

Anomalous pressure dependence of thermal conductivities of large mass ratio compounds

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The lattice thermal conductivities (κ) of binary compound materials are examined as a function of hydrostatic pressure P using a first-principles approach. Compounds with relatively small mass ratios, such as MgO, show an increase in κ with P , consistent with measurements. Conversely, compounds with large mass ratios that create significant frequency gaps between acoustic and optic phonons (e.g., BSb, BAs, BeTe, BeSe) exhibit decreasing κ with increasing P , a behavior that cannot be understood using simple theories of κ . This anomalous P dependence of κ arises from the fundamentally different nature of the intrinsic scattering processes for heat-carrying acoustic phonons in large mass ratio compounds compared to those with small mass ratios. This work demonstrates the power of first-principles methods for thermal properties and advances a broad paradigm for understanding thermal transport in nonmetals.

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Introduction. Pressure dependence gives important insight into the understanding of a range of key materials properties. In 1924, Bridgman published the first measurements of thermal conductivities κ of several nonmetallic solids as a function of compressive hydrostatic pressure P [1]. κ for these compounds increased with P from that at ambient pressure (i.e., $P = 0$). Since then, measurements of the pressure dependence of κ have been made in numerous nonmetals including alkali halides [2–4], CuCl [5], ice [6,7], and those residing in the Earth's interior, such as MgO [8,9]. In all systems not approaching a pressure-induced phase transition (PIPT) [10], κ was found to increase with P , suggesting that this is a universal feature.

Explanations of the pressure dependence of κ in nonmetallic crystalline materials have typically relied on the theory of Liebfried and Schlömann (LS) [11], which predicts that κ must increase with P [12], i.e., $d\kappa/dP > 0$. However, LS gives an empirical description of κ that relies on a number of approximations, which leads one to ask the following questions: Is $d\kappa/dP > 0$ really a universal property of materials, or are there materials for which $d\kappa/dP < 0$? What are the underlying physical properties that govern $d\kappa/dP$?

To address these questions, we have implemented first-principles calculations of pressure-dependent phonon thermal transport in a large range of materials. We show that for a class of materials, κ decreases with P , i.e., $d\kappa/dP < 0$, a behavior not thought possible for materials far from a PIPT. In binary compounds, this anomalous behavior occurs when the mass ratio of the atoms is large enough to eliminate most of the intrinsic scattering of the heat-carrying acoustic phonons by optic phonons.

First-principles thermal transport. Intrinsic thermal resistance in nonmetals arises from the anharmonicity of the interatomic potential, which causes interactions between phonons [13]. For the cubic systems examined here, the thermal conductivity is a scalar,

$$\kappa = \kappa_{\alpha\alpha} = \frac{1}{3} \sum_{\lambda} C_{\lambda} v_{\lambda}^2 \tau_{\lambda}, \quad (1)$$

where the sum is over phonon modes $\lambda = (\mathbf{q}, j)$, with \mathbf{q} the phonon wave vector and j the phonon polarization, ω_{λ} is the phonon frequency, $C_{\lambda} = \hbar\omega_{\lambda}(\partial n_{\lambda}^0/\partial T)/V$ is the mode specific heat at constant volume, $n_{\lambda}^0 = 1/[\exp(\hbar\omega_{\lambda}/k_B T) - 1]$ is the Bose distribution, V is the crystal volume, v_{λ} is the group velocity, and τ_{λ} is the phonon transport lifetime obtained from the nonequilibrium phonon distribution, $n_{\lambda} = n_{\lambda}^0 + n_{\lambda}^1$ [14,15].

The first-principles approach for determining κ has been described previously [16–29], including examination of κ as a function of P [27–29]. This approach has no adjustable parameters and has demonstrated very good agreement with measured κ for a variety of systems, which has validated its predictive capability. The present approach uses a full, iterative solution of the Peierls-Boltzmann equation for n_{λ}^1 . The only inputs are harmonic and anharmonic interatomic force constants (IFCs). The harmonic IFCs were calculated within density functional perturbation theory [30] using the QUANTUM ESPRESSO package [31,32]. Calculations used the local density approximation and norm-conserving pseudopotentials for the core electrons. Typically, an 80–100 Ryd plane-wave energy cutoff was used with $6 \times 6 \times 6k$ -point meshes for the electronic structure and phonon integrations. Anharmonic IFCs were calculated within density functional theory [33,34] using 80–100 Ryd plane-wave energy cutoffs and gamma point calculations in 216 atom supercells with interactions from the third or fifth nearest neighbors of the unit cell atoms. Further details are in Ref. [21]. The harmonic IFCs give the phonon frequencies, velocities, and specific heat. Anharmonic IFCs are required for the three-phonon scattering rates. We take κ to be limited by only intrinsic three-phonon scattering [35]. The ground state is determined by adjusting the lattice constant a to minimize the total energy. For each system, we decrease a from this value to get a hydrostatic pressure [31] at which we calculate the harmonic and anharmonic IFCs.

Results. We have calculated the pressure dependence of κ for a large number of elemental, III-V, and II-VI cubic compounds. We have chosen cubic materials in part because the isotropy of κ gives simpler and more intuitive changes

TABLE I. Calculated values for κ_0 , the mass ratio, the a - o gap (scaled by the largest acoustic frequency), $(d\kappa/dP)/\kappa_0$, comparing *ab initio* values with those from LS theory for several materials at $T = 300$ K [37]. The $\phi_{aaa}^{3\text{-ph}}$ and $\phi_{aao}^{3\text{-ph}}$ are the three-phonon scattering phase spaces [Eq. (2)]. The up and down arrows indicate increases or decreases with P . The double arrows give the type of scattering processes (aaa or aao) that dominate the behavior of κ with P .

	κ_0 (W m ⁻¹ K ⁻¹)	Mass ratio	Scaled a - o gap	$\frac{1}{\kappa_0} \frac{d\kappa}{dP} \Big _{ab\text{initio}}$ (GPa ⁻¹ × 100)	$\frac{1}{\kappa_0} \frac{d\kappa}{dP} \Big _{\text{LS}}$ (GPa ⁻¹ × 100)	$\phi_{aaa}^{3\text{-ph}}$	$\phi_{aao}^{3\text{-ph}}$
MgO	72.45	1.52	0	5.8	3.4	↑	↓
BN	2157	1.30	0	0.9	2.8	↑	↑
GaAs	55.4	1.07	0.09	3.9	19	↓	↑
SiC	572.0	2.34	0.19	2.1	3.5	↑	↓
BP	554.1	2.87	0.36	4.1	4.7	↑	↓
BA _s	3239	6.93	0.97	-1.9	4.1	↑	N/A
BeSe	621.6	8.76	1.08	-1.6	8.3	↑	N/A
BSb	1087	11.26	1.42	-2.4	5.7	↑	N/A
BeTe	367.6	14.16	1.68	-2.8	14	↑	N/A

in κ with P . Here, we provide illustrative results for MgO, GaAs, SiC, BN, BP, BSb, BA_s, BeTe, and BeSe, a set of compounds with widely varying mass ratios (see Table I). The last four have much larger mass ratios of constituent atoms than do the first five. This gives qualitatively different pressure dependences of κ . We note that for the materials considered here, pressures are far less than those at which measured or calculated phase transitions occur, e.g., BA_s (125 GPa) [38], BeSe (56 GPa) [39], BeTe (35 GPa) [39], and BSb ($P = 56$ GPa) [40]. For several compounds we examined—CuCl, Si, Ge, AlSb, InAs, and InSb—calculated transverse acoustic (TA) frequencies soften significantly with small applied P , indicating an approach to a PIPT. We did not include these materials here since the focus of the present Rapid Communication is identifying and explaining thermal transport behavior for which conventional theory fails both qualitatively and quantitatively in predicting $\kappa(P)$.

Figure 1 shows the phonon dispersions for cubic MgO, an important constituent of the Earth's lower mantle whose P -dependent κ has attracted wide attention [8,9,27]. The calculated phonon frequencies for both $P = 0$ and $P > 0$ agree well with measurements, reflecting the accuracy of our harmonic IFCs. Note that increasing P increases phonon frequencies.

The left panel in Fig. 2 gives the calculated and measured [9] κ for MgO as a function of P at $T = 300$ K. Inclusion of phonon scattering by isotopes, impurities, and sample boundaries [44,45] gives reasonable agreement with the measured κ and confirms $d\kappa/dP > 0$. Calculated κ/κ_0 vs P for BSb, BA_s, BeTe, and BeSe are shown in the right panel. Values for κ_0 , the isotopically pure κ at $P = 0$, are given in Table I. Although the changes in the phonon dispersions of these large mass ratio compounds with P are similar to those in MgO (see Fig. S1 in Ref. [46]), κ decreases with P , opposite to the expected behavior. Table I illustrates this distinction between the small and large mass ratio compounds for a number of materials. Note that, in contrast to the *ab initio* results, the LS theory (discussed below) gives $d\kappa/dP > 0$ for all materials. We explain the different behaviors in terms of the fundamentally different scattering processes that govern κ in these systems.

Equation (1) shows that $\kappa(P)$ can be expressed in terms of two features: changes in specific heat and phonon velocities ($C_\lambda v_\lambda^2$), and changes in phonon lifetimes (τ_λ). For the large

range of materials examined here, the change in κ with P is governed primarily by changes in phonon lifetimes. Thus, in order for κ to decrease with increasing P , phonon lifetimes must decrease.

To understand how τ_λ changes with P , we group three-phonon scattering processes according to the number of participating acoustic (a) and optic (o) phonons. While heat is carried primarily by acoustic phonons, their scattering by optic phonons contributes significantly to thermal resistance. Typically, two intrinsic three-phonon scattering processes dominate: $a + a \rightarrow a$ (aaa) and $a + a \rightarrow o$ (aao) (see Fig. S2 in Ref. [46]). For materials with large longitudinal optic (LO)-transverse optic (TO) splitting, aao processes can also give non-negligible contributions at low frequency. However,

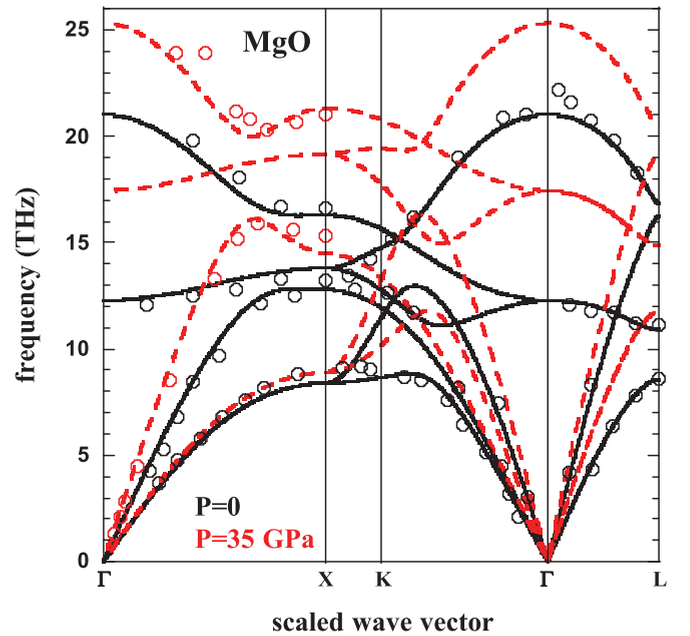


FIG. 1. (Color online) Calculated phonon dispersions for MgO at $P = 0$ (solid black curves) and $P = 35$ GPa (dashed red curves). Black circles give measured data from Refs. [41,42], and red circles give measured data from Ref. [43]. Note the upward frequency shifts of phonons with P .

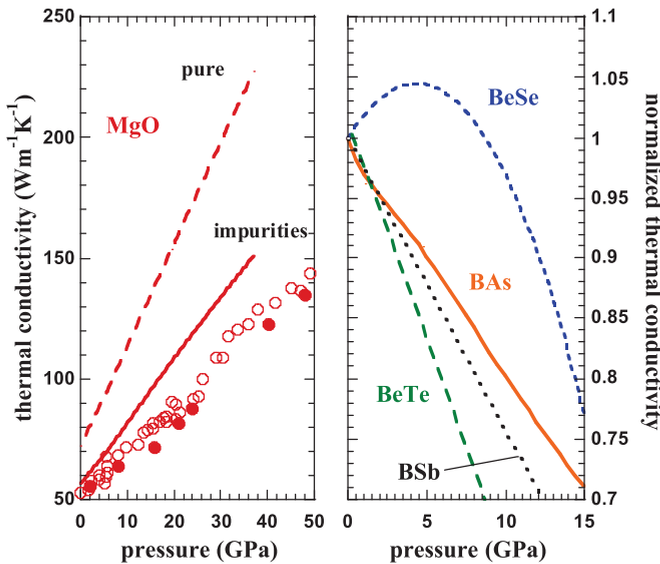


FIG. 2. (Color online) Left panel: Calculated κ (dashed red line) and that including isotopes, impurities, and sample boundaries (solid red line), along with measured thermal conductivity (red circles), vs P for MgO [9]. Right panel: κ/κ_0 for BSb (dotted black), BAs (solid orange), BeTe (dashed green), and BeSe (short dashed blue). BeSe has a relatively large contribution to the three-phonon scattering phase space from processes involving one acoustic phonon and two optic phonons. The decrease in this contribution as P increases from zero gives the initial increase in κ seen here in this figure.

for the materials considered here, the behavior of κ is still dominated by *aaa* and *aaO* processes. Changes in τ_λ can be understood from the *aaa* and *aaO* contributions to the phonon-phonon scattering rates and their P dependence. A key factor is the size of the frequency gap between acoustic and optic phonon branches (*a-o* gap). For small *a-o* gap materials, such as MgO, both *aaa* and *aaO* processes play significant roles in limiting κ . With increasing P , *aaO* scattering rates typically decrease (see Fig. S3 in Ref. [46]) because the upward shift of the optic modes gives decreased optic phonon populations and decreased phase space of *aaO* processes, defined below. This reduction dominates the change in κ , giving $d\kappa/dP > 0$ (see Table I) around room temperature, in qualitative agreement with measured data [2–4,6–10]. It is interesting that the value $(d\kappa/dP)/\kappa_0 = 0.009 \text{ GPa}^{-1}$ given in Table I for BN (also obtained independently from the SHENGBTE code [47]) is significantly larger than that given in Ref. [29], which gives $(d\kappa/dP)/\kappa_0 = 0.003 \text{ GPa}^{-1}$ for $P < 114 \text{ GPa}$. Further, our calculations give a roughly linear dependence of κ with P up to $P = 300 \text{ GPa}$, unlike that in Ref. [29]. The anharmonic IFCs calculated in the present work simultaneously satisfy point-group and derivative permutation symmetries and translational invariance conditions. This was not the case for anharmonic IFCs used in Ref. [29] and resulted in low-frequency scattering rates that were too large, which then gave suppressed and bimodal behavior of $\kappa(P)$ for BN [see Fig. 1(b) of Ref. [29]] compared to the larger $\kappa(P)$ values and linear trend for BN obtained by us.

In contrast to $\kappa(P)$ for the small mass ratio materials, the κ for BSb, BAs, BeTe, and BeSe decrease with P . To understand

this behavior, note that the size of the *a-o* gap is controlled in part by the mass ratio $m_{\text{heavy}}/m_{\text{light}}$ of the atoms. As $m_{\text{heavy}}/m_{\text{light}}$ increases from one (the smallest possible value), the *a-o* gap increases. For sufficiently large $m_{\text{heavy}}/m_{\text{light}}$, the *a-o* gap becomes so large that *aaO* processes are no longer possible because phonon energy and momentum cannot be conserved. This *aaO* freeze-out condition occurs roughly when the *a-o* gap exceeds the maximum acoustic phonon frequency, which is satisfied for the large mass ratio compounds in Fig. 2 (see Table I). In such cases, τ_λ and κ depend on *aaa* scattering. Acoustic phonon branches are typically bunched together in large mass ratio compounds. In these, TA and longitudinal acoustic (LA) branches separate with increasing P , giving increased *aaa* phase space and phonon-phonon scattering rates (see Figs. S1 and S4 in Ref. [46]). The resulting smaller acoustic phonon lifetimes prevail against the increased acoustic velocities, giving decreasing κ with P .

Consider how κ and $d\kappa/dP$ vary as the size of the *a-o* gap is increased [48]. Figure 3 shows the calculated κ at 300 K for $P = 0$ and for $P = 16 \text{ GPa}$ of a hypothetical crystal in which BAs IFCs are retained but m_2/m_1 is varied, keeping the unit cell mass fixed: $m_1 + m_2 = m_B + m_{AS} = 85.73 \text{ amu}$. This changes the phonon frequencies and the size of the *a-o* gap. For $m_2/m_1 = 1$, the *a-o* gap is zero, so both *aaO* and *aaa* scattering rates are strong, giving much reduced κ compared to the actual BAs compound. Then increasing P increases κ since the reduction in *aaO* scattering rates with P has a larger

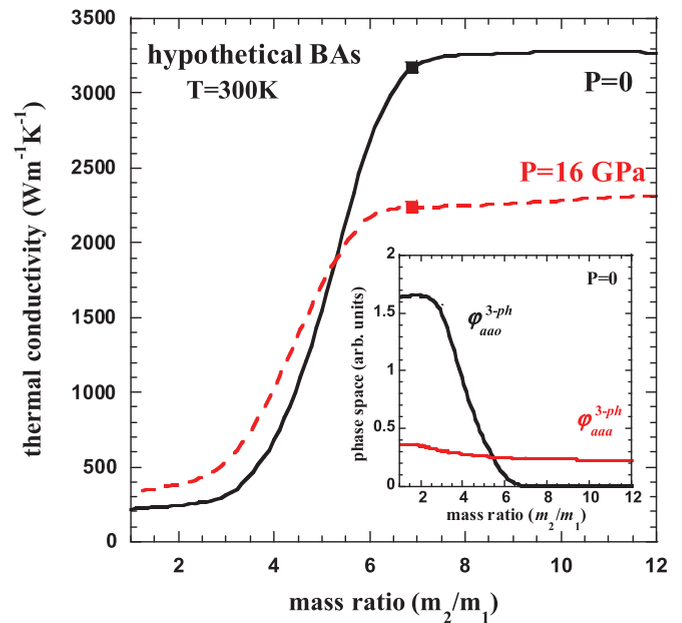


FIG. 3. (Color online) Calculated κ vs mass ratio (m_2/m_1) for a hypothetical BAs material for which m_1 (light mass) and m_2 (heavy mass) are varied, but $m_1 + m_2$ stays fixed at $m_1 + m_2 = m_B + m_{AS}$. Solid black and dashed red curves give κ using the BAs IFCs for $P = 0$ and $P = 16 \text{ GPa}$, respectively. The square symbols on each curve give the calculated κ for the actual BAs mass ratio, $m_{AS}/m_B = 7$, which occurs near the freeze-out condition $\phi_{aaO}^{3-ph} = 0$. The inset gives the phase space for *aaa* scattering, ϕ_{aaa}^{3-ph} (red curve), and *aaO* scattering, ϕ_{aaO}^{3-ph} (black curve), vs m_2/m_1 for the hypothetical material for $P = 0$.

effect than the increase in *aaa* scattering rates. As m_2/m_1 increases, zone boundary LA frequencies decrease while optic frequencies increase. This widens the *a-o* gap, causing the *aaa* scattering rates to decrease and κ to increase. Also, increasing m_2/m_1 pushes the LA and TA branches together, which decreases the phase space for *aaa* scattering. There is a crossover in the $P = 0$ and $P = 16$ GPa curves at $m_2/m_1 \approx 5$. For $m_2/m_1 < 5$, κ increases with P , i.e., $d\kappa/dP > 0$, while for $m_2/m_1 > 5$, $d\kappa/dP < 0$. This reflects the transition between the region where *aaa* scattering dominates ($m_2/m_1 < 5$) and that where *aoa* scattering dominates ($m_2/m_1 > 5$). We note that the large mass ratio compounds considered have strong chemical bonds and reasonably small LO-TO splittings. Compounds with weak bonds and/or large LO-TO splittings would require correspondingly larger mass ratios to achieve the freeze-out condition.

The above behavior can be understood qualitatively by examining the three-phonon scattering phase space [49],

$$\phi_{jj'j''}^{3\text{-ph}} \propto \sum_{\mathbf{q}\mathbf{q}'} \left[\delta(\omega_{\mathbf{q}\mathbf{j}} + \omega_{\mathbf{q}'\mathbf{j}'} - \omega_{(\mathbf{q}+\mathbf{q}'-\mathbf{K})\mathbf{j}''}) + \frac{1}{2} \delta(\omega_{\mathbf{q}\mathbf{j}} - \omega_{\mathbf{q}'\mathbf{j}'} - \omega_{(\mathbf{q}-\mathbf{q}'-\mathbf{K})\mathbf{j}''}) \right], \quad (2)$$

which is constrained by the energy and momentum conservation conditions $\omega_{\mathbf{q}\mathbf{j}} \pm \omega_{\mathbf{q}'\mathbf{j}'} = \omega_{\mathbf{q}''\mathbf{j}''}$ and $\mathbf{q} \pm \mathbf{q}' = \mathbf{q}'' + \mathbf{K}$, where \mathbf{K} is a reciprocal lattice vector. For small mass ratio, the *aoa* processes contribute significantly to $\phi_{aaa}^{3\text{-ph}}$, and these, combined with the *aaa* processes, give a large phase space for three-phonon scattering (see the inset of Fig. 3). With increasing m_2/m_1 , the *aoa* and, to a lesser extent, the *aaa* phase spaces decrease, contributing to the increase in κ seen in Fig. 3. Beyond the *aoa* freeze-out condition, only *aaa* scattering occurs, giving a phase space that weakly decreases with increasing mass ratio. Table I shows that the *aoa* phase space typically decreases with increasing P while the *aaa* phase space typically increases with increasing P (see Figs. S3 and S4 in Ref. [46]).

Why does the LS theory not capture the decrease in κ with P ? It gives [7,11] $\kappa = AV^{1/3}\omega_D^3/(\bar{\gamma}^2T)$, where A is a constant, ω_D is the acoustic Debye frequency, and $\bar{\gamma}$ is a mode-averaged Grüneisen parameter [46]. Upon compression, V decreases, while ω_D increases as phonons are shifted to higher frequencies, and $\bar{\gamma}$ decreases because of the stiffening of the interatomic potential. The decrease in V is more than compensated by the increase from $\omega_D^3/\bar{\gamma}^2$, giving an

increased κ . Phonon-phonon scattering rates are affected by anharmonicity and by the scattering phase space. The LS theory represents the anharmonicity through $\bar{\gamma}$. However, it does not capture the details of the three-phonon phase space. This phase space and its changes with P can be highly variable across materials. With increasing P the energy-momentum conserving surfaces for *aaa* processes become larger. On the other hand, for *aoa* processes, the *a-o* gap increases more rapidly than the maximum acoustic frequency, which typically decreases the *aoa* phase space. Such changes are not accounted for in LS theory. For small *a-o* gap materials, they contribute to give $d\kappa/dP > 0$, consistent with LS theory. For large *a-o* gap materials, they contribute to give $d\kappa/dP < 0$, contrary to LS theory.

Summary. The pressure (P) dependence of the lattice thermal conductivities κ of a large number of cubic binary compounds has been studied using an *ab initio* theory of thermal transport. For compounds with similar masses, κ increases with P because anharmonic three-phonon scattering involving two acoustic phonons and one optic phonon (*aoa* processes) gets weaker. This increase is consistent with previous measurements and with simple theory. In contrast, for large mass ratio compounds having a large frequency gap between the acoustic and optic phonons, *aoa* processes do not occur. Then scattering between acoustic phonons limits κ , which decreases with P due to increased scattering phase space. Such behavior is not predicted by the simple theory [11] commonly used to analyze $\kappa(P)$, and it has not been observed previously in crystals far from a pressure-induced phase transition. These results demonstrate the power of *ab initio* methods for elucidating the underlying physics of thermal transport and motivate the growth of materials with large mass ratios.

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