Importance of non-parabolic band effects in the thermoelectric properties of semiconductors

Xin Chen, David Parker & David J. Singh

Materials Science and Technology Division, Oak Ridge National Laboratory, Oak Ridge, TN 37831-6056.

We present an analysis of the thermoelectric properties of of n-type GeTe and SnTe in relation to the lead chalcogenides PbTe and PbSe. We find that the singly degenerate conduction bands of semiconducting GeTe and SnTe are highly non-ellipsoidal, even very close to the band edges. This leads to isoenergy surfaces with a strongly corrugated shape that is clearly evident at carrier concentrations well below 0.005 e per formula unit (7–9 × 10^{19} cm^{-3} depending on material). Analysis within Boltzmann theory suggests that this corrugation may be favorable for the thermoelectric transport. Our calculations also indicate that values of the power factor for these two materials may well exceed those of PbTe and PbSe. As a result these materials may exhibit n-type performance exceeding that of the lead chalcogenides.

Electronic properties of semiconductors, and thermoelectrics in particular, are typically understood using parabolic band models. Within these models the electronic structure and transport is governed by an effective mass, and anisotropy is quantified by the use of anisotropic effective mass tensors. Non-parabolicity in finite gap semiconductors is normally characterized by an energy and/or temperature dependent mass, and is taken as a weak effect as it must be sufficiently close to the band edge with a finite gap. Transport is then understood in terms of parabolic band expressions with a varying effective mass. In the context of thermoelectrics these expressions imply limitations to performance. Specifically, thermoelectric materials, which are heavily doped semiconductors, require both high thermopower and high conductivity. However, for a given carrier concentration and temperature the thermopower increases with effective mass, while the mobility (and conductivity) decreases.

Here we elucidate a different effect of non-parabolicity that can overcome this difficulty. Within a parabolic band picture, the isoenergy surfaces in the band structure, which govern transport take the shape of ellipsoids. In large Fermi surface metals and in cases where the bands are degenerate at the band edge (e.g. the t_{2g} derived conduction band minimum of some transition metal oxides) more complex shapes are possible, and this can be beneficial in overcoming the above conundrum. Here we show that there can be large deviations from the ellipsoidal shapes that are usually described by an effective mass tensor even for relatively low carrier concentrations of relevance to thermoelectrics. This is the case even without band degeneracy. Furthermore, this electronic feature may be beneficial for thermoelectric properties. Specifically, we discuss two narrow gap semiconductors, GeTe and SnTe, which have conduction band structures that are not only anisotropic (i.e. with different effective masses in different directions) and non-parabolic, but in fact also have a novel corrugated structure in the doping range near the band edges that appears to affect thermoelectric performance. This cannot be captured within parabolic band models, even with an anisotropic effective mass tensor.

We show by first principles calculations that the performance of these n-type materials may potentially outstrip that of the n-type lead chalcogenides PbTe and PbSe. This is of considerable importance given that the best n-type ZT for these chalcogenides has significantly lagged the best reported p-type values of ZT (see for example Ref. 3). We contrast these two materials with PbTe and PbSe, which do not show this corrugated structure.

We begin with the observation that many of the high-ZT materials, such as PbTe, Bi_{2}Te_{3}, and PbSe have highly anisotropic band structures. Typically, they have Fermi surfaces with rather different masses along different directions. When the symmetry of the material is taken into account, Fermi surface pocket degeneracy ensures that all directions (in the case of the cubic rock-salt structure compounds given above) or all planar directions (for rhombohedral Bi_{2}Te_{3}) average transport from both heavy and light bands. This leads to both high thermopower and high electrical conductivity. This thus represents one means of solving the conundrum of attaining both these...
The chalcogenides considered here have conduction band minima at the L-point of the fcc Brillouin zone. The top panel of Fig. 1 shows isoenergy surfaces for SnTe near the conduction band minimum, while the bottom panel of Fig. 1 shows a comparison of the four compounds, GeTe, SnTe, PbTe, and PbSe. These are based on calculations with augmented plane-wave plus local orbital (APW + lo) method, as implemented in the Wien2K code. We included spin-orbit coupling in our calculations and employed the modified Becke-Johnson potential of Tran and Blaha (TB-mBJ), which can generally yield accurate electronic band structures and gaps for simple semiconductors and insulators. Within a parabolic band model, these constant energy surfaces must take the shape of an ellipsoid of revolution about the Γ-L line, which the actual surfaces do for very low carrier concentrations. It is clear from Fig. 1 that the electron pockets in SnTe have a corrugated shape, obviously different from that in PbTe and PbSe, where the pockets are relatively close to ellipsoids. This is even more noticeable in GeTe. Note that there is still substantial Kane-band (i.e. quasilinear dispersion) non-parabolicity in all four materials (as is evident in the band structure plots in Figure 2), but the corrugated character is absent in PbTe and PbSe.

We note also from the Table that the difference in heavy and light masses (a form of anisotropy) is greatest for SnTe and GeTe and significantly smaller for PbTe and PbSe. This is also a contributor to improved transport in these materials, as the lighter band leads to good conductivity and the heavier band to good thermopower, a scenario explored in Ref. 27.

A simple argument (supported by calculations, below) explains the benefits to thermoelectric transport of the corrugated Fermi surfaces. Briefly, the effect of the corrugation is to increase the area of the Fermi surface for a given carrier concentration (the volume contained within the Fermi surface). A larger area implies the presence of more electronic states, or equivalently a larger electronic density of states in the same bulk material. The chalcogenides considered here have conduction band minima at the L-point of the fcc Brillouin zone.

The top panel of Fig. 1 shows isoenergy surfaces for SnTe near the conduction band minimum, while the bottom panel of Fig. 1 shows a comparison of the four compounds, GeTe, SnTe, PbTe, and PbSe. These are based on calculations with augmented plane-wave plus local orbital (APW + lo) method, as implemented in the Wien2K code. We included spin-orbit coupling in our calculations and employed the modified Becke-Johnson potential of Tran and Blaha (TB-mBJ), which can generally yield accurate electronic band structures and gaps for simple semiconductors and insulators. Within a parabolic band model, these constant energy surfaces must take the shape of an ellipsoid of revolution about the Γ-L line, which
scattering time $t$ terms of the effective mass $m$. For an isotropic parabolic band, the conductivity can be expressed in terms of the density of states and the carrier concentration $n$, $\sigma = ne^2/\tau m^*$. Since the electrical conductivity intrinsically involves a scattering time $\tau(E) \sim E$, where $r$ is the scattering parameter, here we adopt two different approximations on $\tau$. One is the constant scattering time approximation (CSTA) ($r = 0$), as widely used in ab-initio calculations, including those in this paper. In the other approximation, $\tau(E)$ is considered to be affected by the acoustic-mode lattice scattering ($r = -1/2$), as considered in Refs. 15–18, which we here note as ASTA. Then, considering the band degeneracy of 8 for our cases here (2 for spin and 4 coming from the number of electron pockets), we can derive the following expressions for thermopower within CSTA and ASTA according to the Mott formula $^{19}$,

$$S = \left(\frac{\pi^2 k_B^2}{3e} T \frac{d\log[\tau(E)]}{dE}\right)_{E=E_F},$$

where $E_F$ is the Fermi energy,

$$S_{\text{CSTA}} = \frac{4\pi^2 k_B^2}{3e} m^* T \left(\frac{4\pi^2}{3n}\right)^{2/3}$$

$$S_{\text{ASTA}} = \frac{8\pi^2 k_B^2}{3e\hbar^2} m^* T \left(\frac{4\pi^2}{3n}\right)^{2/3}$$

It is clear that with parabolic bands the thermopower is proportional to $m^n$ for both approximations. Note that the main effect of the ASTA is a reduction in thermopower relative to the CSTA. For the comparison of SnTe and GeTe with PbTe and PbSe we adopt the CSTA for all materials for consistency and note that the likely effect of using ASTA would simply be a reduction in calculated thermopower for all four materials. As will be seen, the likely advantages of SnTe and GeTe may well dwarf any potential effects of neglecting the relaxation time energy dependence.

Note also that the Mott relation is inherently a single band formula and neglects the effects of bipolar conduction. However, as is the case here, the region of likely optimal doping (i.e., likely largest $ZT$) is generally at concentrations heavier than that where bipolar conduction becomes significant (i.e. to the right of the thermopower maxima in Figure 4), where the single band Mott formula generally applies.

Table 1 | Effective mass for lowest conduction band of GeTe, SnTe, PbTe, and PbSe (in m_e units) along transverse (W–L) and longitudinal (L–L) directions

| Compound | m_W | m_L |
|----------|-----|-----|
| GeTe     | 0.022 | 0.021 |
| SnTe     | 0.49 | 0.18 |
| PbTe     | 0.040 | 0.32 |
| PbSe     | 0.081 | 0.18 |

To ensure a fair comparison with the specific compounds examined here, we also calculate the effective masses for the lowest conduction bands of GeTe, SnTe, PbTe and PbSe, as shown in Table 1. These masses were calculated from a parabolic fitting of the dispersion very near the conduction band minima, with axes chosen as the principal axes of the ellipsoids (which run perpendicular to the Brillouin zone face around the L point).

We note that the effective mass of GeTe is the most different along transverse (W–L) and longitudinal directions (L–L), as reflected in the large anisotropy seen in its electron pockets (see Fig. 1). SnTe also shows very anisotropic, non-ellipsoidal pockets although the difference is not as large as in GeTe. Therefore, we chose the heavy effective mass of SnTe (0.18 $m_e$, with $m_0$ the free electron mass) in the parabolic model calculations.

These differences can be seen more directly in Figure 2, which plots the band structures and densities-of-states of the four compounds. The conduction band DOS rises from the band edge much more rapidly for SnTe and GeTe than the other two compounds, indicative of the larger effective DOS mass of these two compounds. This will be seen to be responsible for larger thermopower in these materials, irrespective of the large mass of GeTe relative to SnTe (at 700 K the thermopower samples electronic structures larger than the energy window within which effective masses were determined). Correspondingly, the conduction band for these two compounds shows only a moderate dispersion of 0.9 eV from the L to W points, while the value is roughly double this for the other two compounds. In addition, SnTe and GeTe show near-degeneracies at the band edge, with at least one additional band minima only 0.15 eV above the CBM in GeTe and 0.4 eV above in SnTe. These degeneracy differences, combined with the fact that the light-mass (as shown in the Table) for these two compounds is lower than for PbSe and PbTe, suggest that the conductivity of SnTe and GeTe is likely to be substantial. Indeed this is the case, as Figure 3 depicts the calculated values of $\sigma/\tau$ for all compounds, showing in fact a slight advantage for SnTe and GeTe.

![Figure 3](https://www.nature.com/scientificreports)
Figure 4 | Calculated 700 K thermopower for n-type GeTe, SnTe, PbTe, PbSe, and general isotropic parabolic system within CSTA and ASTA, compared with the experimental data from Refs. 20–24. Note the excellent agreement between theory and experiment for PbTe and the good agreement (±15 percent) for PbSe.

Figure 4 shows the thermopower, calculated within semiclassical Boltzmann transport theory and the CSTA (see the Methods section) at $T = 700$ K. We have chosen this temperature as it is above the rhombohedral to cubic phase transition temperature of 670 K for GeTe, and is also a typical temperature for high temperature thermoelectric work. While for applications such phase transitions can be a negative, in this work we are studying thermoelectric performance irrespective of any application. Note that our first principles calculations follow fairly closely the existing experimental data on PbTe and PbSe, indicating the validity of our approach.

It is clear that the behavior of $S$ for n-type GeTe and SnTe is significantly superior to that of PbTe and PbSe consistent with previous numerical results. The enhanced thermopower in n-type GeTe and SnTe comes from the corrugated Fermi surface character that increases the Fermi surface area so that more electronic states contribute to transport. Furthermore, the thermopower of GeTe and SnTe obviously exceeds the parabolic band model values obtained within CSTA at doping levels larger than $3.6 \times 10^{20}$ and $4.8 \times 10^{20}$ cm$^{-3}$, for the two compounds, respectively. This is the range where the properties are consistent with good thermoelectric performance. The beneficial effect of the non-parabolic band structure is increased in the ASTA. ($S_{\text{CSTA}}$ is one and a half times larger than $S_{\text{ASTA}}$ according to the Eqs. (1) and (2)). The lead salts PbTe and PbSe also have a larger thermopower than the parabolic case within ASTA at doping levels of $> 5.9 \times 10^{19}$ and $3.9 \times 10^{19}$ cm$^{-3}$, respectively.

It is instructive, in order to isolate the effect of the corrugation, to consider the transport quantity $S\tau$ (Fig. 5), as the effective mass in the expressions of $\sigma$ and $S$ cancel in this product. It is clearly seen from Fig. 5 GeTe and SnTe have values that substantially exceed those of the general parabolic case over a wide range of carrier concentration. This appears to be due to the corrugation. PbTe and PbSe show higher $S\tau$ than the parabolic case at a doping level of the order $10^{20}$ cm$^{-3}$. Remembering the larger $S$ in the specific anisotropic non-parabolic compounds, they have a much enhanced thermoelectric performance (proportional to power factor $S^2\sigma$) than those of the parabolic case although actual values clearly depend on the details of the scattering.

Since for the actual quantitative performance what is most critical is the power factor $S^2\sigma$, rather than $S\tau$, in Figure 6 we present a plot of the power factor (with respect to scattering time $\tau$) $S^2\tau$ for each of the four materials for a range of carrier concentration. While mean scattering times can and do vary from material to material, such a difference is likely overshadowed by the band structure effects depicted in Figure 4, which shows the substantial advantage of SnTe and GeTe relative to PbTe and PbSe. At the heavy dopings around $n = 2–3 \times 10^{20}$ cm$^{-3}$, where thermoelectric performance is likely optimal for SnTe and GeTe, the power factor is approximately seven times as large as for the other two materials. While it is true that the relaxation times may differ, we take this plot as strong evidence for good thermoelectric performance in SnTe and GeTe.

As mentioned previously, GeTe forms in a rhombohedral structure below 670 K and we have therefore computed one transport property, the thermopower, in both the high temperature cubic structure and the low-temperature rhombohedral structure, which we depict in Figure 7. (Note that for the rhombohedral case we have taken the conductivity-weighted average of the thermopower, i.e. $S = (\sigma_{x\bar{x}}S_x + \sigma_{y\bar{y}}S_y + \sigma_{z\bar{z}}S_z)/(\sigma_{x\bar{x}} + \sigma_{y\bar{y}} + \sigma_{z\bar{z}})$). The calculations were done at 300 K. There is some difference in the curves, particularly at light doping, but for the heavy dopings around $2–5 \times 10^{20}$ cm$^{-3}$ (where
Conversely, for the given carrier concentration $n$ value as the isotropic light band case with the same Fermi energy.

In evaluating the transport coefficients for an isotropic parabolic single band system. Related arguments have been made in the context of oxides Na$_2$CoO$_2$, SrTiO$_3$, and KTaO$_3$\textsuperscript{2,27,28}, with complex multiband conduction band minima from degenerate $t_2g$ bands or a highly two dimensional metallic band structure as in Na$_2$CoO$_2$\textsuperscript{29}. Here we find that such effects are important even with a singly degenerate band near the conduction band minimum of SnTe and GeTe.

In conclusion, we find that anisotropy beyond the level possible in parabolic, ellipsoidal band models may strongly influence transport properties even for moderately doped semiconductors with finite gaps, and that this appears to be favorable for thermoelectric performance. This is illustrated by detailed calculations for GeTe and SnTe. Based on our Boltzmann transport calculations, the $n$-type performance of these materials may exceed that of the lead chalcogenides PbTe and PbSe. We hope that our current findings will stimulate future experimental exploration of the thermoelectric properties of these materials, particularly $n$-type SnTe and alloys.

**Methods**

Transport properties of a crystalline solid, including the thermopower and conductivity $\sigma$ (here $\tau$ is a measure of the scattering time) are calculated within Boltzmann transport theory, based on the electronic structure\textsuperscript{30}. We performed electronic transport calculations using full-potential APW + lo method, using WIEN2K\textsuperscript{8} including spin-orbit coupling. Experimental lattice constants were used for the four materials studied, with the cubic lattice constants for GeTe as 6.009 A, PbTe 6.661 A, and PbSe 6.121A. With the exception of GeTe, these 300 K lattice constants (the lattice constant for cubic GeTe was measured at 673 K, just above the phase transition). For the GeTe rhombohedral phase, the (300 K) lattice constant in the rhombohedral coordinate system was 4.281 A and the angle between rhombohedral lattice vectors 58.358°. To obtain reliable transport properties, a dense $k$-point mesh comprising approximately 3000 points in the irreducible portion of the Brillouin zone was used. The expressions for the electrical conductivity and the thermopower are:

$$S(T) = \frac{1}{\sigma(T)} \int (\sigma(E) - \mu) \left( \frac{\partial F}{\partial E} \right) dE$$

$$\sigma(T) = \int (\sigma(E) - \mu) \left( \frac{\partial F}{\partial E} \right) dE$$

where $f$ is the Fermi function, $\mu$ is the chemical potential, and $\sigma(E)$ is the transport function given by $N(E)\nu(E)/\hbar(E)$. The transport coefficients are tensors, but we omit the subscripts in the present work because of the cubic symmetry. Calculations based on these formulas with actual band structure have proven useful in rationalizing and predicting transport properties of known compounds\textsuperscript{11–14}. Here we used the BoltzTraP code\textsuperscript{15}. The conductivity can only be calculated with respect to the relaxation time using the preceding expressions. However, within the constant scattering time approximation (CSTA) the thermopower can be obtained directly from the band structure information without adjustable parameters. This approximation consists of assuming that the energy dependence of the relaxation time at a given doping level and temperature is negligible on the scale of $k_BT$, and has shown considerable success in describing the thermopower of a large number of materials.

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