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Superstructure formation in SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆†

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X-ray pure samples of SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆ were synthesized from appropriate amounts of binary nitrides (Sr₃N₂, Ba₃N₂ and BN in sealed niobium ampoules and EuN, Ba₃N₂ and BN in BN crucibles, respectively) at temperatures up to 1370 K. The structure of SrBa₈[BN₂]₆ was refined from single crystal X-ray diffractometer data: $Fd\bar{3}m$, a=1595.1(1) pm, $wR(F^2)=0.0515$, $387\ F^2$ values and 21 variables. EuBa₈[BN₂]₆ has a lattice parameter of 1595.00(9) pm. Both nitridoborates adopt a new 2 × 2 × 2 superstructure variant of the LiCa₄[BN₂]₃ type, realized through ordering of vacancies and Sr²⁺ and Eu²⁺ cations, respectively. The structures of SrBa₈[BN₂]₆ and LiCa₄[BN₂]₃ are related by a group–subgroup scheme. The Sr²⁺/vacancy ordering leads to an asymmetric coordination (1 × Sr²⁺ and 8 × Ba²⁺ in a distorted, mono-capped square prism) for the [BN₂]³⁻ units with B–N distances of 132 and 136 pm. Vibrational spectra of SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆ confirm the discrete linear [BN₂]³⁻ units and ¹¹B solid state MAS NMR spectra are compatible with single crystallographic sites for the boron atoms. In EuBa₈[BN₂]₆ the spectra are profoundly influenced by interactions of the ¹¹B nuclei with the unpaired electrons of the paramagnetic Eu²⁺ ions.

1. Introduction

The $[BN_2]^{3-}$ anion is one of the basic building units in nitridoborate crystal chemistry. A first proof for the $[BN_2]^{3-}$ units was obtained by Goubeau and Anselment from IR spectra of Li₃[BN₂], Ca₃[BN₂]₂ and Ba₃[BN₂]₂, while their precise crystal structures were determined from single crystal X-ray data only much later and also the europium compound Eu₃[BN₂]₂ was reported. 11,12

Phase analytical studies later on led to a series of nitridoborate halides $M_2[BN_2]X$ (M = Ca, Sr, Eu; X = F, Cl, Br, I) and $Ba_8[BN_2]_5F$.^{13–16} Depending on the metal and halide matrix, the $[BN_2]^{3-}$ units show small distortions, *i.e.* two slightly different B–N distances within the unit and/or N–B–N angles slightly deviating from 180°. Extension of the crystal chemistry

is also possible through the ordered replacement of the alkaline earth metal position by alkali or divalent rare earth metals. Many such nitridoborates, $AM_4[BN_2]_3$ (A = Li, Na; M = Ca, Sr, Ba, Eu),^{17–21} have been characterized structurally. They are electron-precise nitridoborates with linear $[BN_2]^{3-}$ units. Both series of quaternary nitridoborates have been studied intensively with respect to their IR and Raman spectra. ^{16,22,23} Distortions (deviations from linearity) of the $[BN_2]^{3-}$ units are readily visible in the IR spectra and the separate bands for the ¹⁰B/¹¹B isotopologues are resolved. A comparison of the spectra of different compounds reveals a clear correlation between the force constant and the effective partial charge of the $[BN_2]^{3-}$ anion.

Most recent studies on $[BN_2]^{3-}$ containing nitridoborates have concerned their luminescence properties. Polycrystalline samples of $Mg_3[BN_2]N$, 24 $Mg_{8-x}M_x[BN_2]_2N_4$ (M=Al, Si; x=1, 2) 25 and $Mg_3Ga[BN_2]N^{26}$ were studied with and without Eu^{2+} doping. These compounds show broad emission bands with orange-red to deep-red photoluminescence. In view of these interesting results we extended our work to the doping of compounds with the composition $MBa_8[BN_2]_6$ (M=Sr, Eu). The structures of both nitridoborates show a 50% occupancy of the strontium/europium site in order to account for an electron precise composition. 27 X-ray powder patterns of our first samples always showed additional reflections that could not be indexed on the basis of the body-centered cell, indicating superstructure formation. Herein we report on a single crystal

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study, revealing complete strontium/vacancy ordering within a $2 \times 2 \times 2$ superstructure cell.

2. Experimental

Synthesis

SrBa₈[BN₂]₆ was prepared by solid state reaction of stoichiometric quantities of Sr₃N₂ (Materion, 99.5%), Ba₃N₂ (Materion, 99.7%) and h-BN (Alfa Aesar, 99.5%). The binary nitrides were thoroughly ground in an agate mortar and arc-welded²⁸ in a niobium container under 800 mbar of argon, which was purified by a titanium sponge at 870 K, silica gel and molecular sieves. Subsequently, the niobium container was sealed in an evacuated silica tube for oxidation protection and heated at 1370 K for two hours, followed by slow cooling to room temperature at a rate of 20 K h⁻¹. The slightly yellowish sample was well crystallized and contained small transparent single crystalline platelets suitable for structure determination.

EuN, Ba₃N₂ and BN were used in stoichiometric ratios for the synthesis of polycrystalline EuBa₈[BN₂]₆. EuN was synthesized from europium metal (Treibacher, 99.9%), firstly preparing the hydride at 620 K for 2 h in a mixture of argon and hydrogen (Westfalen AG, Ar/W35), followed by subsequent nitridation under N2 (Westfalen AG, 5.0) at 770 K for 4 h. The starting materials were thoroughly ground in an agate mortar. The mixture was then transferred into BN crucibles, capped with a lid and calcined at 1270 K for 4 h under nitrogen. Both gases were purified using an "oxisorb"-cartridge (Spectron), which reduces the oxygen/moisture level to below five and thirty parts per billion, respectively. Further in situ gas purification was carried out using two crucibles filled with Mischmetal placed in front of and behind the reaction vessel. EuBa₈[BN₂]₆ was obtained as a dark purple to black polycrystalline powder, similar to LiEu₄[BN₂]₃,²¹ while the respective Sr compound was almost colorless. Although these two compounds are electron precise, the [Xe]4f⁷ configuration of Eu²⁺ along with low-lying 4f-5d transitions leads to complete absorption in the visible range. Two broad absorption bands are characteristic of the interconfigurational transition from $[Xe]4f^{7}$ to $[Xe]4f^{6}5d^{1}(t_{2g})$ and $[Xe]4f^{6}5d^{1}(e_{g})$ of Eu^{2+} in cubic crystals. 29,30

Due to the moisture-sensitive nature of the precursors and the final products, all handling was carried out in a nitrogen filled glovebox from GS Glovebox Systemtechnik with $\rm H_2O$ levels well below 50 ppm.

EDX data

The $SrBa_8[BN_2]_6$ single crystal studied on the diffractometer was semi-quantitatively investigated by EDX analyses using a Zeiss EVO® MA10 scanning electron microscope in the variable pressure mode with SrF_2 and BaF_2 as standards. The experimentally observed strontium: barium ratio was 21 ± 2 at %: 79 ± 2 at%, in close agreement with the ideal composition. The light elements could not be analysed owing to the limited

instrument resolution. No impurity elements heavier than sodium were observed.

X-ray diffraction

The polycrystalline $SrBa_8[BN_2]_6$ and $EuBa_8[BN_2]_6$ samples were characterized by X-ray powder diffraction using a Guinier camera equipped with an image plate system (Fujifilm, BAS-1800) using $CuK_{\alpha 1}$ radiation and α -quartz (a=491.30, c=540.46 pm) as an internal standard. The complete patterns could be indexed on the basis of large F-centred cubic cells and least squares refinements resulted in the lattice parameters of a=1595.1(1) pm for $SrBa_8[BN_2]_6$ and a=1595.00(9) pm for $EuBa_8[BN_2]_6$. Correct indexing of the patterns was ensured through intensity calculations using the LazyPulverix routine. The powder pattern of $SrBa_8[BN_2]_6$ is exemplarily presented in Fig. 1. The corresponding pattern of the europium compound is available in the ESI.†

Irregularly shaped crystal fragments of $SrBa_8[BN_2]_6$ were glued to thin silica fibers using beeswax. The sample was previously coated with paraffin oil (dried over sodium granules) for protection from hydrolysis. Subsequently the quality of the selected fragments was tested using Laue photographs on a Buerger camera (white molybdenum radiation, the image plate technique, Fujifilm, BAS-1800). Suitable crystals of $SrBa_8[BN_2]_6$ were stored in Schlenk tubes under an argon atmosphere prior to the data collection. Intensity data of the selected crystal were collected at room temperature under a constant nitrogen gas flow using a Stoe StadiVari diffractometer equipped with a Mo microfocus source (λ = 71.073 pm) and a Pilatus 100 K detector with a hybrid-pixel-sensor. A spherical absorption correction was applied to the data set. All relevant crystallographic data and details of the data collection and evaluation are given in Table 1.

Structure refinement

Careful analysis of the $SrBa_8[BN_2]_6$ data set revealed a facecentred cubic lattice and the systematic extinctions were in accordance with space group $Fd\bar{3}m$. The starting atomic para-

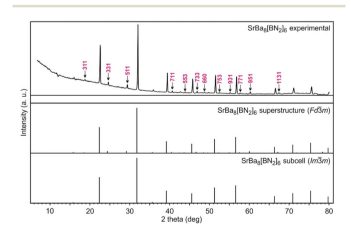


Fig. 1 Experimental (top) and calculated (middle and bottom) X-ray powder patterns of $SrBa_8[BN_2]_6$. The *hkl* indices of the superstructure reflections are marked in the experimental pattern.

Table 1 Crystallographic data and structure refinement for $SrBa_8[BN_2]_6$, space group $Fd\vec{3}m$, Z=8

Formula	$SrBa_8[BN_2]_6$
Molar mass/g mol ⁻¹	1419.3
Lattice parameter/pm (powder data)	a = 1595.1(1)
Cell volume/nm ³	V = 4.0585
Density calc./g cm ⁻³	4.65
Crystal size/µm	$30 \times 40 \times 40$
Detector distance/mm	60
Exposure time/s	43
Integr. param. A, B, EMS	7.0; -6.0; 0.030
h k l range	$\pm 23; \pm 23, \pm 23$
$\theta_{\min}, \theta_{\max} / \circ$	2.21/32.34
Linear absorption coeff./mm ⁻¹	17.9
No. of reflections	23 114
$R_{\rm int}$	0.0198
No. of independent reflections	387
Reflections used $[I \ge 3\sigma(I)]$	284
F(000), e	4800
R factors $R(F)/wR(F^2)$ $[I \ge 3\sigma(I)]$	0.0237/0.0453
R factors $R(F)/wR(F^2)$ [all data]	0.0450/0.0515
Data/parameters	387/21
Goodness-of-fit on F ²	0.96
Extinction coefficient	_
Diff. Fourier residues/e ⁻ Å ⁻³	-1.54/1.93

meters for the structure refinement were obtained using the charge-flipping algorithm Superflip. The structure was subsequently refined with anisotropic displacement parameters for all atoms on F^2 using Jana2006. As a check for the correct composition and site assignment, the occupancy parameters for all atoms were refined in separate least-squares cycles. All sites were fully occupied within three standard deviations. The setting of the structure was finally adjusted to that required for the group–subgroup scheme ($vide\ infra$). The final difference Fourier syntheses revealed no significant residues. The refined atomic positions, displacement parameters and interatomic distances are given in Tables 2–4.

Further details of the structure refinement may be obtained from the Fachinformationszentrum Karlsruhe, D-76344 Eggenstein-Leopoldshafen (Germany), by quoting Registry No. CSD-431340.

IR and Raman spectroscopy

For infrared (IR) spectroscopy, KBr specimens were prepared and analysed in the range of 450–2000 cm⁻¹ employing a Spec-

Table 2 Atom positions and equivalent isotropic displacement parameters (pm²) for $SrBa_8[BN_2]_6$. U_{eq} is defined as one-third of the trace of the orthogonalized U_{ij} tensor

Atom	Wyckoff position	X	у	z	$U_{ m eq}$
Sr	8 <i>a</i>	1/8	1/8	1/8	134(2)
Ba1	32 <i>e</i>	0.25833(2)	\boldsymbol{x}	\boldsymbol{x}	230(1)
Ba2	16 <i>c</i>	0	0	0	251(1)
Ba3	16 <i>d</i>	1/2	1/2	1/2	244(1)
В	48f	0.3671(6)	1/8	1/8	240(30)
N1	48 <i>f</i>	0.0344(5)	3/8	3/8	243(15)
N2	48f	0.2025(5)	3/8	3/8	257(15)

Table 3 Anisotropic displacement parameters (pm²) for SrBa₈[BN₂]₆. Coefficients U_{ij} of the anisotropic displacement factor tensor of the atoms are defined by: $-2\pi^2[(ha*)^2U_{11}+\cdots+2hka*b*U_{12}]$

Atom	U_{11}	U_{22}	U_{33}	U_{12}	U_{13}	U_{23}
Sr	134(4)	U_{11}	U_{11}	0	0	0
Ba1	230(2)	U_{11}	U_{11}	-23(1)	U_{12}	U_{12}
Ba2	250(2)	U_{11}	U_{11}	-61(2)	U_{12}	U_{12}
Ba3	244(2)	U_{11}	U_{11}	-22(2)	U_{12}	U_{12}
В	360(70)	180(30)	U_{22}	0	0	140(30)
N1	170(30)	280(20)	U_{22}	0	0	20(30)
N2	170(30)	300(20)	U_{22}	0	0	-10(30)
	. ,	. ,				. ,

Table 4 Interatomic distances (pm) for SrBa₈[BN₂]₆

Sr:	6	N1	254.3(8)	N1:	1	В	132(1)
Ba1:	3	N2	277.8(3)		1	Sr	254.3(8)
	3	N1	303.6(1)		2	Ba2	287.3(2)
Ba2:	6	N1	287.3(2)		2	Ba1	303.6(1)
Ba3:	6	N2	292.0(2)	N2:	1	В	136(1)
B:	1	N1	132(1)		2	Ba1	277.8(3)
	1	N2	136(1)		2	Ba3	292.0(2)
			. ,				. ,

trum 100 FT-IR spectrometer from Perkin Elmer equipped with the Spectrum 6.0.1 data processing software. The Raman spectrum of $SrBa_8[BN_2]_6$ was recorded using a Jobin Yvon HR 800 dispersive Raman spectrometer equipped with a green Nd: YAG laser ($\lambda = 532$ nm; P = 20 mW).

Solid state MAS NMR spectroscopy

¹¹B solid state MAS NMR spectra were recorded at magnetic field strengths of 4.7 and 11.7 T on NMR spectrometers at resonance frequencies of 64.166 and 160.419 MHz, respectively, using a Bruker DSX 500 spectrometer. Finely powdered nitridoborates were filled into ZrO₂ MAS rotors with 4 mm diameter and spun at a frequency of 12 kHz with nitrogen gas flow to prevent hydrolysis. Chemical shifts were referenced to BF₃·OEt₂ as the reference sample; π /2 pulse lengths of p_1 = 3.25–3.80 μs and a relaxation delay of d_1 = 5 s were used in conventional single-pulse experiments. The spectra were recorded using the Bruker Topspin software³⁴ and analyzed with the *Dmfit* software package.³⁵ The experimental results are given in Table 7.

3. Crystal chemistry

First hints for superstructure formation were already evident from the Guinier powder patterns which revealed weak additional reflections. The latter could be completely indexed with the $2\times2\times2$ superstructure cell for both compounds. As an example, the strongest superstructure reflections for SrBa₈[BN₂]₆ are marked with small arrows in Fig. 1. The Guinier pattern of EuBa₈[BN₂]₆ also shows all of the expected strong superstructure reflections so that both compounds can be considered isotypic. Crystal chemical details of the subcell structure have repeatedly been reported. Herein we focus

Dalton Transactions Paper

only on the crystal chemical discussion of the SrBa₈[BN₂]₆ superstructure which was refined from single crystal data.

The reason for superstructure formation lies in the electron-precise formulation. When sodium in the alkali metal compound NaBa₄[BN₂]₃ ¹⁸ is substituted by strontium, charge balance requires half of the strontium sites to remain unoccupied. A view of the SrBa₈[BN₂]₆ unit cell is presented in Fig. 2. The strontium-vacancy ordering is not simply realized through decentering of the body-centred cell (as is the case for the wellknown example bcc vs. CsCl type³⁶). The eight subcell strontium sites (50% occupied) around each strontium atom are ordered 4 + 4 in the superstructure, with both strontium and vacancies in a tetrahedral arrangement. This results in a doubling of the subcell in all three directions.

The resulting near-neighbor coordination for the strontium atoms and the $[BN_2]^{3-}$ units is presented in Fig. 3. The strontium atoms have end-on octahedral coordination by [BN₂]³⁻ units with Sr-N distances of 254 pm, similar to the shorter Sr-N distances in the structures of Sr_4N_3 (255–258 pm), ³⁷ SrN $(258-265 \text{ pm})^{38}$ and $Sr_2Si_5N_8$ $(254-272 \text{ pm})^{39}$

The most pronounced influence of the strontium-vacancy ordering is readily seen in the near-neighbor coordination of the $[BN_2]^{3-}$ units. In alkali metal compounds like $NaBa_4[BN_2]_3$, ¹⁸ each $[BN_2]^{3-}$ unit has a cation coordination in the form of a bi-capped square prism. Since only half of these sites are filled in the present strontium compound, only the N1 atoms have a strontium neighbor. Consequently we observe a lower number of cations coordinating with the N2 atoms, leading to drastically shortened N2-Ba distances of 278 and 292 pm as compared to the N1-Ba distances of 287 and 304 pm. The barium displacements are readily evident from Fig. 2.

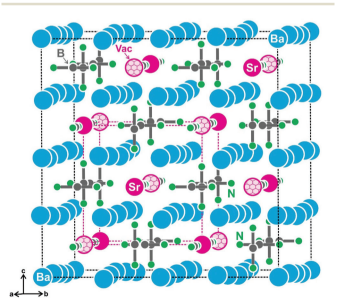


Fig. 2 The unit cell of SrBa₈[BN₂]₆. Strontium, barium, boron, and nitrogen atoms are drawn as magenta, blue, grey, and green circles, respectively. The [BN₂]³⁻ units are emphasized and ordered strontium vacancies are marked by honeycomb nets.

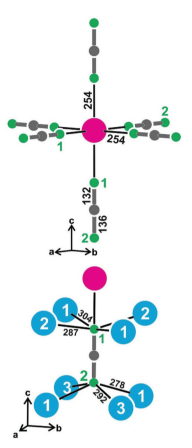


Fig. 3 Near-neighbor coordination of the strontium atoms and the $[BN_2]^{3-}$ units in the structure of $SrBa_8[BN_2]_6$. Strontium, barium, boron, and nitrogen atoms are drawn as magenta, blue, grey, and green circles, respectively. Relevant interatomic distances (pm) and atom designations are given.

The cation ordering leads to asymmetric (but linear) [BN₂]³units with interatomic distances of 132 pm for B-N1 and 136 pm for B-N2.

We have further analyzed the superstructure model of SrBa₈[BN₂]₆ through calculations of the bond valence sums (BVS)⁴⁰ and the Chardi algorithm.^{41,42} The refined numerical values are listed atom by atom in Table 5. The BVS data result from empirical parameters^{43,44} and show strong overbonding for the strontium atoms and strong underbonding for the Ba3 atoms. The charges are much better reproduced with the point-charge approach in Chardi, nicely underlining the elec-

Table 5 Charge distribution analysis of SrBa₈[BN₂]₆ using bond valence sums (BVS) and the Chardi algorithm

	Sr	Ba1	Ba2	Ва3	В	N1	N2
$\sum_{\text{Q (Chardi)}^{40}} \text{Q (Chardi)}^{40}$	2.57	1.96	2.02	1.78	2.85	-3.04	-2.81
	1.92	2.03	1.92	2.09	2.99	-3.13	-2.87

BV = $e^{(r_0-r)/b}$; b=0.37, r_0 (Sr^{II}–N) = 223 pm, r_0 (Ba^{II}–N) = 247 pm and r_0 (B^{III}–N) = 147 pm. ^{43,44}

Paper

bonded $[BN_2]^{3-}$ units.

tron-precise description. This holds also for the covalently

The SrBa₈[BN₂]₆ structure is related with the LiCa₄[BN₂]₃ subcell type through a group-subgroup relation, which is presented in the Bärnighausen formalism⁴⁵⁻⁴⁷ in Fig. 4. The symmetry reduction proceeds in two steps. First we observe a decentering of the body-centred lattice through a klassengleiche transition of index 2 (k2) from Im3m to Pn3m. Although this symmetry reduction leads to splitting of the 8c calcium site, there is still no possibility of ordering of the strontium atoms for the title compound. Subsequently, a second klassengleiche transition of index 2 (k2) from $Pn\bar{3}m$ to $Fd\bar{3}m$ upon doubling of the unit cell in all three directions proceeds and the splitting of the 2a site into two eightfold sites in the superstructure allows for strontium-vacancy ordering. The klassengleiche transitions lead to the superstructure reflections observed in the powder patterns. The symmetry reductions also lead to free x parameters for the Ba1 and B atoms as well as two different x parameters for N1 and N2.

Ordering of cation vacancies has also been determined for the β -Ca₃[BN₂]₂ structure,⁷ but with a different symmetry reduction. This superstructure variant crystallizes with the orthorhombic space group *Cmce*. The smaller calcium atoms and the different pattern of vacancy ordering lead to signifi-

cant tilts and bending of the $[BN_2]^{3-}$ units. For comparison, projections of the $SrBa_8[BN_2]_6$ and β - $Ca_3[BN_2]_2$ structures are presented in Fig. 5.

4. Vibrational spectroscopy

Linear [BN₂]³⁻ nitridoborate anions are the characteristic structural units of SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆. The numerous vibrational modes can be easily detected in either IR and/or Raman spectra. The actual number of modes that can be expected in the solid, dependent on the exact symmetry of the dumbbell-shaped [BN2]3- units, has been discussed in detail before for several similar compounds.^{2,9,12,23,48} Fig. 6 depicts the IR spectra of SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆. The two isotypic nitridoborates exhibit almost identical spectra. The numerical values of the frequencies are listed in Table 6. IR-active, characteristic anti-symmetric stretching modes v_2 of the B-N bonds can be observed in the range between 1600 and 1700 cm⁻¹ (band splitting is a consequence of factor group splitting) and lower-lying doubly degenerate deformation modes v_3 from 500 to 650 cm⁻¹. Additional splitting of bands is due to the presence of ¹¹B and ¹⁰B isotopologues at the natural abundance ratio of 80:20. The assumption of asym-

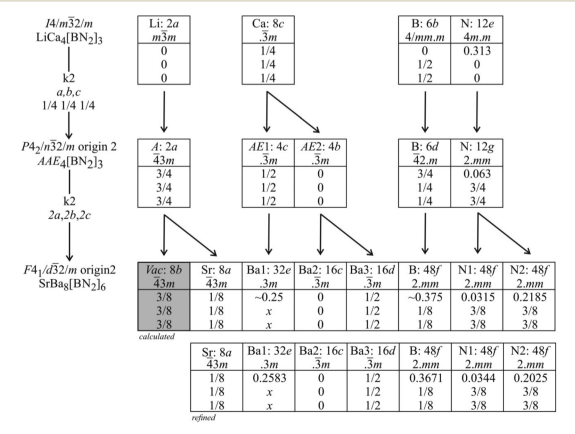


Fig. 4 Group–subgroup scheme in the Bärnighausen formalism $^{45-47}$ for the structures of LiCa₄[BN₂]₃ and SrBa₈[BN₂]₆. The indices for the *klassen-gleiche* (k) symmetry reductions, the unit cell transformations, and the evolution of the atomic parameters are given. No representative is known for the intermediate space group $Pn\bar{3}m$. The alkali and alkaline earth sites for that intermediate are symbolized by A and AE, respectively.

Dalton Transactions Paper

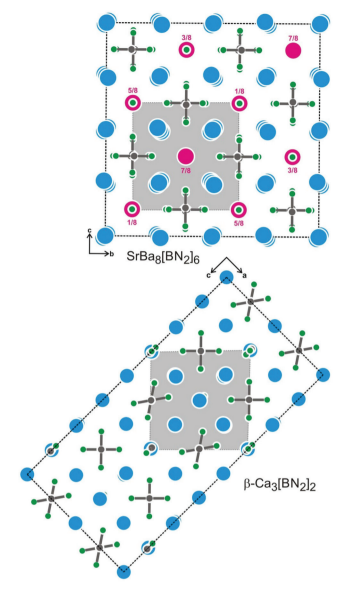


Fig. 5 Projections of the $SrBa_8[BN_2]_6$ and β - $Ca_3[BN_2]_2$ structures. Heights of the strontium atoms in SrBa₈[BN₂]₆ are indicated. Both superstructures are a consequence of vacancy ordering. For further details of the vacancy ordering in β -Ca₃[BN₂]₂, we refer the reader to ref. 7.

metric [BN₂]³⁻ units as confirmed by the crystallographic data is underlined by the presence of the symmetric ν_1 stretching mode in the FTIR spectrum. Further confirmation of the broken linear symmetry, due to different B-N bond lengths, is given by the doublet splitting of the ν_3 deformation modes.

The Raman spectrum of SrBa₈[BN₂]₆ is presented in Fig. 7. EuBa₈[BN₂]₆ was also tested; however, similarly to $\text{LiEu}_{4}[\text{BN}_{2}]_{3}^{23}$ it decomposed during laser exposure. The symmetric B-N stretching splits into two bands in the Raman spectrum at 1024 and 1030 cm⁻¹, similar to the situation in Ca₃[BN₂]I⁴⁸ that also shows two different B-N bond lengths of 139.3 and 136.9 pm.

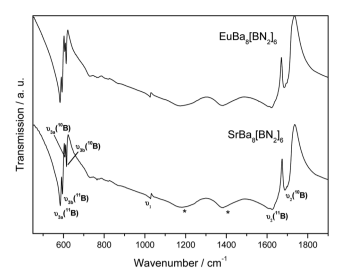


Fig. 6 IR spectra of SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆. Relevant modes are indicated. For numerical values see Table 6. The modes marked with asterisks result from slow decomposition of the samples.

Table 6 Frequencies (cm⁻¹) of the fundamental modes of the IR spectra (Fig. 6) of SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆

	v_1	$ \stackrel{\upsilon_2}{(^{11}\mathrm{B})}$	$ \stackrel{\upsilon_2}{(^{10}{\rm B})}$	v_{3a} $\binom{11}{B}$	v_{3b} $\binom{11}{B}$	v_{3a} $\binom{10}{B}$	v_{3b} (^{10}B)
SrBa ₈ [BN ₂] ₆	1028	1625	1687	585	595	607	615
EuBa ₈ [BN ₂] ₆	1025	1622	1684	583	593	606	613

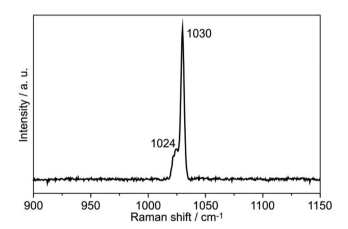


Fig. 7 Raman spectrum of SrBa₈[BN₂]₆.

¹¹B solid state MAS NMR spectroscopy

The ¹¹B solid state MAS NMR spectra of the nitridoborates SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆ are shown in Fig. 8 and 9, respectively. Consistent with the crystal structure the spectra show only one signal whose line shape is strongly influenced by second order quadrupolar perturbations induced by the

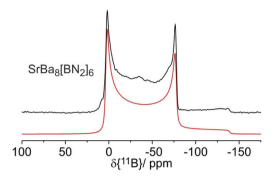


Fig. 8 11 B MAS NMR spectrum of SrBa₈[BN₂]₆ with simulation (red line) recorded at an external magnetic flux density of $B_0 = 4.7$ T and a spinning frequency of 12 kHz.

interaction of the 11 B nuclei with the electric field gradient resulting from their non-spherical electronic environment. Table 7 summarizes the quadrupolar coupling constant $C_{\rm Q}$ and the asymmetry parameter $\eta_{\rm Q}$ obtained by fitting the line shape of the central transition. Consistent with the $C_{\rm cov}$ local symmetry at the boron site, the field gradient is found to be close to axially symmetric in both compounds. The spectra of the Eu²⁺ compound are profoundly influenced by the proximity of the unpaired 4f-electrons to the 11 B nuclei. This paramagnetic interaction manifests itself in (1) a broadening effect

Table 7 11 B NMR spectroscopic parameters of the nitridoborates SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆: isotropic resonance shifts $\delta_{\rm iso}$ (±1 ppm), quadrupolar coupling constant $C_{\rm Q}$ (±0.05 MHz) and electric field gradient asymmetry parameter $\eta_{\rm CQ}$ (±0.03), magnetic flux density of external field $B_{\rm Q}$ (T) and line broadening parameter LB (±1.0 ppm)

	$\delta_{ m iso}$	$C_{ m Q}$	η_{CQ}	B_0	LB
SrBa ₈ [BN ₂] ₆	26.6	3.30	0.02	4.7	2.0
$EuBa_8[BN_2]_6$	26.6	3.10	0.03	4.7	15.2
	24.3	3.10	0.03	11.7	7.0

imposed on the second-order quadrupolar powder pattern characterizing the central $1/2 \leftrightarrow -1/2$ transition and (2) the appearance of intense MAS sidebands, reflecting strong inhomogeneous spectral broadening effects. The latter arise from the anisotropic effective local field generated at the site of the ¹¹B nuclei by proximal electron spins, which are rapidly fluctuating between their Zeeman states. For the quadrupolar ¹¹B nuclei this interaction is superimposed on the anisotropy of the nuclear electric quadrupolar interaction affecting the $\pm 1/2 \leftrightarrow \pm 3/2$ "satellite" Zeeman transitions. Fig. 9 shows the spinning sideband profiles at the external magnetic flux densities of 4.7 and 11.7 T. A detailed analysis of these sideband profiles requires additional experimental inputs from EPR spectroscopy and will be presented in a forthcoming study.

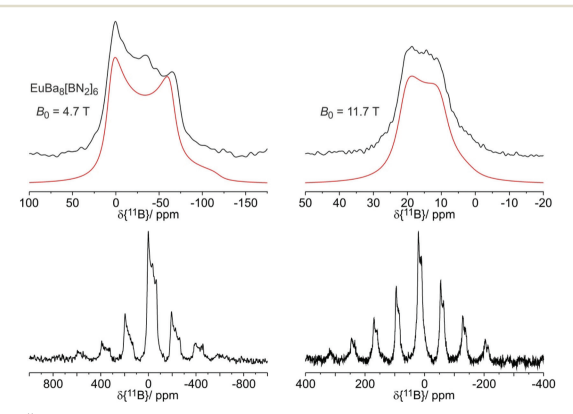


Fig. 9 Enlarged 11 B MAS NMR spectra of EuBa₈[BN₂]₆ recorded at external magnetic flux densities of $B_0 = 4.7$ T (top left) and $B_0 = 11.7$ T (top right) with simulated spectra (red lines) and full spectra with spinning side-band patterns (bottom).

6. Conclusions and outlook

High-resolution powder and single crystal X-ray diffraction data confirm full cation ordering in the nitridoborates SrBa₈[BN₂]₆ and EuBa₈[BN₂]₆. Both compounds crystallize with a new superstructure that derives from the LiCa₄[BN₂]₃ type through a group-subgroup scheme. The cation ordering leads to asymmetric linear [BN₂]³⁻ units with B-N distances of 132 and 136 pm. This is underlined by vibrational spectroscopy. The ¹¹B solid state MAS NMR spectra confirm the presence of single boron atomic sites in both compounds, whose electronic environment is characterized by axially symmetric electric field gradients. The spectrum of the Eu compound is additionally influenced by strong anisotropic paramagnetic interactions, resulting in characteristic field-dependent spectra. SrBa₈[BN₂]₆ is an excellent host structure for doping with divalent or trivalent rare earth cations. Detailed optical, solid state NMR and ESR studies will be published shortly.

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Paper

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