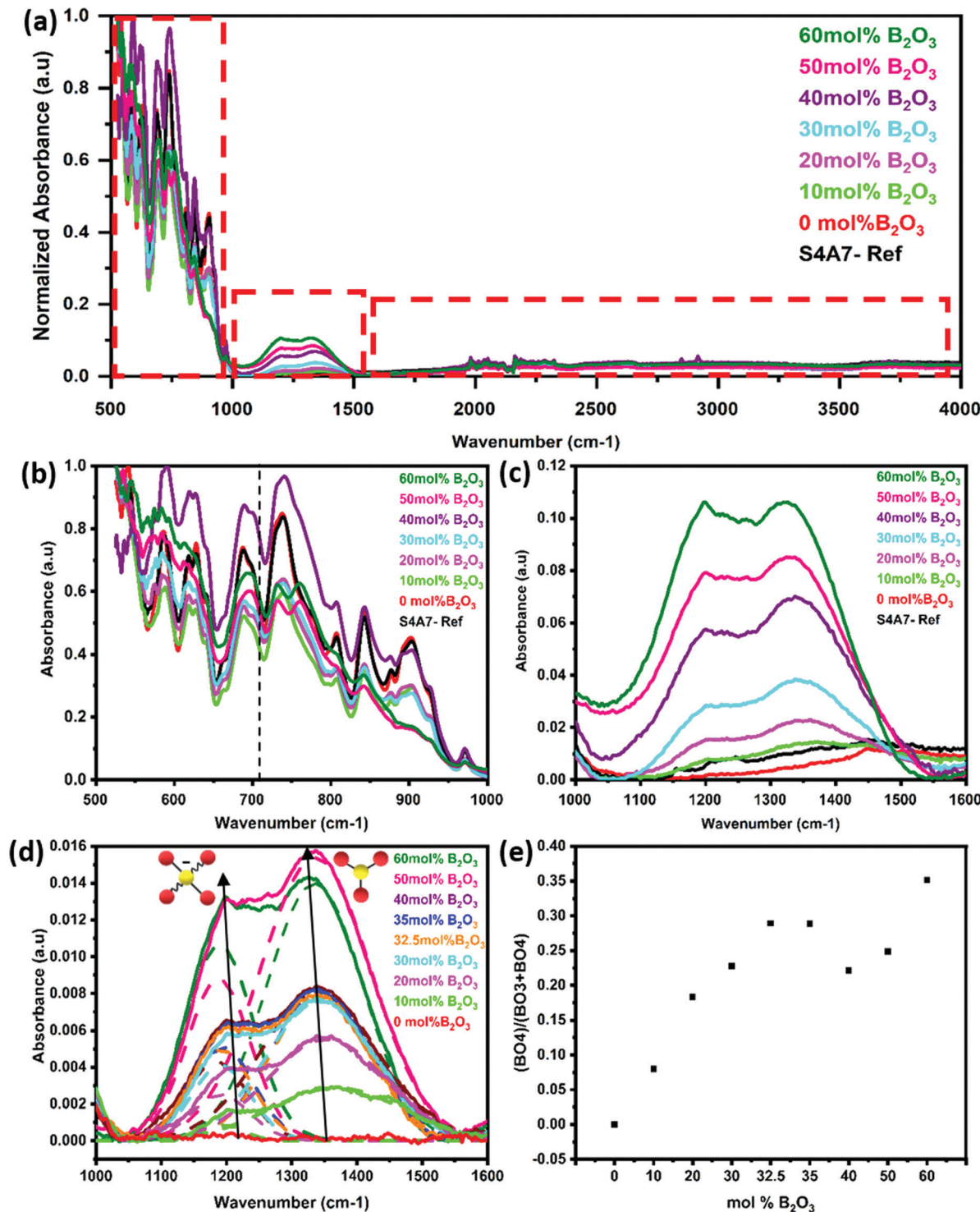


Fig. 5 FTIR spectra summary for S4A7EDxB ($0 \leq x \leq 60$) mol% B (a) full range spectra; (b) $500\text{--}1000\text{ cm}^{-1}$ magnified; and (c) $1000\text{--}1600\text{ cm}^{-1}$ magnified; (d) deconvolution of the $1000\text{--}1600\text{ cm}^{-1}$ magnified region; (e) N_4 vs. mol% B_2O_3 , where $N_4 = (B_4/(B_3 + B_4))$.

The strongest peak at 808 cm^{-1} was assigned to the symmetric breathing vibration of $B\ddot{O}_3$ boroxyl rings, reflecting the presence of vitreous B_2O_3 compounds (Fig. 7a).^{13,44} The second strongest peak at 750 cm^{-1} was assigned to the symmetric breathing vibration of 6-membered rings, containing one $B\ddot{O}_3$

triangle replaced by one $B\ddot{O}_4$ tetrahedron (Fig. 7b).^{18,44} The band at 803 cm^{-1} was assigned to metaborate $B\ddot{O}_2O^-$ units (Fig. 7c). Additional evidence of metaborate formation was provided by the weaker bands at around the 640 cm^{-1} symmetric stretch of B–O–B rings.^{17,50} With elevated B_2O_3 content,

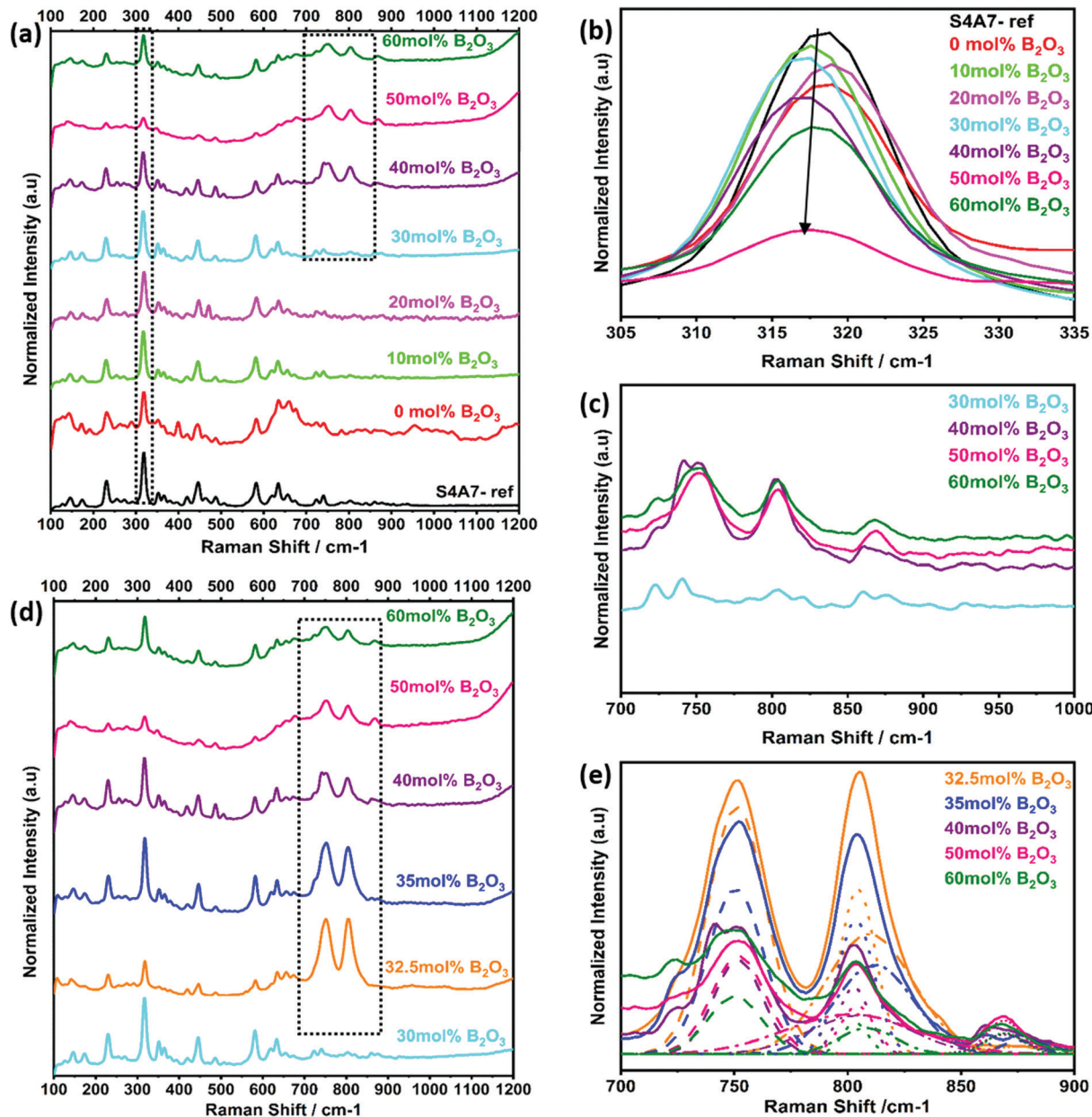


Fig. 6 (a) Raman shift spectra of S4A7EDxB ($0 \leq x \leq 60$ mol% B_2O_3); (b) relative peak shift ca. 318 cm^{-1} ; (c) overlap region for $700\text{--}1000 \text{ cm}^{-1}$; (d) S4A7EDxB ($30 \leq x \leq 60$ mol% B_2O_3); (e) deconvoluted Raman shift spectra for S4A7EDxB ($30 \leq x \leq 60$ mol% B_2O_3).

the formation of orthoborate monomers at 868 cm^{-1} was observed, reflecting the depolymerization of boroxyl rings and rings with BO_4 tetrahedral units to orthoborate monomers (Fig. 7d).^{16,21} Both Raman and IR spectroscopic techniques were applied in a complementary way to reveal the formation and changes of complex boron units.

Raman mapping was performed on sintered polished pellets of S4A7EDxB compounds of $x = 30$ mol% B_2O_3 and 40 mol% B_2O_3 . To analyze the hyperspectral Raman dataset, we applied a multivariate curve resolution-alternating least squares (MCR-ALS) method. The spectral data matrix was analyzed in the Wire 3.4 environment using the MCR-ALS module, in which the data

matrix was decomposed into spectral profiles (“Loadings”) and concentration profiles (“Scores”).

In the $x = 30$ mol% B_2O_3 data set, three physically significant MCR-ALS components were observed; and a good correlation was obtained between the identified loadings of the reference spectra from pure S4A7, B_2O_3 powder, and epoxy. Fig. S3a (ESI[†]) shows the visible-light image of the region of interest for Raman spectral imaging for the $x = 30$ mol% B_2O_3 S4A7EDxB specimen. Fig. S3b (ESI[†]) shows the scan area at a higher magnification. The hyperspectral image of the first component is shown in Fig. S3c (ESI[†]), which is related to the S4A7 grains. Both overlay and color-coded (Fig. S3d, ESI[†]) images show the

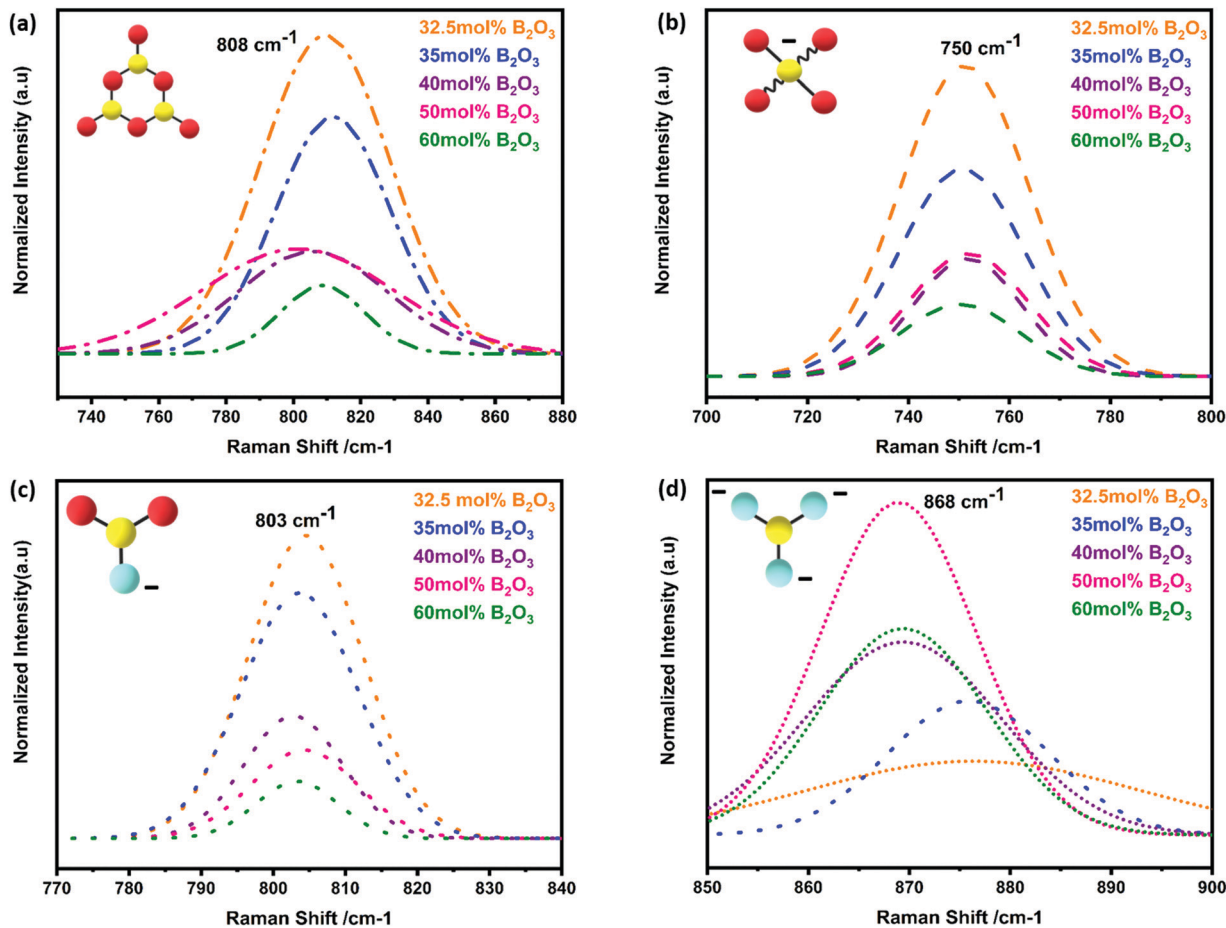


Fig. 7 Deconvoluted Raman-shift spectra of S4A7EDxB ($30 \leq x \leq 60$ mol% B_2O_3) for vibrations of (a) boroxyl rings of corner-sharing $[\text{BØ}_3]^{3-}$; (b) $[\text{BØ}_4]^{3-}$; (c) metaborate $[\text{BØ}_2\text{O}]^{3-}$; and (d) orthoborate monomers of $[\text{BO}_3]^{3-}$.

spatial distribution of S4A7 grains. The representative Raman spectrum from component 2, shown in green, is consistent with the reference Raman spectrum obtained from neat epoxy (Fig. S3e, ESI†). Since the pellet was necessarily embedded in an epoxy mold for polishing, the observation of epoxy at the surface was not surprising (Fig. S3f, ESI†). The last component appearing was attributed to B_2O_3 , which is shown in blue (Fig. S3g, ESI†). Color-coded images formed by boron units revealed the distribution of boron units around the S4A7 grains (Fig. S3h, ESI†). In Fig. S3i (ESI†) the corresponding Raman spectra of the 3 components compared to the reference Raman spectra of pure S4A7, B_2O_3 and the epoxy are shown.

MCR-ALS analysis of Raman shift hyperspectral maps of $x = 40$ mol% B_2O_3 S4A7EDxB are shown in Fig. S4 (ESI†). There were four components, which were consistent with the reference spectra collected from phase pure S4A7, SA6, B_2O_3 powder, and the epoxy. The distribution of complex phases was color-coded in the MCR score—*i.e.*, concentration profile—plot shown. The multi-phase region was selected for mapping to analyze the distribution of crystalline and amorphous components (Fig. S4a and b, ESI†). The first component originated from the epoxy, as was observed in the pellet of $x = 30$ mol%

B_2O_3 (Fig. S4c and d, ESI†). The hyperspectral image that was color-coded red had correlated with large S4A7 grains (Fig. S4e and f, ESI†). The presence of small SA6 grains in the scan area resulted from the third component, which was colored green (Fig. S4g and h, ESI†) and was consistent with the reference spectrum of SA6. Component 4 reflected the presence of polyborate units and their spectral distributions across the mapped region (Fig. S4i and j, ESI†). This component appeared to be superposed with the characteristic vibrations of S4A7, which is due to the borate-rich glassy phase being adjacent to S4A7 grains. In Fig. S4k (ESI†) representative Raman spectra of the 4 components compared to the reference Raman spectra of B_2O_3 , epoxy, pure S4A7, and pure SA6 are shown.

Fig. 8 shows the effect of individual rare earth dopants on the S4A7 unit cell. The S4A7 unit cell doped with only Eu exhibited distortion in the Raman shift spectrum especially in the region 550 – 850 cm^{-1} , while doping with only Dy did not generate such a distortion. This observation can be attributed to the difference in size between Eu and Dy ions, the latter of which, being larger, is more prone to segregate out of the grain above the solubility limit,⁴ as a result, S4A7 doped only with Dy exhibited similar Raman spectra with the non-doped sample.

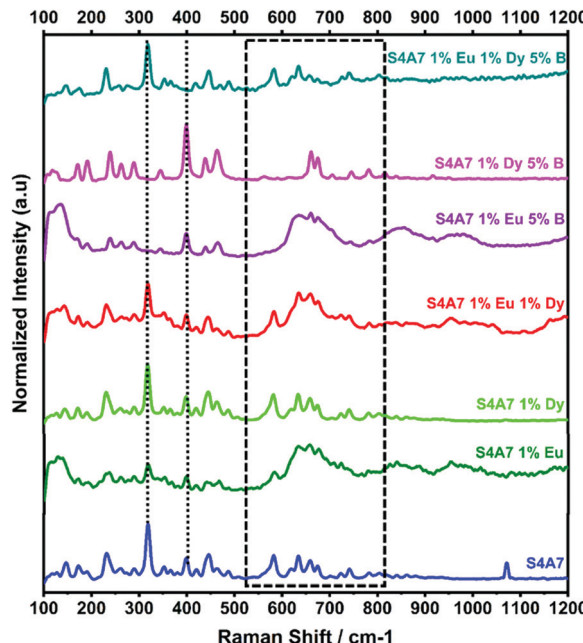


Fig. 8 Raman-shift of the S4A7 unit cell doped with only Eu, Dy and B.

The compound co-doped with Eu and Dy showed a spectrum with high intensity and narrow bands consistent with S4A7 doped only with Eu.

It should be noted that in S4A7 containing 1 at% Eu 5 mol% B_2O_3 and S4A7 1 at% Dy and 5 mol% B_2O_3 , neither compound exhibited the vibration modes of S4A7. The Raman shift spectrum of S4A7 1 at% Eu and 5 mol% B_2O_3 had wider and weaker Raman bands and was missing the shift at 318 cm^{-1} . Meanwhile, the spectrum of S4A7 1 at% Dy and 5 mol% B_2O_3 had Raman bands similar to those of SA2. The changes induced in the Raman spectra suggest that the presence of boron affects the atomic arrangements in each compound differently, which may also differentiate their incorporation into the Sr sub-lattice sites in the unit cell. Finally, the Raman spectra of S4A7EDxB with $x = 5$ mol% B_2O_3 contained known vibrational modes that were observed in other S4A7EDxB compounds, the ones containing S4A7 as a major phase.

Discussion

The optimal B_2O_3 composition for the Pechini-processed S4A7EDxB compound was determined to lie in the range $30\text{ mol\% B}_2\text{O}_3 \leq x \leq 40\text{ mol\% B}_2\text{O}_3$. De-trapping from both shallow-trap and deep-trap states, as extrapolated from the afterglow decay curves, was of the longest duration within this range of boron content and correlated with persistent luminescence of the longest duration. The introduction of B_2O_3 into the S4A7 unit cell appears to be associated with increasing the depth of both types of trap centers, leading to a slower decay rate than that observed in other B_2O_3 -doped compositions.

The variation in strontium aluminate phase distribution was observed with the addition of boron oxide in samples prepared

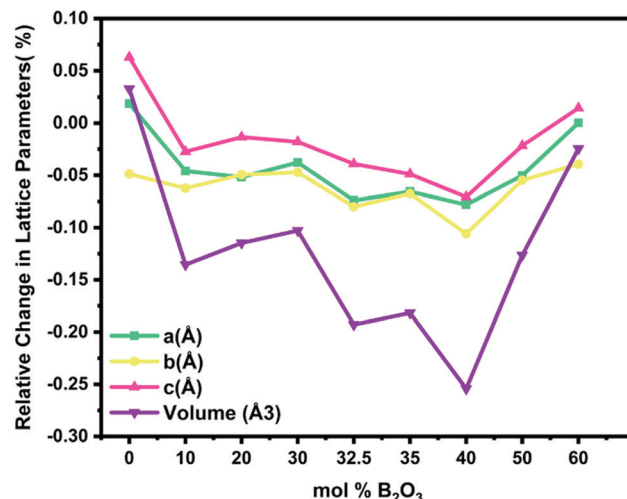


Fig. 9 Changes in the unit cell dimensions of S4A7 with an increase in the B_2O_3 content during Pechini processing.

with identical synthesis and heat treatment conditions (Fig. 4b). We focused our analysis on the atomic positions in the target persistent luminescence phase, S4A7EDxB, where $0 \leq x \leq 60\text{ mol\% B}_2\text{O}_3$, by performing Rietveld structural refinement analysis on the XRD data. As shown in Fig. 9, the refined unit-cell parameters revealed the most significant shrinkage in the S4A7 crystal lattice dimensions (a , b , c , and volume) starting from 30 mol% B_2O_3 and extending to 40 mol% B_2O_3 (Table S3, ESI†). These parameters were compared against those of the pure S4A7 reference. The reason for this apparent change in the unit cell parameters appears to be induced by the presence of borate structures in the lattice.

The effect of B_2O_3 incorporation on the crystalline S4A7 unit cell was visualized with the help of Vesta, using the atomic positions determined by Rietveld refinement analysis (Fig. S5, ESI†). The S4A7 unit cell contains 6 different crystallographically non-equivalent Al sites, and these Al sites are coordinated with 4 or 6 oxygen atoms. The Al1, Al2 and Al3 form AlO_4 tetrahedra, while Al4, Al5 and Al6 form AlO_6 octahedra. The interconnected network of these alumina polyhedra forms a rigid 3-D scaffold, with large cavities filled by Sr cations, providing electrical charge balance. There are two nonequivalent Sr sites: Sr1 coordinated by 10 oxygens (Wyckoff 4j) and Sr2 coordinated by 7 oxygens (Wyckoff 4i) (Fig. 1).

The change in the polyhedral volume of each of the AlO_4 tetrahedra is shown in Fig. 10a. Within the compositional range S4A7EDxB $x = 30\text{--}40\text{ mol\% B}_2\text{O}_3$ the volume of the sites Al2 and Al3 increased, while that of Al1 did not change significantly (Fig. S6, ESI†). With the further addition of up to 60 mol% B_2O_3 , the Al2 polyhedral volume increased by *ca.* 12.6%. Boron appeared not to have caused a significant change in the octahedral AlO_6 chains in the $x = 30\text{--}40\text{ mol\% B}_2\text{O}_3$ range, when compared to the reference sample (Fig. 10b). The overall decrease in the AlO_6 polyhedra correlated with an increase in the B_2O_3 content (Fig. S7, ESI†).



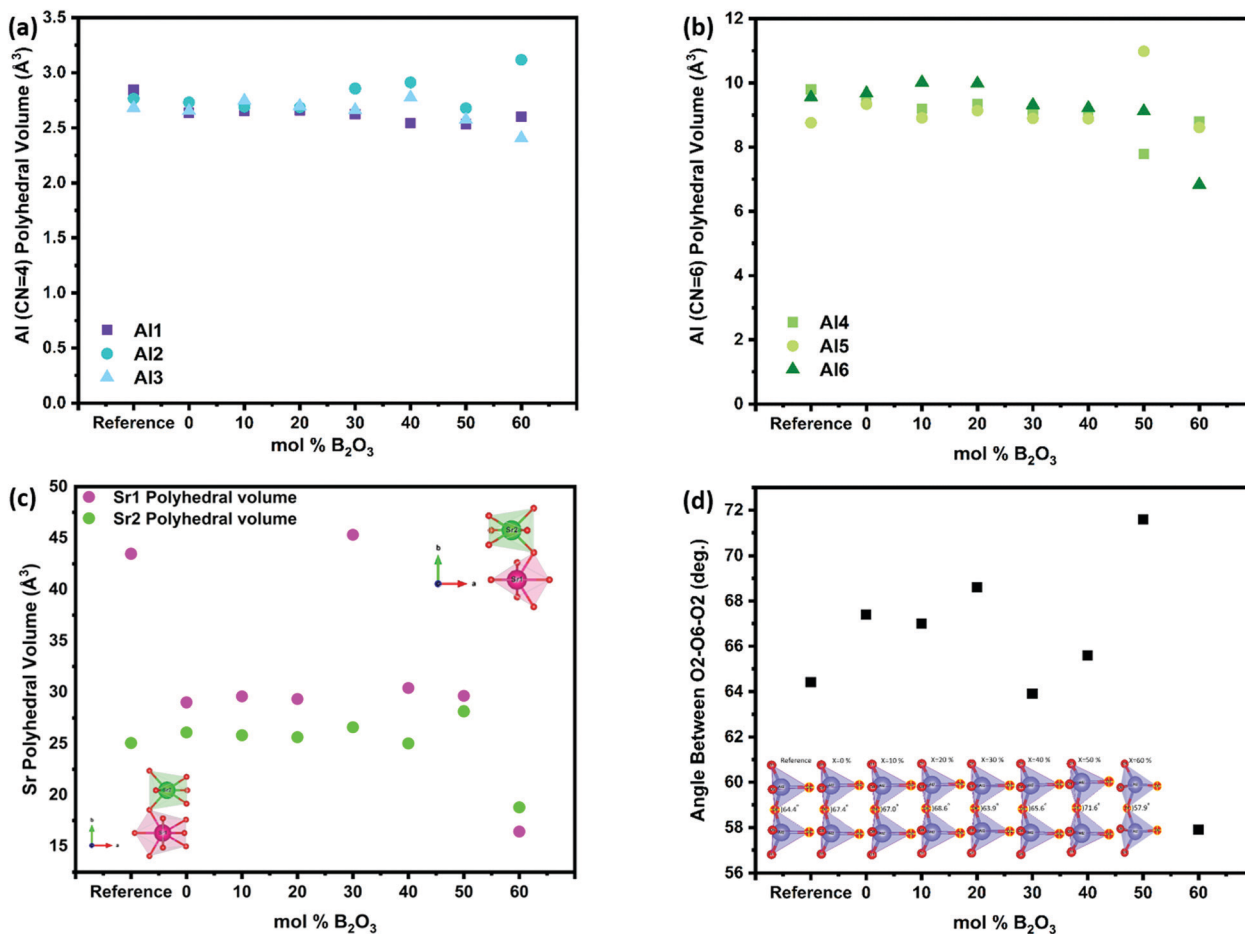


Fig. 10 Volume changes resulting from modification by B_2O_3 addition for S4A7 reference and the representative S4A7EDxB ($0 \leq x \leq 60$ mol% B_2O_3) for (a) Al CN = 4 polyhedra; (b) Al CN = 6 polyhedra; and (c) Sr1 and Sr2 coordination polyhedra; and (d) angle between oxygen sites O2–O6–O2.

The incorporation of Eu into Sr sites has already been demonstrated earlier by our group using AR-STEM imaging.²⁵ In this work, the results of our analyses revealed that both types of Sr sites investigated in compounds containing 30–40 mol% B_2O_3 showed localized expansion of their volume, particularly in the Sr1 polyhedral volume (Fig. 10c and Table S4, ESI†). Sr1 (CN = 10) has lower covalency compared to Sr2 (CN = 7). This expansion would have altered the crystal field coordination environment around the Eu^{2+} ions, resulting in changes to the electronic structure of Eu^{2+} that control the persistent luminescence.

With Sr1 located in the large interstitial cavities of the alumina scaffold, any expansion of their polyhedral volume would deform the scaffold structure (Fig. 10d) (Table S5, ESI†). The angle between the neighboring Al2 tetrahedra within the $x = 30$ –40 mol% B_2O_3 compositional range showed values closest to the undoped reference spectra. In addition, the expansion in Sr sites pushed the 4-coordinated alumina polyhedra towards each other. This observation may explain the decreased d_{421} -spacing between the main (421) planes within the compositional range of 30 $\leq x \leq 40$ mol% B_2O_3 .

Our material system had crystallized directly from an amorphous precursor, the one in which multiply connected and

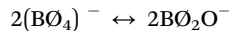
isolated borate units were present throughout the microstructural evolution. Mid-infrared transmission and micro-Raman spectra revealed that S4A7EDxB ($0 \leq x \leq 60$ mol% B_2O_3) compositions contained polyborate supramolecular arrangements like neutral trigonal BO_3 units, charged BO_4^- tetrahedra, metaborate BO_2O^- , and orthoborate $(\text{BO}_3)^{3-}$. The red-shift in the 1000–1600 cm^{-1} absorption band envelope has been reported for other highly modified borate glasses.^{17,18} For the compositions of S4A7EDxB with $10 \leq x \leq 40$ mol% B_2O_3 , the progressive red-shift of the 1342–1370 cm^{-1} peak frequency was attributed to the formation of metaborate triangles BO_2O^- that were manifested by the deconvoluted Gaussian BO_3 component. Changes in the number of $\text{B}-\text{O}^-$ and $\text{B}-\text{O}$ bonds undergoing stretching vibrations in BO_2O^- and BO_3 units resulted in the red-shift and asymmetry of these bands.^{18,49} In particular, the red-shift suggests stronger interactions between the non-bridging oxygens and other cations—Sr²⁺, Al³⁺, Eu²⁺, and Dy³⁺—that consequently result in weaker $\text{B}-\text{O}^-$ bonding.⁴⁹

For the S4A7EDxB compositions of $40 \leq x \leq 60$ mol% B_2O_3 , these trends continued in the 1335–1342 cm^{-1} spectral shift range, which was attributed to the progressive depolymerization in the borate network in superstructural units transforming into metaborate $(\text{BO}_2\text{O})^-$ and then into orthoborate

$(\text{BO}_3)^{3-}$ monomers. The peak at around 710 cm^{-1} indicated an out-of-plane bending mode of borate $(\text{BO}_3)^{3-}$ triangular units. It should be noted that this mode was only present in compositions of $x > 40\text{ mol\% B}_2\text{O}_3$, indicating that orthoborate units of $(\text{BO}_3)^{3-}$ were present significantly in these compositions. In addition, the absorption bands at around $800\text{--}1000\text{ cm}^{-1}$ had weakened, which was indicative of vibration modes damped by the presence of other cations in the structure. The absorption peak component related to BO_4 units ($1189\text{--}1198\text{ cm}^{-1}$) did not show a pronounced overall red-shift compared to that of BO_3 ($1335\text{--}1370\text{ cm}^{-1}$).

The Raman spectra had also showed the characteristic vibrations of boron units, but additionally revealed deeper insight into the polyborate suprastructures. At shift frequencies $< 750\text{ cm}^{-1}$, there were strong lattice vibrations from the different phases of strontium aluminate, which generally obscured the ν_4 in-plane bending mode of orthoborate units that appears at around 625 cm^{-1} . When S4A7EDxB contained $30\text{--}40\text{ mol\% B}_2\text{O}_3$, the simultaneous occurrence of bands at around 808 cm^{-1} , 750 cm^{-1} , and 803 cm^{-1} in the Raman spectra suggested the presence of polymeric polyborate groups in the orthorhombic S4A7 crystal structure. Deconvolution analysis of the $700\text{--}900\text{ cm}^{-1}$ band revealed not only the presence of planar BO_3 rings (808 cm^{-1} band), but also the symmetric breathing vibration of the six-membered ring one replaced by BO_4^- (750 cm^{-1} band). The presence of such rings suggests an interconnected network of borate units in the $32.5\text{ mol\% B}_2\text{O}_3$ and $35\text{ mol\% B}_2\text{O}_3$ compounds. These spectral changes are consistent with the transformation of BO_3 units into BO_4^- tetrahedra, as revealed by FTIR spectra (Fig. 5d and e). The maximum N4 value was attained by processing S4A7EDxB with $32.5\text{--}35\text{ mol\% B}_2\text{O}_3$, which indicated the conversion of boroxyl rings to BO_4^- tetrahedra groups in compounds of this composition range.

Furthermore, the co-existence of metaborate suprastructural units—resonance at 803 cm^{-1} and 640 cm^{-1} —with BO_4^- ones, suggests equilibrium between the isomeric trigonal $2\text{B}\text{O}_2\text{O}^-$ and charged tetrahedral units within S4A7EDxB ($30 \leq x \leq 40\text{ mol\% B}_2\text{O}_3$):



As the B_2O_3 concentration was increased, the borate backbone became further de-polymerized. The number of non-bridging oxygen ions increased with the formation of orthoborate units, as revealed by the Raman peak at 868 cm^{-1} . De-polymerization resulted in the break-up of the B—O—B linkages and an increase in the free volume within the scaffold of coordination polyhedra.

In S4A7EDxB compounds of $>40\text{ mol\% B}_2\text{O}_3$, the intensity decreased for the bands at 808 cm^{-1} , 803 cm^{-1} and 750 cm^{-1} , while the intensity of that at 868 cm^{-1} (ortho) increased. Such a spectral evolution indicates that the number of bridging oxygen per boron decreased, resulting in a dampening and red-shift of the vibration of B—O bonds in the BO_3 , $\text{B}\text{O}_2\text{O}^-$, and BO_4^- units. The frequency of the B—O[—] stretching in orthoborate triangles

BO_3^{3-} was observed to increase with an increase in the non-bridging oxygen content. Meanwhile, the lower intensity and red-shift of the main vibration at 318 cm^{-1} and the distortion in phonon vibrations in the Raman spectra indicated that the covalent nature of Al—O bonding was affected by an increase in non-bridging oxygen.

In compositions with the B_2O_3 content in excess of that with the longest persistent luminescence duration ($30\text{--}40\text{ mol\% B}_2\text{O}_3$), the electron density of the boron ions had increased as a consequence of depolymerization of the boroxyl network, which increased the non-bridging oxygen content. As a consequence, the change in the proportion of different borate units perturbed localized electrical charge distribution within the crystal unit cell, which would in turn modify the luminescence duration. Because the compounds of optimal luminescence contained connected planar triangular units and tetrahedral units, along with metaborate units, these results suggest that the formation of polymerized borate groups from BO_3 triangles and BO_4 tetrahedra contributes to the persistence behavior by stabilizing the Eu^{3+} and Dy^{2+} in the S4A7 unit cell. The coordination environment around the Eu and Dy ions, *i.e.* the crystal field around them, was affected by changes in the polyborate supramolecular structures. After all, the B—O network is known to act as an electron trapping center, which may serve as additional trap centers.⁵¹

The fraction and arrangement of polyborate groups depend not only on the concentration of B incorporated into the crystal structure but also the type of modifier cation, which will affect the structure and optical properties of the overall network. Sr plays a dual role as a network modifier and electrical charge compensator as a function of the $\text{SrO}:\text{Al}_2\text{O}_3$ ratio.⁵² There was a competition between Al and B as network formers for bonding with the oxygen ions, which also contributed to different numbers of bridging oxygens. When there was a coexistence of aluminum and boron oxides, the resulting glass was characterized predominantly by the formation of AlO_4 tetrahedra.²⁰ In a low glass and crystalline system rich in Al^{3+} modifier, AlO_6 formed for charge compensation.¹⁹

The amount of B_2O_3 used determined the way B interacts with the ions in the amorphous pre-ceramic, and consequently the evolution in the crystal structure. Evaluation of the crystal structure of S4A7 revealed that boron incorporation had altered the lattice parameters, to a degree correlated with the amount of B_2O_3 used during synthesis. The integrity of the unit cell was maintained within the S4A7EDxB composition optimal for extreme persistent luminescence— $30 \leq x \leq 40\text{ mol\% B}_2\text{O}_3$ —although some cation coordination polyhedra underwent deformation, particularly the Sr1 polyhedra and the AlO_4 tetrahedra: Al1, Al2, and Al3.

For $30 \leq x \leq 40\text{ mol\% B}_2\text{O}_3$, there was increased free volume around Sr^{2+} and $\text{B}\text{O}_2\text{O}^-$ structures present in the crystal structure, suggesting that the polyhedral scaffold changes around Sr^{2+} may be attributed to the $\text{B}\text{O}_2\text{O}^-$ inducing AlO_x polyhedral rotation. The longest afterglow microstructure existed in this composition range, in which there was a transition in the interaction between BO_3 units and the alumina



polyhedra in the aluminoborate melt. Three-fold boron coordinated with oxygen from alumina polyhedra and the conversion from three- to four-coordinated boron induced a volume change in the alumina polyhedra. It appears that the boron structural anomaly in this composition range was related to the alumina polyhedral rotation.

For $x > 40$ mol% B_2O_3 , the BO_3/BO_4 ratio defined the network character. Strontium metaborate (SrB_2O_4) precipitated, with Sr^{2+} providing charge compensation for the excess non-bridging oxygen. Because Sr^{2+} was occupied in SrB_2O_4 , the stoichiometry of Sr:Al in the melt changed. The Al-rich regions precipitated as SA6, consistent with the increased SA6 content revealed by Rietveld phase fraction analysis.

To summarize, the results presented in this work show a strong correlation between the duration of persistent luminescence in S4A7EDxB (Fig. 2) with the B_2O_3 content during Pechini processing and with the metaborate content (Fig. 5 through 7) appearing in the crystal structure of $\text{Sr}_4\text{Al}_{14}\text{O}_{25}$. Our analysis of these results reveals new details in the state-of-the-art structural model of long afterglow S4A7EDxB, suggesting that metaborate induces the clustering of optically active ionic point defects, which support persistent luminescence.

Conclusions

The longest persistent luminescence in S4A7EDxB compounds synthesized by Pechini processing was observed in compounds prepared with 30–40 mol% B_2O_3 . Micro-Raman analysis of S4A7EDxB grains revealed the presence of borate supramolecular structures in their orthorhombic $\text{Sr}_4\text{Al}_{14}\text{O}_{25}$ crystal structure as BO_3 , BO_2O^- , and BO_4^- units, with an equilibrium between BO_2O^- and BO_4^- units. BO_4^- units replaced BO_3 ones in the network of polymerized $\text{BO}_3/\text{BO}_2\text{O}^-$ groups and induced deformation in the scaffold of interconnected AlO_4 polyhedra that led to increased Sr^{2+} site volume. These results suggest that the enlarged Sr sub-lattice sites neighboring the negatively charged non-bridging oxygen accommodate Eu and Dy dopant incorporation, giving rise to clustering of ionic point defects that support persistent luminescence.

Author contributions

A. C. E.: investigation, methodology, visualization, writing—original draft; S. K.: investigation and methodology; E. D. investigation; C. O.: conceptualization, funding acquisition, resources, methodology, writing—review & editing.

Conflicts of interest

There are no conflicts to declare.

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