Phonon anharmonicity and negative thermal expansion in SnSe

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Abstract

The anharmonic phonon properties of SnSe in the Pnma phase were investigated with a combination of experiments and first-principles simulations. Using inelastic neutron scattering (INS) and nuclear resonant inelastic X-ray scattering (NRIXS), we have measured the phonon dispersions and density of states (DOS) and their temperature dependence, which revealed a strong, inhomogeneous shift and broadening of the spectrum on warming. First-principles simulations were performed to rationalize these measurements, and to explain the previously reported anisotropic thermal expansion, in particular the negative thermal expansion within the Sn-Se bilayers. Including the anisotropic strain dependence of the phonon free energy, in addition to the electronic ground state energy, is essential to reproduce the negative thermal expansion. From the phonon DOS obtained with INS and additional calorimetry measurements, we quantify the harmonic, dilational, and anharmonic components of the phonon entropy, heat capacity, and free energy. The origin of the anharmonic phonon thermodynamics is linked to the electronic structure.

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I. INTRODUCTION

Thermoelectric materials are of current interest for cost-effective, reliable power generation applications, either by utilizing natural sources or recovering man-made waste heat [1, 2]. In order to increase their heat-to-electricity conversion efficiency, thermoelectric materials need to have a low thermal conductivity, while maintaining a high electrical conductivity. A detailed understanding of phonons is necessary to rationalize both the thermodynamics and the thermal transport properties in thermoelectrics. Phonons are the main contributors to the entropy and heat capacity [3–6]. In addition, phonons are the dominant heat carriers in semiconductors, and understanding deviations from harmonic lattice dynamics is necessary to account for the bulk lattice thermal conductivity, \( \kappa_{\text{lat}} \). From a fundamental standpoint, phonons are sensitive to the nature of chemical bonding [7–10] and the temperature dependence of phonon spectral functions often provides important insights into the anharmonicity of the interatomic potential, directly affecting thermal resistivity through phonon-phonon scattering [10, 12]. The coupling between phonons and electronic structure also has an important effect on phonon frequencies (and linewidths) [5, 13–16], and electronic instabilities can produce large anharmonicity, helping to suppress \( \kappa_{\text{lat}} \) and improve thermoelectric efficiency [9, 10, 17].

Tin-selenide was recently reported to achieve an outstanding thermoelectric figure-of-merit, with Na-doped samples maintaining a high power factor over a broad range of temperatures [18–21]. At high temperature, SnSe undergoes a continuous structural phase transition between the low-symmetry Pnma (below \( T_c \approx 805 \text{K} \)) and the higher-symmetry Cmcm phases (above \( T_c \)). This phase transition has been the subject of a number of early crystallographic and thermomechanical experimental investigations [22, 23]. More recently, the ultra-low lattice thermal conductivity of SnSe has attracted great interest for thermoelectric applications, and has been studied both experimentally and through \textit{ab-initio} simulations [18–20, 22, 24]. The coupling between lattice distortion and electronic structure was recently shown to be responsible for the large anharmonicity, which persists in a broad range of temperatures, and leads to the low thermal conductivity [18, 17].

The crystal structure of SnSe is illustrated in Fig. 1. Importantly, x-ray and neutron diffraction studies...
atoms are in grey and Se atoms in green.

FIG. 1. Crystal structure of SnSe in Pnma (a) and Cmcm (c) phases, illustrating the double bilayer structure and Pnma distortion, and corresponding Sn-Se bonding state (b,d). Sn atoms are in grey and Se atoms in green. $d_1$ and $d_2$ are labels for bonds in out-of-plane and in-plane directions. The orientation of crystal axes in the Cmcm structure is chosen to match the Pnma phase, in order to facilitate the comparison. The rotation of pyramids (shaded in (a)) $\theta$ and the angle between two $d_2$ bonds $\phi$ in the Pnma structure are also shown in (a). The conventional unit cell edges are indicated by black square box.

showed that, in the Pnma phase, the $c$ axis, parallel to the direction of corrugation of Sn-Se bilayers below $T_c$, exhibits a negative thermal expansion (NTE) coefficient $|\Delta c/\Delta T|$. The thermodynamic and atomistic origins of this effect in SnSe remain to be established. The NTE implies that the thermodynamic generalized Grüneisen tensor must have negative values for the diagonal element corresponding to the in-plane layer [5]. Yet, this is in contrast to positive values predicted from quasiharmonic $ab$-initio simulations [10]. Further, our inelastic neutron scattering measurements showed pronounced differences in the behaviors of phonons propagating along in- and out-of-plane crystallographic axes [10]. Since the coupling between the lattice distortion and the electronic structure is a strong source of anharmonicity [17], it is important to understand its role in the unusual thermal expansion of SnSe.

Here, we investigate the origin of the NTE and anharmonic phonon thermodynamics in SnSe, focusing on the Pnma phase, using inelastic neutron scattering (INS) and nuclear resonant inelastic X-ray scattering (NRIXS) measurements, and first-principles simulations. Our measurements show a strong temperature dependence of the phonon frequencies, revealing a high degree of anharmonicity in the interatomic potential. We show how this anharmonicity is related to the structural phase transition. We model the thermal expansion and its anisotropic behavior, from first-principles, especially the lattice contraction parallel to the Sn-Se layers, and find good agreement with reported lattice parameter measurements. In addition, we quantify the harmonic, dilational, and anharmonic contributions of phonons to the entropy, internal energy and free energy.

II. SAMPLE PREPARATION

Single crystals of SnSe were synthesized from high purity Sn and Se (Alfa Aesar, 99.999%) in fused silica ampoules. After an initial reaction, a polycrystalline precursor was melted and subsequently crystallized while cooling at 0.4-0.5°C/h; a 24 hr annealing at ~830°C occurred before cooling to room temperature. One of the single crystals was ground into a fine powder for neutron scattering measurements of the phonon density of states. Some properties of these samples were previously reported, and further details can be found in Ref. [10].

III. SPECTROSCOPY

A. Inelastic Neutron Scattering

We measured the phonon DOS of polycrystalline SnSe (mass ~ 8 g) using the time-of-flight wide angular-range chopper spectrometer (ARCS) at the Spallation Neutron Source (SNS) at Oak Ridge National Laboratory [38]. The powder sample was contained in a standard thin-walled aluminum can. Measurements at low (5 $\leq$ $T$ $\leq$ 300 K) and high temperature (300 $\leq$ $T$ $\leq$ 750 K) were performed using aluminum can inside a closed-cycle helium refrigerator and low-background resistive furnace. At low temperatures, we filled the sample chamber with low pressure of helium to facilitate cooling. We used two incident energies, $E_i$ = 30 meV and 55 meV, when combined it provided an entire phonon spectrum and high resolution datasets to distinguish between closely spaced phonon peaks. An oscillating radial collimator was employed to minimize scattering from the sample environment. We followed the similar steps in data normalization, reduction to $Q$-$E$ (momentum and energy transfer) space, background subtraction, multiphonon scattering, and removal of elastic peak as described in our previous work [16].

The neutron scattering cross-sections are different for different elements. The phonon DOS derived from INS measurements is weighted by the respective scattering cross-sections ($\sigma$) and mass (m). The neutron-weighting factors for Sn, and Se are $\sigma_{Sn}/m_{Sn} = 0.0412$, and $\sigma_{Se}/m_{Se} = 0.1051$, respectively (in units of barns/amu). Due to the larger neutron-weighting factor for Se in comparison to Sn, phonon DOS is over-weighted by Se vibrational contributions. Accordingly, the neutron-weighted
The energy range of 160 meV with 0.25 meV steps. Reactions at 20-130 nsec. The inelastic spectra were collected over a signal, delayed in time relative to the prompt pulse by 0.25 meV. The inelastic signal collected was reduced to 1.2 meV using a high resolution cryostat. Since SnSe exhibit anisotropic behavior, we could extract the direction-projected phonon DOS by changing the orientation of the sample with respect to the incident x-ray beam. For the a-axis projected phonon DOS, the sample was mounted such that the incident x-ray beam was parallel to a-axis, while for the projected phonon DOS perpendicular to a-axis, the incident beam was parallel to the b-c plane. The incident x-ray energy was 23.88 keV, the nuclear resonance energy of 119Sn. The energy bandpass of incident beam was reduced to 1.2 meV using a high resolution crystal monochromator. The inelastic signal collected is comprised mainly of 23.88 keV nuclear fluorescence signal, delayed in time relative to the prompt pulse by 20-130 nsec. The inelastic spectra were collected over an energy range of 160 meV with 0.25 meV steps. Repeat scans were taken and data were subsequently added. The 119Sn specific phonon DOS was extracted from the NRIXS data using the PHOENIX package. The samples were single-crystal SnSe platelets (no isotopic enrichment), cut from INS crystals. The platelet samples were mounted with either transmission geometry (beam along a-axis) or grazing geometry (beam nearly parallel to b – c plane). Because of the crystalline anisotropy, the direction-projected partial DOS for each configuration provides information about different Sn motions (∥a or ⊥ a, respectively).

IV. MODELING

A. Structural Relaxation and Phonon Dispersions

First-principles simulations were performed in the framework of density functional theory (DFT) as implemented in the Vienna Ab initio Simulation Package (VASP 5.3). A 6 × 12 × 12 Monkhorst-Pack electronic k-point mesh was used for all of our simulations, with a plane-wave cut-off energy of 500 eV. The projector-augmented-wave potentials explicitly included four valence electrons for Sn (5s25p2), and 6 for Se (4s24p4). The lattice parameters and atomic positions were optimized until forces on all atoms were smaller than 1 meV Å⁻¹. We carefully evaluated the accuracy of our phonon calculations, comparing the local-density approximation (LDA) exchange-correlation (XC) functional [49] and the generalized gradient approximation (GGA) with XC functional given by the Perdew-Burke-Ernzerhof (PBE) parametrization [50]. The relaxed unit cell parameters are compared with the experimentally reported structure (Ref. 29) in Table I. The agreement between the LDA and PBE structures and the experimental structure is good, with, as expected, LDA underestimating the lattice constants and PBE overestimating them, by opposite amounts.

The phonon dispersions were calculated in the harmonic approximation, using the finite displacement approach as implemented in Phonopy [51], with the atomic forces in the distorted supercells obtained with VASP. To construct the force-constant tensor using the finite displacement approach in Phonopy [51], we computed 8 independent atomic displacements, each with atomic displacement amplitudes of one hundredth of lattice constants. The LDA phonon calculations used a 2 × 4 × 4 supercell of the primitive cell, containing 256 atoms, while GGA calculations used up to 3 × 5 × 5 supercells containing 600 atoms. We compare the phonon dispersion and phonon DOS of SnSe at 300 K with experimental phonon dispersions measured with INS in Fig. 2 and with the NRIXS and INS phonon DOS in Fig. 11 and 6, respectively. The experimental phonon dispersions in Fig. 2 were extracted from INS datasets originally reported by our group in Ref. 10. From these comparisons with INS measurements, one can see how the PBE simulations systematically underestimate the phonon frequencies in this material, while LDA simulations are much more accurate. Even with the larger supercell, the GGA phonon frequencies show a worse agreement with INS data than the LDA calculations. The mean phonon energy calculated from experimental phonon DOS at 300 K (corrected for neutron weighting), GGA simulations, and LDA simulations are 12.92, 12.15, and 12.90 meV, respectively. Additionally, the GGA phonon group velocities vq,j are on average ∼10% lower than the LDA values near the Γ point. Thus, first-principles calculations of the lattice thermal conductivity (κlat) using GGA (Refs. 32, 33) are expected to carry a sizeable systematic error, since κlat is proportional to vq,j, with a further dependence of phonon scattering rates on the dispersions, while LDA calculations are expected be more accurate [30].

B. Negative Thermal Expansion

Non-cubic materials often exhibit anisotropic thermal expansion coefficients, reflecting the anisotropy in crystal structure and bonding. In the case of orthorhombic SnSe, the anisotropy is particularly striking, with negative thermal expansion (NTE) of the c axis, parallel to the direction of corrugation of Sn-Se layers (also corresponding to the direction of atomic motions across the Sn-Se bonds).
phase transition), while the other in-plane axis (b) and the out-of-plane a-axis show normal, positive expansion [22,24]. The relative change of lattice parameters with temperature reported by Adouby et al. in Ref. [23] are shown in Fig. 3. This distortion can also be analyzed as a rotation of SnSe$_3$ tetrahedra, as illustrated in Fig. 1 and further discussed below. We proceed to rationalize this behavior based on first-principles thermodynamics and a microscopic analysis of bonding.

We start by noting that simple Grüneisen parameter calculations are insufficient to explain the anisotropy and NTE along one or more crystallographic axes. Instead, one needs to compute the 3-D total free energy, whose temperature dependence is dominated by the phonon entropy. In order to clearly make this point, we start by performing the computation of Grüneisen parameters. A generalized Grüneisen tensor $\gamma_{ij}^{\text{avg}}$ for a given polarization index $p$ and wave vector $q$ is given by:

$$\gamma_{ij}^{\text{avg}}(q) = -\left\{ \frac{1}{\omega_p(q)} \frac{\partial \omega_p(q)}{\partial \eta_{ij}} \right\}_T \bigg|_{\eta=0},$$

where, $\eta_{ij}$ is the $(i,j)$ component of applied isothermal strain, and $\omega_p(q)$ is the atomic vibration frequency. Summation over polarization index and wave vector, weighted by heat capacity at constant configuration $C_\eta$, yields an average value of a generalized Grüneisen tensor. We have calculated the generalized Grüneisen tensor from our DFT simulations of phonons $\gamma_{ij,DFT}^{\text{avg}}$. In the case of SnSe, considering the symmetry of the second order tensor for the orthorhombic unit cell, non-diagonal components are exactly zero. Thus, the only isothermal strains required are $\eta_{11}$, $\eta_{22}$, and $\eta_{33}$, corresponding to stretching or compressing the lattice along crystallographic axes $a$, $b$, and $c$, respectively. Subsequently, the generalized Grüneisen tensor, $\gamma_{ij,DFT}^{\text{avg}}$, is obtained by averaging over all polarizations and wave vectors, with each mode weighted by its heat capacity at constant configuration as follows:

$$\gamma_{ij,DFT}^{\text{avg}} = \begin{bmatrix} 1.44 & 0.0 & 0 \\ 0 & 1.35 & 0 \\ 0 & 0 & 0.64 \end{bmatrix}.$$

We have also calculated the Grüneisen tensor along high symmetry directions by averaging over polarizations only. Results are listed in Table I. As can be seen from Table I, isothermal strains along $a$, $b$, and $c$ axes lead to positive Grüneisen parameters i.e., softening of phonon branches along all directions, except for isothermal strain along the $c$ axis, which produces a stiffening of phonon modes along $\Gamma - X$ and $\Gamma - T$. However, summation over the entire Brillouin zone at constant configuration yields positive effective Grüneisen parameters for all cases of isothermal strain as noted in Eq. (3) and Table II which is at odds with the experimentally observed negative thermal expansion along the $c$ axis.

Alternatively, the relationship between the thermodynamic generalized Grüneisen tensor and the thermal expansion tensor, $\alpha_{kl}$, can be expressed in terms of the isothermal bulk modulus $B_T$, volume $V$, and heat capacity at constant configuration $C_\eta$, as following:

$$\gamma_{ij,DFT}^{\text{avg}} = \frac{V}{C_\eta} \sum_{kl} B_T^{T DFT} B_{ijkl} \alpha_{kl}. \quad (4)$$

The isothermal bulk modulus tensor, $B_{ijkl}^{T}$, can be determined either from the phonon group velocities along different crystallographic directions, or from DFT simulations by applying isothermal longitudinal and transverse strains. We have calculated $B_{ijkl}^{T}$ with both approaches. The independent components of $B_{ijkl}^{T}$, constrained by the orthorhombic symmetry, are shown in Table III. The phonon group velocities at 300 K were obtained from INS on single crystals, previously reported by our group in Ref. [10]. The tensor components not
Thus, $\gamma_{ij,Exp}$ are substantial negative, while we do observe a positive prediction. Importantly, while we do observe a negative $\gamma$ along $c$ as a whole, the DFT prediction is positive. Importantly, while we do observe a substantial negative $\gamma$ along $\Gamma - X$ by applying the strain along $c$ (cf Table III), this alone is insufficient to produce negative $\gamma$ along $c$ as a whole, once the integration is performed over the entire Brillouin zone. We note that this is not a failure of the DFT calculation per se, but rather arises because of the omission of couplings between different crystallographic axes.

The difficulty to capture the NTE along $c$ from the Gruneisen parameter prompted us to instead compute the three-dimensional free energy surface, on a 3D grid of isothermal strains. To construct the total 3-D free energy surface, we created a $3 \times 3 \times 3$ grid of lattice parameters with 1% expansion and compression of lattice parameter along each crystallographic axes with total of 27 grid points. The inherent advantage of this approach is that it captures the evolution of both electronic and vibrational free energy by varying lattice parameters individually (i.e., $\{a \pm \Delta a, b, c\}$, $\{a, b \pm \Delta b, c\}$, and so on), as well as coupled terms (i.e., $\{a \pm \Delta a, b \pm \Delta b, c \pm \Delta c\}$, and so on), and determine the total free energy minima. The total free energy is a sum of electronic free energy, $F_{el}$, and vibrational free energy, $F_{vib}$, which can be expressed as following:

$$ F_{el} = E_{gs} - TS_{el}, $$

and

$$ F_{vib} = \frac{1}{2} \sum_i^{3N} \varepsilon_i + k_B T \sum_i^{3N} \ln \left\{ 1 - \exp\left( - \frac{\varepsilon_i}{k_B T} \right) \right\}, $$

where electronic entropy, $S_{el}$, and the Fermi-Dirac occupation function, $f$, are given by:

$$ S_{el} = -k_B \int_{-\infty}^{\infty} [(1 - f) \ln(1 - f) + f \ln f], $$

and

$$ f = \frac{1}{\exp\{(E - E_F)/k_B T\} + 1}, $$

respectively. Here, $E_{gs}$ is ground state energy, $\varepsilon_i$ is phonon energy of $i^{th}$ mode, $E_F$ is the energy at Fermi level, $k_B$ is Boltzmann’s constant, and $N$ is the number of atoms in the unit cell. We obtain the ground-state electronic energy, and the electronic density of states (eDOS) from DFT simulations for all points on the $(a, b, c)$ grid. The electronic entropy was determined from eq. (5), and (7). The vibrational free energy was obtained from eq. (7), using the phonon density of states (phonon DOS) calculated as described in section IV A.

The measured thermal expansion between 300K and 813K along $a$, $b$, and $c$ is approximately 1.8%, 3.5%, and -3.1%, respectively [23]. Additional points were added to extend our grid to larger expansions/compressions of 3% along $a$, 5% along $b$, and 5% along $c$. However, phonons become unstable for extended grid points. Thus, we limit our procedure to the original 27-point grid, extended with the seven new points for which phonons are stable, shown in Fig. 3(b). To capture the NTE along $c$ over a wider

| $\gamma_{ij,Exp}$ | $\gamma_{ij,DFT}$ |
|-------------------|-------------------|
| $\gamma_{ij,Exp}$ | $\gamma_{ij,DFT}$ | $\gamma_{ij,DFT}$ |
| $\gamma_{ij,Exp}$ | $\gamma_{ij,DFT}$ | $\gamma_{ij,DFT}$ |
| $\gamma_{ij,Exp}$ | $\gamma_{ij,DFT}$ | $\gamma_{ij,DFT}$ |

Thus, $\gamma_{ij,Exp}$ and $\gamma_{ij,DFT}$ are quite different. The Gruneisen parameter estimated from experimental data is negative along the $c$ axis, but the DFT prediction is positive. Importantly, while we do observe a substantial negative $\gamma$ along $\Gamma - X$ by applying the strain along $c$ (cf Table III), this alone is insufficient to produce negative $\gamma$ along $c$ as a whole, once the integration is performed over the entire Brillouin zone. We note that this is not a failure of the DFT calculation per se, but rather arises because of the omission of couplings between different crystallographic axes.

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| $\gamma_{ij,Exp}$ | $\gamma_{ij,DFT}$ | $\gamma_{ij,DFT}$ |
| $\gamma_{ij,Exp}$ | $\gamma_{ij,DFT}$ | $\gamma_{ij,DFT}$ |
| $\gamma_{ij,Exp}$ | $\gamma_{ij,DFT}$ | $\gamma_{ij,DFT}$ |
range of temperature, we performed a quadratic extrapolation of the free energy surface (see Eq. (10)). We note that the computational cost scales cubicly with the linear grid size, hence the practicality of finer grids is limited.

We determine the equilibrium lattice parameters, $a_{eq}$, $b_{eq}$, and $c_{eq}$ by minimizing a second-order polynomial fit to the free-energy, $F(a, b, c, T)$, with respect to lattice parameters at $T = T_0$.

\[
F(a, b, c, T_0) = C_0 + C_1a + C_2b + C_3c + C_4ab
+ C_5bc + C_6ca + C_7a^2 + C_8b^2 + C_9c^2
\]

\[
\frac{\partial F(a, b, c, T_0)}{\partial a} = 0
\]

\[
\frac{\partial F(a, b, c, T_0)}{\partial b} = 0
\]

\[
\frac{\partial F(a, b, c, T_0)}{\partial c} = 0
\]

The results are shown in Fig. 3(a). As can be seen on this figure, the experimental temperature dependence of lattice parameters is qualitatively captured, including the NTE along $c$. This indicates an interplay between electronic and vibrational components of the free energy, which ultimately controls the minima of $F$. The deviation between calculated and experimental lattice parameters at high $T$ is partially due to the fact that the theoretical lattice parameters at $T = 0$, $a = 11.309$, $b = 4.119$, $c = 4.300\, \text{Å}$, are smaller than the experimental values: $a = 11.502$, $b = 4.153$, $c = 4.450\, \text{Å}$, leading to shorter theoretical bond lengths, especially in the $b - c$ plane.

In addition, strong anharmonic effects, not included in our quasiharmonic calculations of phonon spectra for strained cells, are seen to occur in the last $\sim 200\, \text{K}$ below $T_c$ in our heat capacity measurements (see below). One could also include anharmonic effects in the phonon calculations, for example with ab initio molecular dynamics or other methods incorporating anharmonic phonon renormalization occurring because of the thermal bath [10, 11, 32]. However, our goal is not to exactly match the measured thermal expansion coefficient, but rather to rationalize the origin of the anisotropy and NTE along $c$, which can be explained with this simple and insightful approach.

The electronic ground state energy at $T = 0\, \text{K}$ is minimum for the fully relaxed configuration (relaxed lattice parameters and atomic positions), $(a_0, b_0, c_0)$, and increases with any strain. We note that, for the strained configurations, the minimum in the electronic ground state energy is related to the lattice parameters such that there are two possible scenarios: $(\Delta a, \Delta b > 0, \Delta c < 0)$ or $(\Delta a, \Delta b < 0, \Delta c > 0)$. Hence, the electronic ground state energy alone can not uniquely define the equilibrium lattice structure. Thus, it is the vibrational free energy that brings the minimum of $F$ towards $(\Delta a, \Delta b > 0, \Delta c < 0)$. The calculated phonon DOS for three different configurations is shown in Fig. 4. From this figure, we observe that $(\Delta a, \Delta b > 0, \Delta c < 0)$ leads to additional spectral weight at low energy in the phonon DOS ("softening"), while $(\Delta a, \Delta b < 0, \Delta c > 0)$ results in a stiffening. The softening in the first case increases the phonon entropy, and thus lowers $F$. As further discussed below, the configuration $(\Delta a, \Delta b > 0, \Delta c < 0)$ leads to considerable softening of high energy phonons.

### C. Structural Evolution and NTE

We now investigate the structural changes inside the unit cell in connection with the NTE. In particular, we show that the NTE of SnSe is closely related to the electronic instability and its coupling to the anharmonic vi-
FIG. 4. Phonon density of states calculated from DFT for three different configurations of lattice parameters. The ground-state equilibrium lattice parameters are \(a_0 b_0 c_0\). The other two configurations are \(\Delta a/a = \Delta b/b = +1\%\), \(\Delta c/c = -1\%\) (\(a↑ b↑ c↓\)) and \(\Delta a/a = \Delta b/b = -1\%, \Delta c/c = +1\%\) (\(a↓ b↓ c↑\)).

TABLE IV. Rotation of pyramidal induces NTE along \(c\). Bond length and lattice constant in Å. All data is from Ref. [23] except for the calculated value \(b\) and \(c\). The small discrepancy between calculated \(c\) and \(c_{\text{exp}}\) results from the ignorance of tilt of two \(d_2\) bonds plane.

|       | \(d_2\) | \(\theta\) | \(d_2\) | \(\phi\) | \(c\) | \(b\) | \(c_{\text{exp}}\) | \(b_{\text{exp}}\) |
|-------|---------|------------|---------|--------|------|-----|---------------|---------------|
| Cmcm  | 2.75    | 0          | 3.05    | 89.85  | 4.32 | 4.31| 4.31          | 4.31          |
| Pnma  | 2.72    | 7.71       | 2.80    | 95.67  | 4.49 | 4.15| 4.45          | 4.15          |

Within a close approximation, we have:

\[ c \simeq 2d_3 \sin \theta + 2d_2 \cos \phi/2 \]  \(\text{(11)}\)

\[ b = 2d_2 \sin \phi/2 \]  \(\text{(12)}\)

From Table IV we can see that \(2d_2 \cos(\phi/2)\) decreases from Cmcm to Pnma structure, and therefore, the expansion along \(c\) can be seen to result from the rotation of pyramids (\(\theta\)), which couples strongly to the \(Y + A_g\) mode (\(d_1\) almost remains constant). From the plots in Fig. 5, we can confirm that the NTE along \(c\) is indeed induced by the rotation of pyramids, which is strongly coupled to the anharmonic modes (soft-modes in Cmcm phase generating zone-center soft-modes in Pnma phase) \([17]\). The strongly anharmonic character of these modes partly explains the shortcomings of the quasiharmonic free-energy calculations near the phase transition, seen in Fig. 3.
V. ANHARMONIC PHONON THERMODYNAMICS

A. Phonon DOS and Phonon Entropy

Much of the free-energy discussion above was limited to a quasiharmonic approximation. In order to estimate the anharmonic component of the vibrational entropy, we investigate the temperature dependence of the measured phonon DOS. Fig. 5a and 5b show the experimental phonon DOS measured at different temperatures for $E_i = 55$ meV and $30$ meV, respectively. The low and high energy phonon peaks soften by $\sim 0.3$ and $\sim 1.0$ meV, respectively as the sample temperature is increased from 10 to 750 K. One can observe a drastic broadening of the spectrum with increasing $T$, in particular for optical modes above 13 meV. While the lowest energy zone-center TO mode has been shown to soften strongly with increasing $T$ in Pnma in our previous INS dispersion measurements (Ref. 10), that mode is confined to the zone-center and has a limited spectral weight in the DOS. We note that the neutron-weighted phonon DOS from harmonic DFT simulations agrees well with the total and direction-projected phonon DOS measured at 300 K (Fig. 11 and 9). The direction-projected experimental phonon DOS measurements (Fig. 11) provide substantial information in separating the two low energy Sn dominated phonon peaks at $\sim 6$ and $\sim 11$ meV. The low energy ($\sim 6$ meV) phonon peak include both out-of-plane (along $a$-axis) and in-plane (b-c plane) Sn vibrations, while $\sim 11$ meV phonon peak primarily originates from in-plane (b-c plane) Sn vibrations. Both Sn dominated phonon peaks show substantial broadening in the phonon spectrum with increasing temperature (Fig. 9).

With increasing temperature, it is the coupling between lattice distortion (especially the large Sn in-plane displacements) and the electronic structure that generates strong anharmonicity (leading to phonon-phonon interactions) 10, primarily responsible for a significant phonon softening of high energy phonons and phonon broadening of entire phonon spectra seen in Fig. 8. This conclusion is further supported by our DFT simulations of NTE modeling where we identify that the minimization of electronic free energy alone is not sufficient to obtain the experimental lattice parameters with temperature, and including the change in vibrational free energy due to lattice distortions is necessary to reach equilibrium. As we have detailed in Ref. 17 the origin of the anharmonicity for Sn in-plane motions is the Jahn-Teller instability involving the Se $p$-states.

To quantify the anharmonic contribution to $S_{vib}$, we estimated a phonon broadening parameter, $\Gamma_Q^p$, by convolving the generalized (neutron-weighted) phonon DOS from DFT, $g(E)$, with a damped harmonic oscillator function, $B(E, E', T)$, 554

$$g(E, T) = \int B(E, E', T)g(E' - \Delta E'(T)) \, dE'$$

and matching the result with the measured phonon DOS. Here,

$$B(E, E', T) = \frac{2\Gamma_Q^T}{(\pi E')^2}\left[\frac{E'}{2\Gamma_Q^T} + \frac{1}{2}\left(1 + \frac{E'}{2\Gamma_Q^T}\right)^2\right]$$

and $\Delta E'$ is a shift in phonon energy calculated by fitting the Lorentzian to the peaks in the measured phonon DOS. The resulting phonon DOS, $g(E, T)$, with temperature-dependent phonon broadening parameter, $\Gamma_Q^T$, is shown in Fig. 7. This model reproduces the temperature dependence of the experimental phonon DOS well, and enables us to assess the anharmonic $S_{vib}$. For this purpose, we use the non-neutron-weighted version of $g(E, T)$.

The vibrational entropy $S_{vib}$ can be expressed as

$$S_{vib} = 3Nk_B$$

$$\times \int g(E, T) \left[(1 + n_T)\ln(1 + n_T) - n_T\ln(n_T)\right] \, dE$$

where $n_T(E) = [e^{E/k_BT} - 1]^{-1}$ is the Bose-Einstein occupation factor for phonons, and $N$ is the number of atoms in the crystal.

The harmonic phonon entropy, $S_{vib,h}$, was calculated by convolving the harmonic phonon DOS from DFT with a damped anharmonic oscillator function $B(E, E', T)$ at $T = 10$ K. The resulting phonon DOS was substituted in Eq. (15) to evaluate $S_{vib,h}$ at different temperatures. To evaluate the total entropy, $S_{vib}$, at different temperatures, the $T$-dependent phonon oscillator function $B(E, E', T)$ was convolved with the DFT phonon DOS, and substituted in Eq. (15). The vibrational entropy due to thermal expansion of the lattice is given by

$$S_{vib,d} = \int_0^T \left(\frac{C_p - C_v}{T'}\right) \, dT'$$

$$= \int_0^T \frac{\alpha_{vib}(T')^2V(T')}{\chi(T')} \, dT'$$

where $\alpha_{vib}$ is the volumetric thermal expansion coefficient, $\chi$ is the compressibility (inverse of bulk modulus), and $C_p$ and $C_v$ are specific heat capacity of material at constant pressure and temperature.

Using the total $S_{vib}$, harmonic $S_{vib,h}$, and dilational $S_{vib,d}$ components of the vibrational entropy, the nonharmonic and anharmonic entropy were calculated as: $S_{vib,nh} = S_{vib} - S_{vib,h}$, and $S_{vib,ah} = S_{vib} - S_{vib,h} - S_{vib,d}$, respectively. For the calculation of dilational entropy, we used the thermal expansion coefficient from Wiedemen et al. 24, and temperature dependent compressibility from He et al. 56. In addition, we have also estimated...
the total entropy from our experimental heat capacity measurements ($S_{DSC}$) using the following expression,

$$ S_{DSC} = \int_{T_0}^{T} \frac{C_p}{T} dT. \quad (17) $$

The calculated harmonic, dilational, non-harmonic, and total vibrational entropy values are shown in Fig. 8.

As we can observe from Fig. 8, the main entropic contribution is the harmonic term. However, as temperature increases, the nonharmonic contribution becomes significant and cannot be ignored. The nonharmonic entropy is a summation of dilational and anharmonic entropy contributions. We have also calculated nonharmonic contributions to the vibrational internal energy and vibrational free energy following methods recently reported in Ref. 57. We should note that since tin-selenide is a indirect band gap semiconductor (band gap of 0.86 eV in Pnma phase at room temperature) [18], we could expect the contribution of electronic excitations to specific heat capacity/thermal expansion coefficient to be negligible at room temperature. The band gap in SnSe is temperature dependent, and reduces to $\sim 0.4$ eV in Cmcm phase [15]. While the temperature dependence of band gap may affect our results slightly, we do not expect this effect to be significant. From these calculations, a similar trend is observed in vibrational internal energy (Fig. 9) and vibrational free energy (Fig. 10), with the anharmonic contribution rising as $T_c$ is approached. As we can observe, at high $T_c$, thermal expansion (dilation) alone cannot account for nonharmonic contributions, and anharmonicity plays a large role near the instability.

It is interesting to note that in the experimental phonon DOS (Fig. 6), there is considerably larger softening, $\Delta E/(E) \sim -5.2\%$, of high energy phonons ($E \geq 13$ meV) compared to the softening of low en-

FIG. 6. Neutron-weighted phonon DOS of SnSe measured with inelastic neutron scattering at different temperatures for incident neutron energies of a) 55 meV and b) 30 meV, compared with neutron weighted and experimental resolution convoluted DFT simulations.

FIG. 7. Generalized phonon DOS of SnSe calculated with DFT, weighted with experimental neutron cross-sections. The phonon DOS is convolved with a damped harmonic oscillator function, as detailed in the text.
energy phonons, $\sim -3.8\%$, between $T = 10$ and 750 K.

As evident from Fig. 11a, low energy and high energy modes are dominated by Sn and Se, respectively. We selectively investigate the change in low and high energy phonon energies by applying 1\% isothermal strain along $\Delta a/a = \Delta b/b = +1\%$, $\Delta c/c = -1\%$ (3-D free energy surface minima direction as described earlier), and find that high energy phonons soften by $\sim -2.1\%$ in comparison to $\sim -1.0\%$ for low energy phonons.

The values of Grüneisen parameter along high-symmetry directions are reported in Table V. The large values of Grüneisen parameter across the various high symmetry directions in SnSe reflect the pronounced anharmonicity, which is also shown quantitatively in the anharmonic vibrational entropy, internal energy and free energy calculations, while the significant variation in values across different directions is attributed to the structural anisotropy in this material.

### B. Partial Phonon DOS and Thermal Displacement Parameters

Fig. 11a shows the partial phonon DOS of tin and selenium calculated from DFT simulations for $a_0b_0c_0$, the ground state equilibrium lattice parameters for fully relaxed structure with Sn and Se atoms predominantly occupying the low and high energy phonon spectrum, respectively. To enable the direct comparison with experimentally measured NRIXS Sn partial phonon DOS par-
TABLE V. Grüneisen parameter ($\gamma$) of tin-selenide calculated by applying 1% isothermal strain along $\Delta a/a = \Delta b/b = +1\%$, $\Delta c/c = -1\%$ ($a \uparrow b \uparrow c \downarrow$). Average along particular direction such as $\Gamma - X$ represents the values of $\gamma$ averaged over $q$-point along that direction weighted by specific heat capacity. More details in text.

| $\gamma_{\text{avg}}$ | Isothermal strain along: ($a \uparrow b \uparrow c \downarrow$) |
|------------------------|--------------------------------------------------|
| All modes              | modes $< 13$ meV | modes $\geq 13$ meV |
| $\Gamma - X$           | 5.51          | 4.49          |
| $\Gamma - Y$           | 2.79          | 2.06          |
| $Y - P$                | 1.91          | 2.46          |
| $\Gamma - P$           | 2.46          | 2.86          |
| $\Gamma - A$           | 2.20          | 2.63          |
| A - Z                  | 0.63          | 1.68          |
| $\Gamma - Z$           | 2.21          | 2.61          |
| $\Gamma - T$           | 2.90          | 3.23          |

Parallel and perpendicular to $a$-axis, we have also calculated the projected phonon DOS of Sn as shown in Fig 11 b, and 11c. The good agreement between experiment and simulations further validates the accuracy of DFT-LDA. Additionally, from the partial phonon DOS and $f$-factor (Lamb-Mössbauer factor) measured from NRIXS [15, 54], the mean square thermal displacement of Sn can be calculated. The $f$-factors corresponding to parallel phonon DOS and perpendicular to $a$-axis are 0.1370 and 0.1487, respectively. We should note that, the exact sample orientation for in-plane ($\perp a$) was not known; however, the in-plane NRIXS measurement agrees well with the simulation for the $b$-axis and we believe this was the likely orientation.

Furthermore, we have calculated the mean square thermal displacement parameter, $\langle |U_\alpha(j)|^2 \rangle$, from DFT phonon spectra, as:

$$\langle |U_\alpha(j)|^2 \rangle = \frac{\hbar}{2Nm_j} \sum_{q,s} \omega_s(q)^{-1}(1+2n_s(q))|e_{s,\alpha}(j,q)|^2,$$

where, $\alpha$ denotes thermal displacement direction, $m_j$ is mass of atom at location $j$, $\omega_s(q)$ is phonon frequency of $s$th phonon branch at wave vector $q$, $n_s$ is the mean Bose-Einstein occupation factor, and $e_{s,\alpha}(j,q)$ is phonon wave vector. The results for $T = 300$ K, listed in Table VI, are consistent with values measured by Chattopadhyay et al. at room temperature with neutron diffraction [22]. The thermal displacement parameters for Sn and Se have similar magnitudes at 300 K, thus indicating that the average bond stiffness at each site is comparable. Indeed, our DFT simulations give a small increase (~12%) in the trace of Se on-site force-constant matrix in comparison to Sn. The ratio of average frequency for Se and Sn vibrations calculated from DFT is 1.38, close to the effect of mass ratio, $\sqrt{M(\text{Sn})/M(\text{Se})} \approx 1.23$. It is worth emphasizing, that while the magnitude of thermal displacements of Se atoms reported by Chattopadhyay et al. increases linearly with temperature, off-centered Sn atoms show a superlinear increase for in-plane thermal displacements near the phase transition [22, 23], which reflects the strong anharmonicity in this regime.

C. Heat capacity

To further investigate the nature of the phase transition and anharmonic effects in SnSe, we measured the heat capacity. The measurements were performed with a Netzsch DSC 404C differential scanning calorimeter, with the sample loaded inside a Pt crucible, under an ultrapure Ar purge gas cycled through a Ti gettering furnace. The scans were performed after careful evacuation and purging of the sample chamber. The heating and cooling rates were 20 K/min and 20 K/min, respectively. A sapphire standard and empty-crucible baseline measurements were performed in identical conditions. The heat capacity curves measured during heating and cooling are shown in Fig. 12. The heat capacity exhibits a lambda shape in the vicinity of the phase transition, akin to the classic case of liquid helium [58]. This behavior is generally indicative of a second-order phase transition, in agreement with the nearly continuous evolution of structural parameters observed with diffraction [22, 24]. The c-polarized lowest-energy transverse optic soft-mode was also shown to continuously condense across $T_c$ in our previous INS study [10]. The phase transition temperature, $T_c \approx 795 \pm 4$ K, obtained from our heat capacity measurements is in good agreement with values (802–813 K) reported in the literature [22, 24]. A slight hysteresis in $T_c$ of about 4 K was observed in our DSC. Some of the hysteresis may possibly be caused by some Se evaporation at high $T$, oxidation, or temperature lags in the instrument. However, it is also possible for the transition to exhibit a partially first-order character, since there is some reported evidence for a small latent heat [27, 29]. We point out that the values of $C_p$ from our DSC measurements (Fig. 12), while in excellent agreement with those of Sassi et al. [19], are significantly larger than the linear estimate from laser flash measurements reported in Ref. [18] which also misses the phase transition behavior. For example, at 700 K, our $C_p$ measurement is 16% larger than the linear estimate of Zhao et al. [18], and at 790 K the discrepancy approximately reaches a factor of two. However, the discrepancy is more minimal above the phase transition, in the Cmcm phase.

We calculated the harmonic and dilational heat capacity from INS phonon DOS and thermal expansion coefficient measurements. The harmonic ($C_v$) and dilational ($C_d$) heat capacity are given by:

$$C_v = k_B \left( \frac{\epsilon_i}{k_BT} \right) \frac{\exp(\epsilon_i/k_BT)}{(\exp(\epsilon_i/k_BT) - 1)^2},$$

and

$$C_d = C_p - C_v = \frac{\alpha_{\text{vib}}(T^3V(T))T}{\chi(T)},$$
FIG. 11. a) Partial phonon DOS of tin and selenium calculated from DFT simulations for \(a_0b_0c_0\), the ground state equilibrium lattice parameters for fully relaxed structure, b) projected phonon DOS of Sn parallel to \(a\) axis, and c) projected phonon DOS of Sn parallel to \(b\) and \(c\) axis compared with partial phonon DOS of Sn from NRIXS measurements at room temperature.

TABLE VI. Mean square thermal displacement parameter \(<|U_\alpha(j)|^2>\) of tin and selenium calculated from DFT (LDA) simulations at \(T = 300\) K compared with experimental neutron diffraction and NRIXS data.

|          | Sn      | Se      |
|----------|---------|---------|
|          | This work (DFT) | This work (NRIXS) | Ref. [22] | This work (DFT) | Ref. [22] |
| \(U_{11}\) | 0.0165  | 0.0136  | 0.0151 | 0.0143 | 0.0150 |
| \(U_{22}\) | 0.0148  | 0.0130  | 0.0130 | 0.0118 | 0.0107 |
| \(U_{33}\) | 0.0170  | –       | 0.0177 | 0.0126 | 0.0143 |

respectively. To calculate the harmonic heat capacity from Eq. (19), we have used the non-neutron-weighted version of phonon DOS at 10 K, as described earlier for the entropy calculations. The comparison of experimental measurements with harmonic and dilational heat capacity is shown in Fig. 12. For \(T < 500\) K, the quasi-harmonic (harmonic+dilational) heat capacity accounts well for the measured \(C_p\). However, as the phase transition is approached, the difference between the two curves increases significantly, reflecting the growing contribution of anharmonicity. A similar behavior was also seen in entropy, internal energy and free energy (Fig. 8, 9, and 10).

VI. CONCLUSION

We have investigated the anharmonic thermodynamics and negative thermal expansion in SnSe with a combination of INS, NRIXS, calorimetry measurements, and first-principles simulations. We identified a pronounced contribution of anharmonicity at high temperature, especially within \(\sim 200\) K of the structural phase transition. The NTE along the \(c\)-axis, the direction of corrugation of Sn-Se bilayers in the Pnma phase, can be qualitatively accounted for with a quasi-harmonic free energy minimization, but deviations arising from strong anharmonicity are evident at high temperatures. The structural distortion accompanying the NTE can be traced to rotations of \(\text{SnSe}_3\) tetrahedra, which overlap with the strongly anharmonic soft-modes condensing at the phase transition.

FIG. 12. Heat capacity of SnSe measured with DSC on single-crystal samples (both heating and cooling), and its harmonic, dilational, and nonharmonic components, indicating the strong anharmonicity around the phase transition. The green dots are data measured on polycrystalline samples by Sassi et al. [19]. The cyan triangles are data measured on single crystal samples by Zhao et al. [18].
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