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# The contribution of phonons to the thermal expansion of some simple cubic hexaboride structures: SmB<sub>6</sub>, CaB<sub>6</sub>, SrB<sub>6</sub> and BaB<sub>6</sub>†

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We have performed first-principles calculations of the structure and lattice dynamics in the metal hexaborides SmB<sub>6</sub>, CaB<sub>6</sub>, SrB<sub>6</sub> and BaB<sub>6</sub> using Density Functional Theory in an attempt to understand the negative thermal expansion in the first of these materials. The focus is on the role of Rigid Unit Modes involving rotations of the B<sub>6</sub> octahedra similar to the rotations of structural polyhedra connected by bonds in Zn(CN)<sub>2</sub>, Prussian Blue and Si(NCN)<sub>2</sub>. However, it was found that there is very low flexibility of the network of connected B<sub>6</sub> octahedra, and the lattice dynamics do not support negative thermal expansion except possibly at very low temperature. Thus the negative thermal expansion observed in SmB<sub>6</sub> probably has an electronic origin.

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### 1 Introduction

The simple cubic hexaborides XB6, which are formed from some transition metals (X = Sm, Y, La for example) and some alkali earth elements (X = Ca, Sr, Ba) are shown in Fig. 1.<sup>1,2</sup> The crystal structure consists of B6 octahedra, linked together by B-B bonds. This is similar to the cubic perovskite structure, but without the octahedral cation, and more importantly with the shared oxygen atom replaced by a B-B bond. In this sense it is like Zn(CN)<sub>2</sub>, which has Zn-centred tetrahedra connected by shared CN bonds, and even more similar to analogues of Prussian Blue in which metal-centred octahedra are connected by shared CN bonds in exactly the same topological layout.<sup>4,5</sup> The main difference is that the essential network in SmB<sub>6</sub> is entirely made from boron atoms, rather than metal-centred oxide or halide polyhedra, and thus cubic metal hexaboride materials potentially represent a new type of network material.

Over the past two decades we have seen a rapid increase in the number of conventional network materials that show negative thermal expansion (NTE), that is, over a range of

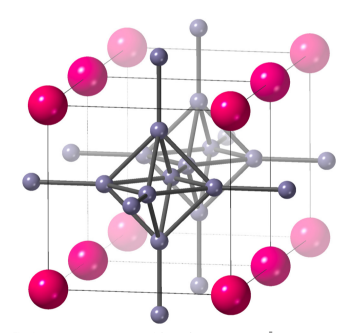


Fig. 1 The crystal structure of SmB<sub>6</sub> (space group Pm3̄m, lattice parameter 4.134 Å at temperature of 20 K). B atoms are the small grey bonded spheres, and Sm atoms are the larger pink atoms at the corners of the unit cell.

temperatures the material will shrink rather than expand on heating.<sup>6-9</sup> Many of these materials are oxides or analogues based on replacing the oxygen anion by a halogen anion or even a molecular anion.10 In these cases there are structural polyhedra that are connected at corners to form infinite networks. Examples are ReO<sub>3</sub><sup>11,12</sup> and its analogue ScF<sub>3</sub>, <sup>13,14</sup> Zn(CN)<sub>2</sub><sup>3</sup> as

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an analogue of the cubic cristobalite phase of SiO2, and the most famous NTE material ZrW2O815 with corner linked WO4 tetrahedra and ZrO6 octahedra.

NTE in network materials is commonly associated with a mechanism known as the 'tension effect'. 8 One simple example is cubic ScF<sub>3</sub>. <sup>16</sup> The crystal structure has cubic symmetry, and consists of corner-sharing ScF<sub>6</sub> octahedra with linear Sc-F-Sc bonds – essentially it has the cubic perovskite structure without the 12-coordinated cation. The Sc-F bonds are relatively rigid (and have positive thermal expansion), and so rotations of the Sc-F bonds will necessarily reduce the corresponding Sc-Sc distances and hence will lead to overall reduction in the crystal size. Such rotations occur as thermally-excited vibrational motions, and their effect in reducing the Sc-Sc distance exceeds that from the thermal expansion of the Sc-F bonds. 14,16 Thus the effect becomes larger with higher temperature, leading directly to NTE. Because atomic motions are correlated, the important question is the extent to which the structure allows for low-frequency rotational vibrations. In structures consisting of connected polyhedra, the question becomes focussed on the balance between the phonons in which the polyhedra rotate as nearly-rigid objects - the Rigid Unit Modes (RUMs)17,18 - and the phonons that lead to bending of the bonds of polyhedra. This point has been discussed in detail elsewhere.<sup>19</sup>

Of the metal hexaborides, as far as we can tell lowtemperature lattice parameters have only been measured for SmB<sub>6</sub>, which itself is now best known as a topological insulator.20-28 Experimental data show the existence of NTE in SmB<sub>6</sub> at low temperatures, and we have recently confirmed this with our own diffraction data shown in Fig. 2. On the other hand, there are higher-temperature data for other metal hexaborides, which all show positive thermal expansion above room temperature.2

In this paper we investigate the role of phonons and the standard Grüneisen and tension effect mechanisms<sup>8</sup> using density functional theory simulations of SmB6 and analogue

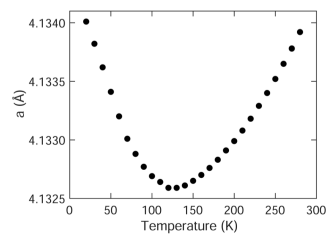


Fig. 2 Unpublished experimental data for the temperature-dependence of the lattice parameter of SmB<sub>6</sub>, obtained by the corresponding author with Zhongsheng Wei and Dean Keeble at the XPDF beam line at the Diamond synchrotron facility in the UK.

materials with alkaline earth cations, CaB<sub>6</sub>, SrB<sub>6</sub> and BaB<sub>6</sub>. In fact it will be shown here that although there is a phonon mechanism to generate NTE, it is not sufficient in these materials to give NTE over a wide range of temperatures. In the case of SmB<sub>6</sub> it is impossible to reproduce the observed NTE from a phonon mechanism, and thus we will conclude that the mechanism for NTE in SmB6 may more likely have an electronic origin.

### 2 Methods

#### 2.1 DFT methods

Simulations were performed using the CASTEP program,<sup>29</sup> version 19. This uses a density-functional-theory method based on periodic structures that make use of plane-wave basis sets. The effects of core electrons were handled using normconserving pseudopotentials supplied as part of the CASTEP package (NCP19).

The calculations were performed using the standard PBE generalised-gradient approximation30,31 for CaB6, SrB6 and BaB<sub>6</sub>, assuming the electronic states are those of an insulator. The integration of the electronic states was performed using a Monkhorst-Pack grid<sup>32</sup> of  $7 \times 7 \times 7$  wave vectors. A cut-off energy of 800 eV was chosen for the plane wave electronic basis set based on convergence tests for energy differences. For structure optimisation, calculations were performed to a convergence of energies to 10<sup>-8</sup> eV per atom, convergence of force to  $10^{-4}$  eV  $\text{Å}^{-1}$ , and convergence of stress to  $10^{-4}$  GPa. All limits were chosen on the basis of tests on convergence of energy differences for two similar volumes.

For the calculations on SmB<sub>6</sub>, on the other hand, the system was treated as a metal. A cut-off energy of 900 eV was used for the electronic basis set. Other details were the same as for the other materials.

We did not consider contributions from magnetic spin moments because it has been shown that SmB<sub>6</sub> remains nonmagnetic down to very low temperatures.<sup>21</sup>

#### 2.2 Structure optimisation

The optimised values of the lattice parameters and interatomic distances for all systems are compared with experimental data, 1,33 in Table 1. In each case agreement on lattice parameters is good, to within 1%.

The bond distances are shown in Fig. 3, where we are able to make one important point. Increasing the size of the metal atom clearly, from Table 1, creates some strain in the crystal structure. This strain is accommodated mostly by expansion of the length of the B-B bond connecting two octahedra, and to a much lesser extent by increasing the size of the octahedra.

#### 2.3 Phonon methods

Phonon frequencies for the alkali earth hexaborides were calculated using the density functional perturbation theory (DFPT) method.34,35 An interpolation method was used to compute frequencies for any wave vector based on calculations of the

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Table 1 Calculated and experimental crystal structure data for the four hexaborites examined in this study. a is the cubic lattice parameter, and x is the fractional coordinate of the boron atom. Bond length values are in Å. B-B (1) is the bondlength within the  $B_6$  octahedra, and B-B (2) is the bondlength between octahedra. M-B is the bond length between the metal cation and the nearest boron atom. Experimental data from SmB<sub>6</sub> are from the neutron diffraction measurements of Trounov et al. measured at a temperature of 23 K, and for the other phase data are taken from a compilation of Schmitt et al. 33 for measurements performed at room

	a (Å)	В х	В-В (1)	B-B (2)	М-В
SmB <sub>6</sub> exp	4.133	0.200	1.755	1.652	3.037
SmB <sub>6</sub> calc	4.129	0.200	1.752	1.652	3.034
CaB <sub>6</sub> exp	4.152	0.202	1.750	1.677	3.053
CaB <sub>6</sub> calc	4.122	0.203	1.732	1.673	3.032
SrB <sub>6</sub> exp	4.198	0.203	1.764	1.704	3.088
SrB <sub>6</sub> calc	4.194	0.203	1.762	1.702	3.085
BaB <sub>6</sub> exp	4.276	0.205	1.784	1.753	3.148
BaB <sub>6</sub> calc	4.262	0.206	1.775	1.752	3.138

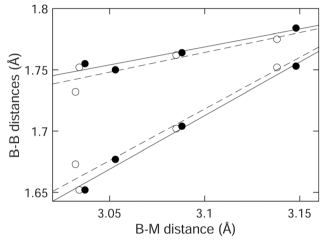


Fig. 3 Comparison of the two experimental<sup>1,33</sup> (filled circles) and calculated (open circles) B-B distances in the four hexaborides, plotted as a function of the metal (M) to boron distance. From left to right the metal atoms are Sm, Ca, Sr and Ba. Note that there is a close overlap of the experimental and calculated distances in the case of the SrB<sub>6</sub>. The upper data are for the B-B distances within the octahedra and the lower data are for B-B distances connecting two octahedra. The lines are fits to the data, and highlight the fact that most of the strain imposed by increasing the size of the metal atom is accommodated though expansion of the B-B bond connecting two octahedra

dynamical matrices performed for a Monkhorst-Pack grid of wave vectors of size  $7 \times 7 \times 7$  in reciprocal space. A convergence tolerance for force constants during the DFPT calculations of  $10^{-5}$  eV  $\text{Å}^{-2}$  was used. The phonon acoustic sum rule was enforced. For SmB<sub>6</sub> the dynamical matrix was calculated using a supercell method. We have checked in the case of CaB<sub>6</sub> that the two methods give substantially the same results.

Volumetric thermal expansion<sup>8</sup> is given by the formula  $\alpha_{\rm V} = C_{\rm V} \bar{\gamma}/BV$ , where  $C_{\rm V}$  is the heat capacity at constant volume, and  $B = V \partial P / \partial V$  is the bulk modulus. The key quantity is the overall Grüneisen parameter  $\bar{\gamma}$ , which can be constructed from the individual mode Grüneisen parameters defined as  $\gamma_i = (\partial \omega_i / \partial V) \times (V/\omega_i)$ , where  $\omega_i$  is the angular frequency of the phonon labelled i and V is the crystal volume. The overall Grüneisen parameter is formed as

$$\overline{\gamma} = \frac{1}{C_{V}} \sum_{i} \hbar \omega_{i} \frac{\partial n(\omega_{i}, T)}{\partial T} \gamma_{i}$$
 (1)

where  $n(\omega_i, T)$  is the normal Bose–Einstein distribution.

For each system a set of calculations was performed with the crystal structure in its lowest-energy configuration, then a second set of calculations was performed using a lattice parameter expanded by about 1%. Mode Grüneisen parameters were obtained as  $\gamma_i = (\Delta \omega_i / \omega_i) / (\Delta V / V)$  from lattice dynamics calculations using a set of random wave vectors. An eigenvectormatching algorithm36 was used to ensure that the correct pairs of modes were compared in the calculation of  $\Delta\omega_i$ .  $C_V$  was calculated from the phonon frequencies using the standard formula based on differentiation of the Bose-Einstein distribution function.

## 3 Phonon dispersion curves of CaB<sub>6</sub> and SmB<sub>6</sub>

Here we consider the main details of the phonon dispersion curves. These are shown for SmB6 and CaB6 along the main symmetry directions in reciprocal space in Fig. 4. Corresponding data for SrB<sub>6</sub> and BaB<sub>6</sub> are given in the ESI.†

At zero wave vector, the mode decomposition gives acoustic modes with irreducible representation  $T_{1u}\text{, and }A_{1g}$  +  $E_g$  +  $T_{1g}$  +  $T_{2g} + 2T_{1u} + T_{2u}$  for the optic modes. In CaB<sub>6</sub> the two optic  $T_{1u}$ modes show splitting of the frequencies of the longitudinal and transverse modes, which is not present in SmB<sub>6</sub> because in this case it is a metal. The values of frequencies at zero wave vector are given in Table 2.

Images of the mode eigenvectors are shown in the ESI.† The lowest frequency T<sub>1u</sub> mode involves simple opposite displacements of the metal cation and B<sub>6</sub> octahedron, creating a local electrical dipole moment. The T<sub>2u</sub> mode consists of sideways displacements of the B-B bond connecting two octahedra in the perpendicular direction. This necessarily causes deformation of the B<sub>6</sub> octahedra. The T<sub>1g</sub> mode involves the rotation of the B<sub>6</sub> octahedra without distortion (see the discussion in Section 4), together with a counter rotation of the linking B-B bonds. The higher-frequency T<sub>1u</sub> mode involves displacements of the linking B-B bond along the direction of the bond, causing bond-bending distortions of the B<sub>6</sub> octahedra. The E<sub>g</sub> mode involves asymmetric stretching of the B-B bonds within the B<sub>6</sub> octahedra. Finally, the A<sub>1g</sub> mode is the totally-symmetric stretch of all bonds within the B<sub>6</sub> octahedra accompanied by an opposite contraction of the linkage B-B bonds.

The low-frequency part of the dispersion curves of SmB<sub>6</sub> have been measured by inelastic neutron scattering along the three symmetry directions from zero wave vector at room temperature.<sup>37</sup> The measurements included the acoustic modes in each direction, and the lowest set of optic modes that come together as the T<sub>1u</sub> triplet at just below 5 THz at zero wave vector. Agreement between calculation and experiment is

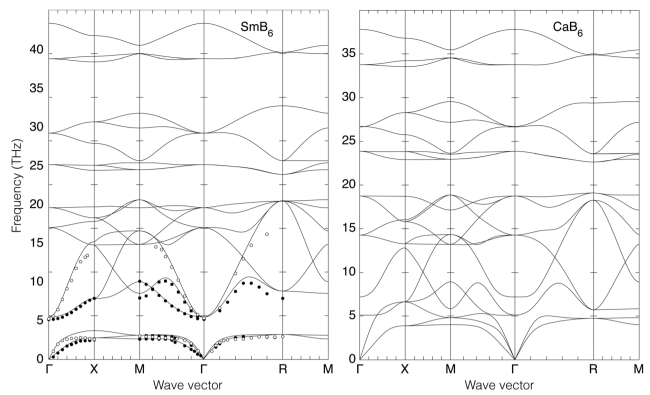


Fig. 4 Calculated phonon dispersion curves of SmB<sub>6</sub> (left) and CaB<sub>6</sub> (right). The wave vector labels have conventional meaning:  $\Gamma$  represents the wave vector (0, 0, 0), X represents  $(\frac{1}{2}, 0, 0)$ , M represents  $(\frac{1}{2}, \frac{1}{2}, 0)$  and R represents  $(\frac{1}{2}, \frac{1}{2}, \frac{1}{2})$ . Results from a single-crystal neutron inelastic scattering experiment<sup>37</sup> are represented as points, using different symbols to aid clarity.

Table 2 Calculated phonon frequencies for the four metal hexaborides of this study, SmB<sub>6</sub>, CaB<sub>6</sub>, SrB<sub>6</sub> and BaB<sub>6</sub> at zero wave vector, in units of THz. In each case other than SmB<sub>6</sub>, which are ionic insulators, the T<sub>1u</sub> modes show longitudinal/transverse splitting, and we give the frequencies of both components together. As a metal, SmB<sub>6</sub> does not show this splitting

Mode	$SmB_6$	CaB <sub>6</sub>	$SrB_6$	$BaB_6$
$\overline{T_{1u}}$	4.86	5.15, 7.22	4.68, 6.27	5.63, 6.79
$T_{2u}$	15.11	14.32	14.10	13.80
$T_{1g}$	17.37	18.76	18.41	18.52
$\begin{array}{c} T_{1g} \\ T_{2g} \end{array}$	22.32	23.93	23.31	22.50
$T_{1u}$	25.94	26.73, 26.80	25.84, 25.90	24.86, 24.93
$E_g$	34.44	33.94	32.33	29.72
$A_{1g}$	38.50	37.99	36.33	33.54

excellent, including the lack of LO/TO splitting of the T<sub>1u</sub> modes at zero wave vector, the significant flattening of acoustic mode dispersion curves, the subtle cross-over of the transverse and longitudinal acoustic modes for wave vectors around  $(\frac{1}{4}, \frac{1}{4}, 0)$  (half way between the points labelled  $\Gamma$  and M in Fig. 4), the rapid rise of the acoustic modes in increasing wave vector from zero wave vector  $(\Gamma)$ , and the maxima in the frequencies of the low-frequency optic modes at the points  $(\frac{1}{4}, \frac{1}{4}, 0)$  (half way between  $\Gamma$  and M) and  $(\frac{1}{4}, \frac{1}{4}, \frac{1}{4})$  (half way between  $\Gamma$  and R). The calculated frequencies of the acoustic modes are in excellent agreement with experiment across the whole Brillouin zone for all three directions. The lowest frequency TO modes agree very well with experiment, including the values of the maximum frequencies. The maximum frequencies of the low-frequency LO modes are calculated to have slightly higher frequencies than experiment at the zone boundaries, but the fact that the frequency of the LO mode at wave vector X is lower than that at wave vector M, which in turn is lower than that at wave vector R, is reproduced in the calculation. On the basis of this level of agreement between calculation and experiment we conclude that our DFT approach is reproducing most of the relevant behaviour of the phonon dispersion curves.

For SmB<sub>6</sub> and CaB<sub>6</sub> we can compare the calculated frequencies for zero wave vector with the results from Raman spectroscopy, 25,38 as summarised in Table 3. We consider the agreement to be good.

It is interesting to draw attention to the phonon anti-crossing at low frequencies, involving the acoustic and low-lying optic modes.

Table 3 Experimental and calculated Raman frequencies, given in the units of measurement, cm<sup>-1</sup>. Experimental data for SmB<sub>6</sub> are from Nyhus et al.<sup>25</sup> (older data are given by Mörke et al.<sup>39</sup>), and were measured at a temperature of 300 K. Experimental data for CaB<sub>6</sub> are from Ogita et al., <sup>38</sup> and were measured at a temperature of 13 K. The exact agreements of pairs of values are coincidental

Mode	SmB <sub>6</sub> Expt	SmB <sub>6</sub> Calc	CaB <sub>6</sub> Expt	CaB <sub>6</sub> Calc
A <sub>1g</sub>	1280	1284	1291	1264
$E_{\alpha}$	1148	1148	1149	1129
$egin{array}{l} A_{1g} \ E_{g} \ T_{2g} \end{array}$	730	743	780	780

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**Table 4** Values of elastic constants  $C_{11}$ ,  $C_{44}$  and  $C_{12}$ , and bulk modulus B, deduced from the slopes of the acoustic modes in the limit of small wave vector as described in the ESI

	$C_{11}$ (GPa)	$C_{44}$ (GPa)	$C_{12}$ (GPa)	B (GPa)
SmB <sub>6</sub>	440	71	22	161
CaB <sub>6</sub>	433	54	14	153
$SrB_6$	394	66	52	166
$BaB_6$	418	98	34	162

This is particularly evident in the directions  $\Gamma$ –X, where as a result both the transverse and longitudinal acoustic modes are seen to flatten quickly on increasing wave vector, and the optic modes rise sharply. The effect is much larger in SmB<sub>6</sub> than in CaB<sub>6</sub>, leading to lower frequencies at the X point in SmB<sub>6</sub>. The same effect is seen in other directions  $\Gamma$ –M and  $\Gamma$ –R. It is noticeable that in each case the extrapolation to the zone boundary of the un-crossed acoustic modes gives similar frequencies.

It is also noticeable from Table 2 that there is a significant and consistent decrease in the calculated value of the A<sub>1g</sub> symmetric stretch mode from SmB<sub>6</sub> to BaB<sub>6</sub>. This is commensurate with a change in B-B bond lengths (Table 1), that is the longer bond has a weaker force constant. However, this trend is not seen in the independent data for SmB<sub>6</sub> and CaB<sub>6</sub> shown in Table 3, for reasons that are not clear.

The slopes of the acoustic modes were used to calculate the elastic constants and bulk modulus. The method is described in the ESI,† and key data are provided there. The results are shown in Table 4. The results for SmB<sub>6</sub>, CaB<sub>6</sub> and SrB<sub>6</sub> are consistent with results given by the Materials Project as computed from analysis using finite strains, although data for BaB<sub>6</sub> are not available there.

## 4 Flexibility analysis

The Rigid Unit Modes (RUMs)17,18 discussed previously in regard to framework structures composed of corner-linked structural polyhedra, such as silicate and perovskite structures, are the phonons that can propagate without any distortion of the polyhedra. In the case of oxides with corner-sharing polyhedra, the RUMs typically propagate with low frequency. The concept of RUMs can be extended to systems composed of structural polyhedra connected via a shared molecular ligand, as in the related materials Zn(CN)23,40 and Si(NCN)2,41 or analogues of Prussian Blue,  $\mathrm{Fe_4^{\,III}}\ [\mathrm{Fe^{II}}(\mathrm{CN})_6]_3.^{40}$  However, as was found in the case of Zn(CN)2, the RUMs that involve bending of the linear Zn-CN-Zn linkage have relatively high frequency, so one should not assume that in these extended network systems the RUMs will have low frequencies,3 which is the pertinent issue for the cubic metal hexaborides.

We have evaluated all possible RUMs for the cubic metal hexaboride structure, based on rigid-body motions of the B<sub>6</sub> octahedra and with a rigid connecting B-B bond, and disregarding any role of the metal atoms. We begin with a simple count of degrees of freedom and constraints. Each octahedron has 6 degrees of freedom. Each connecting B-B provides one

Table 5 Number of RUMs for special wave vectors. For all other wave vectors, including the general wave vector without symmetry, the number of RUMs is equal to 3

Wave vector	Number of RUMs	
(0, 0, 0)	6	
$(\xi, 0, 0)$	5	
(1/2, 0, 0)	5	
$(1/2,  \xi,  0)$	4	
$(\xi, \xi, 0)$	4	
(1/2, 1/2, 0)	4	
$(\xi, \zeta, 0)$	4	

constraint, which is shared between the two connected octahedra. Thus we have 3 constraints for each octahedron. The difference between the numbers of degrees of freedom and constraints implies there are three RUMs for each wave vector, a result confirmed by calculation using a dynamical matrix. It is often found that symmetry leads to the presence of additional RUMs at special wave vectors, and this was found to be the case here using a simple flexibility mode (discussed below). The key results are given in Table 5, and are consistent with those obtained by Goodwin<sup>40</sup> for the analogous Prussian Blue crystal.

The additional three RUMs at wave vector  $\mathbf{k} = (0, 0, 0)$  are the trivial acoustic modes. The rest of the results given in Table 5 can be accounted for simply by noticing that there are planes of wave vectors of the form  $(\xi, \zeta, 0)$  containing one additional RUM each. Thus, for example, the wave vector  $(\xi, 0, 0)$  lies on the intersection of two such planes and thus has an additional two RUMs, as seen in Table 5. This mode is a shear acoustic mode.

To map the RUMs onto the phonon dispersion curves, we use the rigidity analysis described previously by Rimmer and Dove.<sup>36</sup> The eigenvectors from a full lattice dynamics calculation are compared with those from a calculation with a model designed to reproduce the spectrum of RUMs. The calculations were performed using the GULP simulation package. 42,43 Strong interatomic distance potentials were applied to the B-B bonds, and bond-bending potentials were applied to the B-B-B angles within the octahedra. No other potentials were applied to the network, meaning that the octahedra can twist about the connecting B-B bonds with no energy cost. The metal atoms were not included in the model.

For a given phonon wave vector  $\mathbf{k}$  and label i, we form the comparison of the phonon eigenvector  $\mathbf{e}_i(\mathbf{k})$  from the full lattice dynamics calculation with the eigenvectors  $\mathbf{e}'_{i}(\mathbf{k})$  for the flexibility model through the vector product relationship defined by Rimmer and Dove:36

$$m_i(\mathbf{k}) = \Omega^2 \sum_j \frac{\left| \mathbf{e}_i(\mathbf{k}) \cdot \mathbf{e}'_j(\mathbf{k}) \right|}{\Omega^2 + \omega_j^{'2}(\mathbf{k})}$$
(2)

where  $\omega_i'(\mathbf{k})$  is the (angular) frequency of the *j*-th mode in the model calculation, the sum is over all modes in the model calculation, and  $\Omega$  is an arbitrary value designed simply to prevent divergence when  $\omega'_i(\mathbf{k}) \simeq 0$ , but because we needed to include forces for the motion of the metal atom the RUMs will

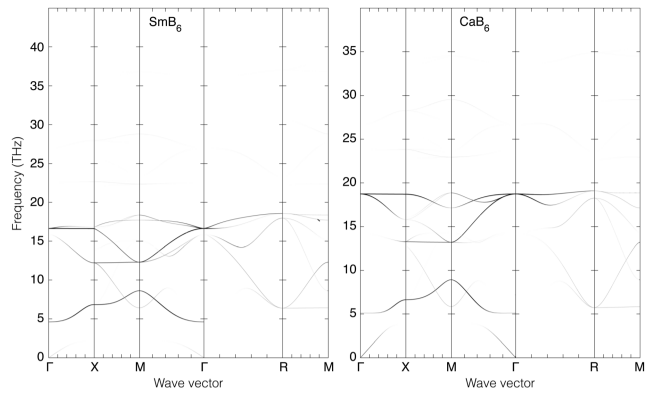


Fig. 5 Calculated phonon dispersion curves of SmB<sub>6</sub> (left) and CaB<sub>6</sub> (right) coloured between white and black to indicate the degree to which the mode eigenvectors can be described as a combination of the RUMs identified in a simple flexibility model, where black indicates close alignment of the phonon eigenvectors with the RUMs. The wave vector labels have conventional meaning:  $\Gamma$  represents the wave vector (0, 0, 0), X represents  $(\frac{1}{2}, 0, 0)$ , M represents  $(\frac{1}{2}, \frac{1}{2}, 0)$  and R represents  $(\frac{1}{2}, \frac{1}{2}, \frac{1}{2})$ .

not quite have zero frequency so it was necessary that the value of  $\Omega$  is significantly larger than the RUM frequencies. From the orthonormality of the phonon eigenvectors we have  $m_i(\mathbf{k}) \leq 1$ for all modes. The case  $m_i(\mathbf{k}) \sim 1$  corresponds to the case where there is a close matching of the eigenvector of the phonon in the full calculation to that of a RUM in the model calculation; in this case one of the products of the eigenvectors is close to a value of 1 and the frequency  $\omega_i'(\mathbf{k}) \simeq 0$ .

The results of the flexibility for SmB<sub>6</sub> and CaB<sub>6</sub> are shown in Fig. 5, where we plot the dispersion curves of Fig. 4 and shade the curves towards black where there is a perfect match between the phonon and RUM eigenvectors – where, in eqn (2),  $m_i(\mathbf{k}) = 1$  – and white where there is no match  $(m_i(\mathbf{k}) = 0)$ .

What we have previously noted is that when the phonon modes that are nominally RUMs are of low frequency, and separated from other modes, we see a clear RUM character. 18 However, when they are of higher frequency, or when there are other low-energy modes, the RUM eigenvectors can mix with those of other phonons. This has been discussed previously, 19 and is seen in the case of Zn(CN)2 where the RUM eigenvectors are spread over a few modes with frequencies around that of the nominally-RUM phonon,<sup>3</sup> and in ZrW<sub>2</sub>O<sub>8</sub> where there is a surface of RUMs in reciprocal space and such a broad distribution of phonon frequencies that the RUM is simply not visible at all.8

In Fig. 5 we see that the RUMs are identified with phonons of relatively high frequency, as in Zn(CN)2.3 In the latter case,

this arises from the resistance of the chemical bonds against flexing of the linear Zn-C-N-Zn connection. This is in contrast to the analogous material Si(NCN)2, which we have recently studied using an approach similar to the one here. 41 In that case, the Si-N-C linkage is much more flexible, and the RUMs are well separated from the frequencies of all other phonons and are clearly identified as such.

The high frequencies of the RUMs identified in Fig. 5 - see also the discussion of mode eigenvectors for phonons at zero wave vector in Section 3, where we identified the  $T_{1g}$  phonon as having RUM character - show that the torstional flexing of the B<sub>6</sub>-B<sub>6</sub> linkage is of high energy, and thus the picture of freelyjointed B<sub>6</sub> octahedra is not represented in the phonon spectrum. This is unlike the case of the cubic perovskites, where there is clear evidence from calculated44 and measured45 phonon dispersion curves of a high degree of flexibility of the linked octahedra. In the case of perovskite the RUMs have wave vectors only along the line M–R in reciprocal space,<sup>17</sup> that is, for wave vectors between  $(0, \frac{1}{2}, \frac{1}{2})$  and  $(\frac{1}{2}, \frac{1}{2}, \frac{1}{2})$ , and evidence from inelastic neutron scattering data and calculations show a low frequency along this branch. It is clear in the case of the metal hexaborides, Fig. 4 and 5, that the corresponding line in the dispersion curves does not show low frequencies.

We conclude from this analysis that in fact the network of connected B<sub>6</sub> octahedra does not have the flexibility seen in more traditional RUM systems such as cubic perovskites<sup>17</sup> and

phases of silica or silicates. 18 In fact this might have been anticipated from the data shown in Table 1 and Fig. 3. Incorporation of the larger cations causes the B<sub>6</sub> octahedra and their connecting B-B bonds to become strained, and hence the whole network will become more taut and less flexible.

## 5 Thermodynamic properties from vibrational spectra

The phonon densities of states of SmB<sub>6</sub> and CaB<sub>6</sub> are shown in Fig. 6a and b respectively, and corresponding data for SrB<sub>6</sub> and BaB<sub>6</sub> are given in the ESI.† The phonon densities of states show the gaps at frequencies of around 20 THz and just above 30 THz as seen in the phonon dispersion curves, Fig. 4. The same gaps are seen in SrB<sub>6</sub> and BaB<sub>6</sub>.

Of relevance for the thermal expansion are the mode Grüneisen parameters  $\gamma_i$  as defined earlier in Section 2.3. We have

computed these for a random set of wave vectors, and we plot a histogram of the distribution of values of  $\gamma$  as spread across the range of phonon frequencies. Fig. 6c and d show these distributions for SmB<sub>6</sub> and CaB<sub>6</sub> respectively, and corresponding data for SrB<sub>6</sub> and BaB<sub>6</sub> given in the ESI.† It is clear that there are phonons in SmB<sub>6</sub> and CaB<sub>6</sub> that have negative values of  $\gamma_i$ , but there are many more that have positive values, particularly at low frequency. However, in SrB6 and BaB6 there are many fewer phonons with negative values of  $\gamma$ .

Fig. 7 shows the distribution of significant values of  $\gamma$  across the dispersion curves of SmB<sub>6</sub> and CaB<sub>6</sub> shown in Fig. 4; corresponding diagrams for SrB6 and BaB6 are again given in the ESI.†

The overall Grüneisen parameters, γ, of SmB<sub>6</sub>, CaB<sub>6</sub>, SrB<sub>6</sub> and BaB<sub>6</sub>, as calculated from eqn (1), are shown as functions of temperature in Fig. 8. Not surprisingly, in the light of the data shown in Fig. 6, and of the corresponding data shown in the ESI,† even at low temperature the overall Grüneisen parameters

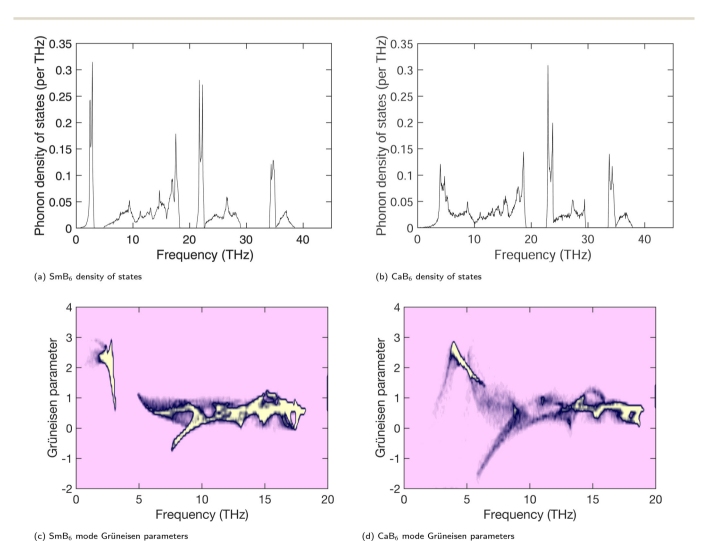


Fig. 6 Calculation of phonon densities of states for SmB<sub>6</sub> (a) and CaB<sub>6</sub> (b), and distribution of values of mode Grüeisen parameters across the range of frequencies for SmB<sub>6</sub> (c) and CaB<sub>6</sub> (d). In the latter two, pink represents a zero of phonon modes with a particular pair of values of frequency and Grüeisen parameters, yellow corresponds to the maximum in the distribution function, and the colour scheme passes from pink to yellow through dark blue. In order to highlight the existence of the distribution function for which there are fewer modes we saturate the scale of the histogram.

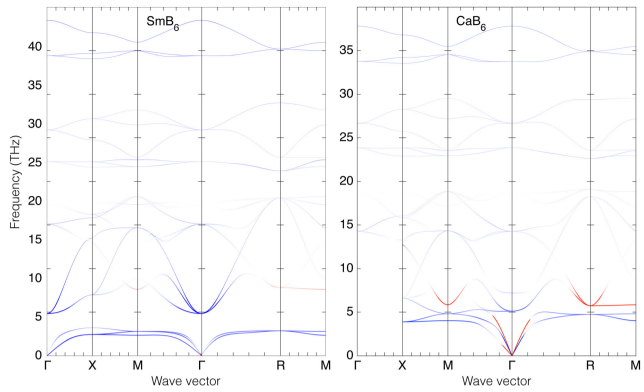


Fig. 7 Calculated phonon dispersion curves of SmB<sub>6</sub> (left) and CaB<sub>6</sub> (right) coloured between red and blue to indicate the values of the mode Grüneisen parameters  $\gamma_b$  where the strongest red indicates values of  $\gamma_i < -1$  and the strongest blue indicates values of  $\gamma_i > +3$ . The wave vector labels have conventional meaning:  $\Gamma$  represents the wave vector (0, 0, 0), X represents  $(\frac{1}{2}, 0, 0)$ , M represents  $(\frac{1}{2}, \frac{1}{2}, 0)$  and R represents  $(\frac{1}{2}, \frac{1}{2}, \frac{1}{2})$ .

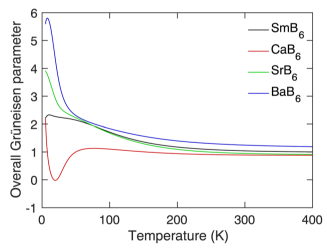


Fig. 8 Calculation of the temperature-dependence of the Grüneisen parameters of SmB<sub>6</sub>, CaB<sub>6</sub>, SrB<sub>6</sub> and BaB<sub>6</sub>.

are dominated by the distribution of positive-valued mode Grüneisen parameters at low frequencies.

What we see is that in SrB<sub>6</sub> and BaB<sub>6</sub> there is not the slightest hint of a negative thermal expansion. On the other hand, CaB<sub>6</sub> does show negative thermal expansion for a small range of temperatures at low temperature as indicated by a negative value of  $\bar{\gamma}$ . The interesting case for us is SmB<sub>6</sub>, which we have to conclude does not show negative thermal expansion as arising from the phonons. The accuracy of our calculated phonons is attested by the good agreement with the inelastic neutron scattering data shown in Fig. 4, and so we can conclude that the negative thermal expansion shown by the data in Fig. 2 must have a different origin that of the normal phonon mechanism.8

The values of the coefficients of volume thermal expansion for the four hexaborides, calculated as described in Section 2.3, are shown in Fig. 9. The data, of course, follow the trend seen in

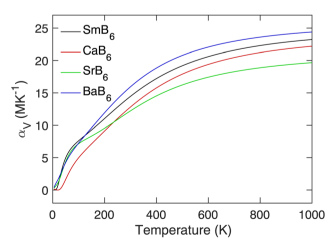


Fig. 9 Calculation of the temperature-dependence of the volume thermal expansion of SmB<sub>6</sub>, CaB<sub>6</sub>, SrB<sub>6</sub> and BaB<sub>6</sub>.

the overall Grüneisen parameters (Fig. 8), with a small range of NTE seen in  $CaB_6$  only.

### 6 Discussion

The work of this paper has been concerned with analysing the flexibility of the network of connected  $B_6$  octahedra in simple cubic metal hexaborides. The key finding from the DFT phonon calculations presented here is that in each case the network is not particularly flexible, with an energy cost to flex the orientation of one  $B_6$  octahedron about the linkage B–B bond connecting two tetrahedra. This was seen in the calculation of RUMs, where their frequencies are far from being low. As a result, we concluded that the NTE observed in Sm $B_6$  (Fig. 2) does not have its origin in the tension effect and phonons. On the other hand, there is a weak effect in Ca $B_6$  which would be interesting to investigate using low-temperature diffraction methods.

The methods employed in this study are based on the quasiharmonic model of Grüneisen in which the important anharmonic effects are coupling of interatomic force constants with volume. It might be asked whether NTE could arise if account were taken of direct phonon–phonon anharmonic interactions as accounted for, for example, in renormalised phonon theory. Most evidence, and in this we include our own theoretical study that addresses this point exactly, <sup>46</sup> is that anharmonic effects will lower the magnitude of the mode Grüneisen parameters of the renormalised phonons because their frequency will increase, and cannot change the sign of mode Grüneisen parameters from positive to negative. We do not believe that there is evidence to suggest that the current state of anharmonic phonon theory will indicate that there is a possibility to find NTE when Grüneisen theory doesn't show NTE.

It is known that there are electronic mechanisms for NTE,  $^{47}$  whether directly so or via magnetic interactions. In this regard, we note that although the samarium atom is magnetic, magnetic ordering in SmB<sub>6</sub> has been shown to be absent down to a temperature of 19 mK.  $^{21}$  Thus in our calculations we did not include magnetic interactions, and the good agreement of our calculated phonon dispersion curves with experimental data shows that this was reasonable. It is also observed that samples of the alkali-earth hexaborides can show magnetism, but this is believed to have its origin in a mechanism that is not directly a bulk property, such as broken B–B bonds at surfaces and interfaces, and magnetism of B<sub>6</sub> octahedra without saturated connectivity.  $^{48}$ 

In our calculations on SmB<sub>6</sub> we attempted to investigate effects of electron entropy through calculation of changes of electron entropy with volume. However, in our calculations on SmB<sub>6</sub> we observed that electron entropy consistently increases with volume across a range of Fermi-Dirac smearing temperatures from 100–1000 K, which means the electron entropy favours positive thermal expansion just as the phonon entropy does. Recently one of us (LL) had calculated the band structure for a range of volumes,<sup>49</sup> showing that over a contraction of

several percent there is no clear change in the electronic density of states.

At this point we are forced to conclude that the NTE in  $SmB_6$  remains of unknown origin.

## Author contributions

Li Li: investigation, formal analysis, validation, writing (review & editing), visualisation. Keith Refson: software, writing (review & editing). Martin T Dove: conceptualisation, methodology, software, validation, formal analysis, resources, writing (original draft), visualisation, supervision, funding acquisition.

#### Conflicts of interest

There are no conflicts to declare.

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