Ab-initio electronic transport in narrow gap semiconductors using the Wigner distribution

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Abstract

Electronic transport properties of semiconductors with small band gaps are often not well described by semiclassical methods, such as the Boltzmann transport equation, because of the missing interaction between carriers whose band energy differences are closer than their linewidth. This limits accuracy of predictions for many topological insulators and materials with complex structure and disorder. We develop a new first-principles formalism that uses the Wigner transform to generalize semiclassical transport models to this regime by including additional quantum effects. We apply the formalism to Bi$_2$Se$_3$, and show that its bulk electronic transport properties at low doping concentrations are dominated by the Zener effect, a mechanism in which carriers transfer charge by tunnelling across the band gap.

Recent years have seen the emergence of topological insulators (TI) as a new important class of materials, thanks to their variety of interesting physical properties and promising applications, such as low-power electronics and robust quantum bits [1, 2, 3, 4, 5]. Most studies focus on the surface states that result from the spin-orbit induced inversion of the bulk band gap, so that TIs behave as insulating crystals in the bulk, and as metals on surfaces. For applications in electronics, it is important to have a thorough understanding of both surface and bulk transport properties of TIs. In fact, several TIs are characterized by a bulk quasiparticle band gap that is much smaller than that of silicon. It is important to note that small band gap systems may behave differently than a silicon-based device. For example, small band gap graphene devices may display a phenomenon called Zener (or Klein) tunneling [6, 7], in which the tunneling of carriers through the band gap can substantially contribute to the electrical conductivity.

Simulations are a convenient tool to estimate the transport properties of a small band gap system. The de-facto tool of choice for first-principles studies of electronic transport properties is the Boltzmann Transport Equation (BTE), which provides estimates of transport properties with remarkable agreement with experimental measurements (e.g. [8, 9, 10, 11, 12, 13]). However, this semiclassical approach is not always sufficient to model electronic transport properties: Zener tunneling, for example, is not captured. Some studies have used the equation of motion for the density matrix as an alternative description that holds for small band gap systems [14], however making difficult to understand the link with semiclassical models. Here instead we aim to extend the range of applicability of the BTE to complex materials, including small band-gap semiconductors such as Bi$_2$Se$_3$, with the benefit of retaining the capability to interpret the results in a simple fashion.

In this work, starting from the Moyal bracket, we derive an equation of motion for electrons, termed the Wigner transport equation (WTE). This equation allows the computation of the full set of
electronic transport coefficients, and in particular the electrical conductivity and Seebeck coefficient, and reduces to the BTE in the limit where the Wigner function is diagonal. We implement this equation with full ab-initio parameters and apply the formalism to the topological insulator Bi$_2$Se$_3$.

We show that at small doping concentrations, the estimates of bulk electronic transport properties deviate significantly from semiclassical estimates, due to the presence of Zener tunneling, which is instead captured by the WTE.

We start by considering the ground state Hamiltonian $H_0$ of a crystal, which we assume to be an independent-particle Hamiltonian with eigenvalues $\epsilon_b(\mathbf{k})$ and Bloch states $\psi_{bk}(\mathbf{x})$, where $\mathbf{x}$ is the position, $\mathbf{k}$ the wavevector and $b$ the band index (for simplicity, we omit the spin index). The ground-state is perturbed by a constant electric-field $E$ which couples with the carriers’ charge $e$ and by the electron-phonon interaction $H_{el-ph}$, so that the total Hamiltonian is $H = H_0 + e \mathbf{x} \cdot \mathbf{E} + H_{el-ph}$.

To derive an equation of motion for such a system, we use the single-particle Wigner function $f$ of the system \[1\], defined as the Wigner transform of the density matrix $\rho$ as

$$f_{bb'}(\mathbf{x}, \mathbf{k}, t) = \sum_{\Delta \mathbf{k}} e^{i\Delta \mathbf{k} \cdot \mathbf{x}} \rho_{bb'}(\mathbf{x} + \frac{\Delta \mathbf{k}}{2}, \mathbf{k} - \frac{\Delta \mathbf{k}}{2}, t),$$

(1)

where $t$ is the time. We build the Wigner function through a transformation of the density matrix $\rho_{bb'}(\mathbf{k}, \mathbf{k}')$ in the reciprocal space representation, although one may also start from a real-space representation. Such Wigner transform consists in a rotation of variables $\mathbf{k}, \mathbf{k}' \rightarrow \frac{\mathbf{k} + \mathbf{k}'}{2}, \frac{\mathbf{k} - \mathbf{k}'}{2}$ combined with a Fourier transform on one variable. The Wigner function operates in a phase-space representation, which is especially useful to draw connections between quantum and classical mechanics.

The evolution of the Wigner function \[16, 17\] is found through a Wigner transform of the equation of motion of the density matrix, and has been shown to be

$$\frac{\partial f_{bb'}(\mathbf{x}, \mathbf{k}, t)}{\partial t} = - \{ \{ f(\mathbf{x}, \mathbf{k}, t), H(\mathbf{x}, \mathbf{k}) \} \}_{bb'},$$

(2)

$$:= \frac{i}{\hbar} \left( f(\mathbf{x}, \mathbf{k}, t) \star H(\mathbf{x}, \mathbf{k}) - H(\mathbf{x}, \mathbf{k}) \star f(\mathbf{x}, \mathbf{k}, t) \right)_{bb'},$$

(3)

where $\{ \{ f, H \} \}$ is the Moyal bracket (the quantum mechanical extension of the Poisson bracket) and the Moyal product $\star$ is defined as:

$$f \star H = f(\mathbf{x}, \mathbf{k}) \exp \left( \frac{i}{2} \left( \frac{\partial}{\partial \mathbf{x}} \cdot \frac{\partial}{\partial \mathbf{k}} - \frac{\partial}{\partial \mathbf{k}} \cdot \frac{\partial}{\partial \mathbf{x}} \right) \right) H(\mathbf{x}, \mathbf{k}),$$

(4)

where $H(\mathbf{x}, \mathbf{k})$ is the Wigner transform of the Hamiltonian, and the arrows indicate that the derivative operator is acting to the left/right operators. We mention in passing that the Moyal bracket implicitly assumes that such derivatives exist; in the supplementary material we discuss how to ensure this by choosing a wavefunction gauge.

We now simplify the Hamiltonian supposing that the electron-phonon interaction is weak and solve the Moyal bracket for the single-particle part of the Hamiltonian; the electron-phonon interaction will be added later as a perturbation. Since we are interested in macroscopic properties, we can make the assumption that only slow spatial variations of the Wigner function are relevant. Therefore, we approximate the Moyal product with the lowest orders in Taylor series and, as detailed in the supplementary material, we find an equation of motion which we term the Wigner Transport Equation
(WTE), that is
\[
\frac{\partial f_{bb}(x, k, t)}{\partial t} + i \frac{1}{\hbar} \left[ \mathcal{H}(k) + d(k) \cdot E, f(x, k, t) \right]_{bb} + \frac{1}{2} \left\{ v(k), \frac{\partial f(x, k, t)}{\partial x} \right\}_{bb} - eE \cdot \frac{\partial f_{bb}(x, k, t)}{\partial k} = - \frac{\partial f_{bb}(x, k, t)}{\partial t} \right|_{\text{coll}} .
\]

where \(\{,\}\) is an anticommutator, \(\mathcal{H}_{bb}(k) = \delta_{bb} \epsilon_{kk}\) is a matrix of electronic energies, \(d_{bb}(k) = (1 - \delta_{bb}) \langle kb | e\mathbf{x} | kb' \rangle\) is a matrix of electric dipoles between two Bloch states (typically used to describe optical excitations), and \(v_{bb}(k)\) is the velocity operator. The electron-phonon scattering operator \(\frac{\partial f_{bb}(x, k, t)}{\partial t}\) \(_{\text{coll}}\) is added as a perturbation to the WTE and is built, as detailed in the appendix, using scattering rates from the Fermi Golden rule [18, 19, 20, 21, 22].

The WTE needs to be solved to obtain an estimate of the single-particle Wigner distribution function. As a first comment, we note that the BTE can be recovered as a simplified limit of the WTE, when the off-diagonal terms \(b \neq b'\) are set to zero. This corresponds physically to a situation when different bands do not couple. This can happen, for example, when the energy difference is too large: if this is the case, neither thermal excitation nor dipole interaction provide sufficient energy to allow for the hopping of one particle from one band to the other. Therefore, the more interesting terms to discuss in the WTE are the off-diagonal terms, which introduce the possibility of additional electronic transitions, or couplings, between different electronic states at a given wavevector \(k\). We also note that, if we neglect the space derivative term, the WTE reduces to the quantum master equation studied in other works [23, 24, 25, 14, 26]; we will later discuss that this additional term allows us to compute the Seebeck coefficient. Additionally, we note that the electronic WTE is conceptually similar to a formalism developed for phonon transport in Ref. [18], although here we develop a simplified derivation and treat dynamics of electrons under an electric field.

To better understand these off-diagonal terms, it is illustrative to estimate the electronic conductivity \(\sigma\). Similar to the procedure used to estimate \(\sigma\) with the BTE, we set to zero time and space derivatives in order to compute the bulk steady-state transport properties, and the resulting WTE can be solved using techniques analogous to those developed for the solution of the BTE (detailed in supplementary material). The WTE can be solved, and transport properties are readily obtained. For example, the charge current density can be computed as
\[
j = \frac{e g_s}{2VN_k} \sum_{kb} \left\{ v(k), f(k) \right\}_{bb},
\]

where \(g_s\) counts the spin degeneracy, \(V\) is the crystal unit cell volume and \(N_k\) counts the number of wavevectors. As detailed in the appendix, one can compute the electronic heat current as well, and thus the complete set of transport coefficients, including electrical conductivity and Seebeck coefficient. The diagonal terms \(b = b'\) correspond to the estimates of electrical conductivity tensor within the BTE formalism \(\sigma^{BTE}_{ij}\), where \(i, j\) are cartesian labels, and is briefly discussed in the supplementary material for completeness. Importantly, the semiclassical result is corrected by an additional term as \(\Delta \sigma^{ij} = \sigma^{BTE}_{ij} + \Delta \sigma^{ij}\) that is
\[
\Delta \sigma^{ij} = \frac{2g_s e^2}{V N_k} \sum_{kb} \frac{\dot{f}_{bb}(k) - \ddot{f}_{bb}(k)}{\epsilon_{bb}(k) - \epsilon_{bb}(k)} \times \frac{\epsilon_{bb}(k) \epsilon_{bb}(k) (\Gamma_{bb}(k) + \Gamma_{bb}(k))}{4(\epsilon_{bb}(k) - \epsilon_{bb}(k))^2 + (\Gamma_{bb}(k) + \Gamma_{bb}(k))^2},
\]

where \(\Gamma_{bb}(k)\) is the electronic linewidth (here, due to electron-phonon interaction) and \(f_{bb}(k)\) the Fermi–Dirac occupation number. We can now make a few key remarks. First, the correction \(\Delta \sigma\) is
positive (note that \( f_b(k) \) is a monotonic function of \( \epsilon_b(k) \)), and therefore the WTE will always adjust the semiclassical prediction of conductivity to higher values. Second, the expression of the electrical conductivity better illustrates the role of the off-diagonal components of the Wigner distribution function. Whenever the energy difference between an electron and a hole is comparable to their linewidth, the two carriers interact. The strength of such interaction is determined by the velocity matrix element \( v_{bb'}(k) \), i.e. the matrix element for the optical transition. As a result, the system allows an additional transport mechanism, known as Zener tunneling, in which electrons propagate by tunneling through the band gap.

All quantities appearing in the WTE are available from first-principles codes and we can therefore apply this formalism using fully ab-initio parameters. In particular, we use the Quantum ESPRESSO [27, 28] software suite for the calculation of electronic and phonon properties [29], Wannier90 [30] to interpolate electronic energies and a linear interpolation of the electron-phonon matrix elements [12] (see Supplementary material for details).

We now apply the formalism to study the intrinsic lattice-limited electronic transport of bulk Bi\(_2\)Se\(_3\). In Fig. 1 we report the band structure [31, 32] and the density of states (DOS) for this narrow-gap semiconductor. Using an LDA functional, we estimate a quasiparticle gap of 0.2 eV, in agreement with experimental estimates [33]. It is worth noting that the DOS increases away from the Fermi level and flattens at energies of approximately -0.8eV and 1.0eV in both valence and conduction bands, indicating that the subvalleys are at approximately 1.8 eV in energy.

In Fig. 2a, we estimate the electron-phonon limited electrical conductivity \( \sigma \) of Bi\(_2\)Se\(_3\) in the in-plane direction as a function of temperature for different values of n-type doping concentrations (p-type is reported in the supplementary material). In particular, dashed lines is the semiclassical estimates \( \sigma^{BTE} \), while solid lines the estimates using the WTE. For the highest doping values, when the chemical potential shifts into the conduction band, the conductivity has the typical metallic-like behavior of decreasing with temperature. Under these conditions, BTE and WTE do not differ significantly, except at larger temperatures.

For smaller doping concentrations the chemical potential lies in the band gap and we thus observe a semiconducting behavior of \( \sigma \) increasing with temperature. The semiclassical model predicts a smaller conductivity than the WTE estimate. In fact, when only a few carriers from the bottom of
Figure 2: Computational estimates of the in-plane electrical conductivity (panel a) and Seebeck coefficient (panel b) of Bi$_2$Se$_3$ as a function of temperature, for different values of electron doping concentration. Solid lines are estimated using the Wigner transport equation, while dotted lines are semiclassical estimates obtained solving the Boltzmann transport equation. For small doping concentrations, the interaction between electrons and holes significantly affect the estimates of transport coefficients. For comparison, we plot the experimental in-plane conductivity of single-crystals at negative dopings of $3 \times 10^{19}$cm$^{-3}$ [34], $7 \times 10^{17}$cm$^{-3}$ [35], $4.4 \times 10^{16}$cm$^{-3}$ and $1.1 \times 10^{17}$cm$^{-3}$ [36] (35-1 and 35-2 respectively).
the conduction band are excited, the average carriers’ group velocity is small, due to the quadratic nature of the band minimum. Therefore, the semiclassical contribution to electrical conductivity tends to be rather small. The WTE corrects this picture, including the Zener tunneling effect [7]. As carriers from valence and conduction band are close in energy, they can interact and contribute to the electrical transport through the tunneling effect, as discussed above. For small dopings, the WTE correction is significant, and can be much larger than the BTE conductivity value. For the smallest value of doping reported (10^{16} \text{cm}^{-3}), this correction is largest at lower temperatures. The doping of 10^{18} \text{cm}^{-3} is an intermediate case, with metallic behavior at low temperatures (and thus smaller WTE correction) and semiconducting (with larger WTE correction) at higher temperatures as the chemical. We can thus conclude that a substantial portion of electrical currents at low doping is carried through the Zener tunneling included in the WTE formalism: the current is not only caused by the carriers traveling at a finite group velocity, but also by carriers’ tunneling between single-particle Bloch states. To validate our estimates, we compare conductivity predictions with experimental measurements on single crystals at low temperatures, showing good qualitative agreement. It’s worth noting that we expect computational estimates of conductivity to be above the experimental measurement, since our simulations include the effect of electron-phonon scattering but neglect the impact of impurities (especially relevant at low temperatures). Therefore, the present simulations must represent a theoretical upper bound for the conductivity of the intrinsic crystal. Indeed, the WTE correctly overestimates the conductivity for every case where experimental data on single crystals is available, while the BTE underestimates it, particularly at low carrier concentrations.

In Fig. 2b, we report the Seebeck coefficient \( S \), solid/dashed lines are WTE/BTE estimates. The negative values indicate that a majority of carriers are electrons, although, in a small band gap system, deviations from this behavior can occur. Under a BTE-like model, \( S \) is approximately proportional to the logarithmic derivative of the density of states [37]. Therefore, at low temperatures, one expects the Seebeck coefficient to increase as the doping concentration is decreased. This phenomenon is crucial to optimize the thermoelectric efficiency, where the goal is to maximize the power factor \( \sigma S^2 \). However, within the WTE, \( S \) is not anymore simply related to the density of states, as additional terms in the transport equations appear (as detailed in the supplementary information). As a result, the large increase of the Seebeck coefficient expected by a semiclassical model at low dopings is completely suppressed by the WTE, and \( S \) becomes comparable to its values of large dopings. We thus conclude that the tunneling effects captured by the WTE can alter considerably the predictions of thermoelectric properties in narrow-gap semiconductors.

We now analyze the energy of the carriers contributing to transport. In Fig. 3a, we analyze the contributions to the BTE electrical conductivity as a function of the carriers’ energy at a doping concentration of 10^{18} \text{cm}^{-3}, and temperature of 700 K. This histogram is built such that the area under the curve integrates to the total electrical conductivity. Within the semiclassical relaxation time approximation, the quantity plotted is an energy-resolved histogram of \[ \sum_{\nu} \frac{\partial f_{\nu}(k)}{\partial\epsilon_{\nu}} \frac{\partial^2 f_{\nu}(k)}{\partial\epsilon^2_{\nu}} \left( \frac{1}{\Gamma(k)} \right), \] i.e. the contribution of a single mode to the BTE electrical conductivity. As expected, the dominant contributions to electrical conductivity come from carriers whose energy is close to the chemical potential \( \mu \) (set at 0 eV). The contributions of other carriers decay exponentially as their energy gets further from \( \mu \).

The WTE correction \( \Delta \sigma \) cannot be resolved in terms of a single carrier’s energy, since it involves the tunneling between two states at different energy. Therefore, in Fig. 3b, we plot the contributions to the electrical conductivity as a function of two interacting carriers energies. On the diagonal, we find again the BTE-like terms shown in Fig. 3a. In addition, we can see important off-diagonal contributions to the electrical conductivity. In particular, there are two peaks of contributions to electrical conductivity, that couple electrons of energy 1.0 eV with holes at -0.8 eV. These two values correspond to the average energies of the top/bottom of valence/conduction bands, when the DOS
reaches maximum values. Therefore, in contrast to the typical intuition of the Zener tunneling, we find that at large temperatures the most significant coupling between carriers takes place far from the chemical potential, with carriers of energy much larger than the thermal energy. For this material, the dipole interaction between carriers in subvalleys of the valence and conduction is thus particularly strong, allowing for high-energy carriers to contribute to transport. As a result, we speculate that Zener tunneling may take place also in semiconductors with a wider gap and contribute significantly to electrical conductivity, provided that the dipole interaction is sufficiently strong.

In conclusion, we have shown that the Moyal equation of motion for the Wigner function leads to a straightforward extension of the Boltzmann transport equation, which takes into account additional quantum transport effects such as Zener tunneling. With this new Wigner transport equation formalism it is possible to compute the full set of Onsager transport coefficients for thermal and electric transport from first principles, starting with density functional perturbation theory. We implemented this equation and applied it to the topological insulator Bi$_2$Se$_3$. We have shown that, while at large doping concentrations the BTE provides a fairly accurate description of transport, it fails at low doping concentrations. At low dopings, the Zener tunneling effect contributes significantly to the electronic transport, changing estimates for both electrical conductivity and Seebeck coefficient. Lastly, we have shown that Zener tunneling does not just take place across the states closest to the band gap, but can involve states that are significantly further apart in energy, provided that the dipole interaction is sufficiently strong. As a result, we have extended the range of applicability of ab-initio transport simulations to materials where quantum tunneling effects couple carriers, and a single-particle description is no longer adequate.
Figure 3: Panel a: histogram of contributions to semiclassical electrical conductivity as a function of the carrier energy, for Bi$_2$Se$_3$ at 700K and n-doping at $10^{18}$ cm$^{-3}$. Panel b, 2D histogram of contributions to the electrical conductivity for the same system as estimates with the Wigner transport equation against the energy of two coupled carriers. Off-diagonal contributions represent electrical conductivity arising from the coupling between electrons and holes.
1 Supplementary material

1.1 Equation of motion of the Wigner function

In this section, we detail the derivation of the Wigner transport equation discussed in the main text.

We start from the single-particle Hamiltonian $H$ of an electron in a crystal in presence of an electric field, that is

$$ H = H_0 + e \mathbf{x} \cdot \mathbf{E} = H_0 + \mathbf{d} \cdot \mathbf{E} , \quad (8) $$

where $H_0$ is the Hamiltonian of a crystal in its ground state, $e$ the electronic charge, $\mathbf{x}$ the position operator, $\mathbf{E}$ the electric field and $\mathbf{d} = e \mathbf{x}$ the dipole operator. We further make the hypothesis that the electric field can be added as a perturbation, so that the eigenvectors $|\psi_{kb}\rangle$ of $H$ are approximately the eigenvectors of $H_0$ as well ($\mathbf{k}$ is the wavevector and $b$ the band index are Bloch quantum numbers). The eigenvalues of $H_0$ are denoted as $\epsilon_b(\mathbf{k})$. We also stress that the single-particle approximation is consistent with the numerical implementation using parameters computed from density-functional theory.

Before proceeding, it is important to choose a wavefunction gauge such that the derivative $\frac{\partial |\psi_{kb}\rangle}{\partial \mathbf{R}}$ exists and is continuous. To this aim, we recall the maximally localized Wannier functions, defined as:

$$ |Rn\rangle = \frac{V}{(2\pi)^3} \int dke^{-i\mathbf{k} \cdot \mathbf{R}} \sum_b U_{k,bn} |\psi_{kb}\rangle = \frac{V}{(2\pi)^3} \int dke^{-i\mathbf{k} \cdot \mathbf{R}} |\tilde{\psi}_{kn}\rangle , \quad (9) $$

where $\mathbf{R}$ labels a Bravais lattice site and $U_{k,bn}$ is a matrix fixing the wavefunction gauge. The matrix $U_{k,bn}$ is chosen as the one that maximally localizes Wannier functions \[38\], which has also the benefit of making $|\tilde{\psi}_{kn}\rangle$ a smooth wavefunction across different wavevectors (otherwise, wavefunctions at different wavevectors assume random phases).

Next, we briefly recall that the definition of the Wigner transform. Given an operator in the real-space (position) representation $A(x, x')$, we can transform it to the phase-space representation through the Wigner transform $W[A]$, defined as

$$ W[A]_{nn'}(x, k) = \int dx' e^{-i2\mathbf{k} \cdot \mathbf{x}'} A_{nn'}(x + x', x - x') . \quad (10) $$

Similarly, if we start from an operator in the momentum representation, we can transform it into the phase-space representation as

$$ W[A]_{nn'}(x, k) = \int dk' e^{i2\mathbf{k} \cdot \mathbf{x}'} A_{nn'}(k + k', k - k') . \quad (11) $$

We now want to describe the equation of motion of the system under the Hamiltonian of Eq. \[9\]. We can build the density matrix of the system using the basis of wavefunctions in the Wannier gauge $|\tilde{\psi}_{kn}\rangle$ introduced above. We can thus represent the state of the system using the single-particle density matrix operator $\rho$, whose matrix elements are $\rho_{nn'}(k, k'; t) = \text{Tr} \{ \hat{\rho}(t) \hat{c}_{k,n}^{\dagger} \hat{c}_{k',n} \}$, where $\hat{c}_{k,n}$ and $\hat{c}_{k,n}$ are creation/annihilation operators of an electronic state $|\tilde{\psi}_{kn}\rangle$. As described in Ref. \[18\], one may use the equation of motion of the density matrix, and Wigner transform it to derive an equation of motion for the system. Here, we derive a simplified equation of motion for the state of the system using the Wigner function of the crystal \[15\], which is defined through the Wigner transform of the density matrix as

$$ W_{nn'}(x, k, t) = \sum_{\Delta k} e^{i2\Delta k \cdot x} \rho_{nn'}(k + \Delta k, k - \Delta k; t) , \quad (12) $$
where we used the rotation of coordinates \( k, k' \to \frac{k + k'}{2}, k' = k \).

Note that that the position \( x \) appearing in the Wigner transform is, to be precise, a Bravais lattice vector and thus a discrete variable. However, when studying transport properties, we are only interested in the macroscopic behavior of the system. Under this macroscopic limit, we only study the changes of \( x \) on a length-scale much larger than the lattice parameter, so that \( x \) can be approximated as a continuum variable. As a result, \( W \) admits a continuous derivative with respect to \( x \) and, thanks to the gauge choice on the wavefunction \( W \) is also differentiable with respect to \( k \).

As demonstrated by Moyal [16][17], the Wigner function obeys the following equation of motion:

\[
\frac{\partial W_{nn'}(x, k, t)}{\partial t} = -\{ \{ W(x, k, t), H(x, k) \} \}_{nn'} = \frac{i}{\hbar} \left( W(x, k, t) \star H(x, k) - H(x, k) \star W(x, k, t) \right)_{nn'},
\]

where \( \{ \{ f, g \} \} \) indicates the Moyal bracket between two operators \( f \) and \( g \), and we defined the Moyal product \( \star \) as:

\[
f \star g = f(x, k) \exp \left( \frac{i}{2} \left( \frac{\partial}{\partial x} \cdot \frac{\partial}{\partial k} - \frac{\partial}{\partial k} \cdot \frac{\partial}{\partial x} \right) \right) g(x, k),
\]

where the left/right arrow indicates that the derivative operator acts on the operator to the left/right.

The equation of motion for \( W \) is the phase-space analogous of the Liouville-Von Neumann equation of motion for the density matrix, and it therefore has a complexity comparable to that of the Schroedinger’s equation.

Now, we can further simplify this equation of motion by making the hypothesis that both \( H \) and \( W \) are slowly varying functions of \( x \) and \( k \). We then expand the exponential appearing in the Moyal product in Taylor series and approximate the equation of motion as:

\[
\frac{\partial W_{nn'}(x, k, t)}{\partial t} \approx i \left( W(x, k, t)H(x, k) - H(x, k)W(x, k, t) \right)_{nn'} - \frac{1}{2} \left( W(x, k, t) \left( \frac{\partial}{\partial x} \cdot \frac{\partial}{\partial k} - \frac{\partial}{\partial k} \cdot \frac{\partial}{\partial x} \right) H(x, k) - H(x, k) \left( \frac{\partial}{\partial x} \cdot \frac{\partial}{\partial k} - \frac{\partial}{\partial k} \cdot \frac{\partial}{\partial x} \right) W(x, k, t) \right)_{nn'}
\]

\[
= -i \left[ H(x, k), W(x, k, t) \right]_{nn'} - \frac{1}{2} \left\{ \frac{\partial H(x, k)}{\partial k}, \frac{\partial W(x, k, t)}{\partial x} \right\}_{nn'} + \frac{1}{2} \left\{ \frac{\partial H(x, k)}{\partial x}, \frac{\partial W(x, k, t)}{\partial k} \right\}_{nn'}.
\]

Note that, if \( H \) and \( W \) commute (for example, if the two are diagonal in the band index \( n \)), this equation reduces to the Poisson bracket, i.e. the time evolution of a classical Hamiltonian.

The equation of motion is almost in the final form reported in the main text. However, it is still expressed in terms of the basis set \( \{ \psi_{bn} \} \). While convenient for the derivation, it is more practical to work with an equation in terms of the Bloch index \( b \), rather than the Wannier index \( n \) (the Wannier function basis set doesn’t in general diagonalize the Bloch Hamiltonian). Therefore, we rotate results in the \( |\psi_{bn} \rangle \) basis set and write \( W \) and \( H \) as:

\[
H_{bb'}(x, k) = \sum_{nn'} U_{bn}^\dagger(k) H_{nn'}(x, k) U_{bn'}(k),
\]

and

\[
f_{bb'}(x, k) = \sum_{nn'} t_{bn}^\dagger(k) W_{nn'}(x, k) U_{bn'}(k).
\]
The equation of motion can thus be written as:

\[
\frac{\partial f_{bb'}(x, k, t)}{\partial t} = -i \left[ H(x, k), f(x, k, t) \right]_{bb'} - \frac{1}{2} \left\{ \frac{\partial H(x, k)}{\partial k}, \frac{\partial f(x, k, t)}{\partial x} \right\}_{bb'} + \frac{1}{2} \left\{ \frac{\partial f(x, k, t)}{\partial x}, \frac{\partial H(x, k)}{\partial k} \right\}_{bb'}. \tag{19}
\]

We now want to manipulate the matrix elements of the Hamiltonian entering the equation of motion for the Wigner function. First, we note that the Wigner transform of the Hamiltonian at Eq. 8 is Eq. 8 itself, because \( H_0 \) (a Bloch Hamiltonian) is diagonal in \( k \) and the coupling with the electric-field is diagonal in \( x \).

\[
\langle \psi_{bb'} | H(x, k) | \psi_{bb'} \rangle = \epsilon_b(k) \delta_{bb'} + d_{bb'}(k) \cdot E = [\mathcal{E}(k) + D(k) \cdot E]_{bb'}, \tag{20}
\]

where we introduced two matrices \( \mathcal{E}(k) \) and \( D(k) \) containing the single-particle energies \( \epsilon_b(k) \) and dipoles \( d_{bb'}(k) \). The dipole operator requires some care, since the position operator is not well-defined in a periodic system. The off-diagonal terms \( b \neq b' \) satisfy:

\[
d_{bb'} = \langle \psi_{bb'} | e r | \psi_{bb'} \rangle = \epsilon_b(k) \delta_{bb'} + d_{bb'}(k) \cdot E = \frac{\epsilon_b(k)}{\epsilon_b(k) - \epsilon_{b'}(k)} v_{bb'}(k), \quad \text{for } b \neq b', \tag{21}
\]

where \( v_{bb'}(k) \) is the velocity operator. The diagonal terms \( d_{bb}(k) \) are ill-defined \[39\]. Luckily, these terms appear only inside a commutator, so that the diagonal terms don’t contribute. We thus set \( d_{bb}(k) = 0 \) without altering results.

The derivatives of the Hamiltonian are readily computed as

\[
\langle \psi_{bb'} | \frac{\partial H(x, k)}{\partial x} | \psi_{bb'} \rangle = eE \delta_{bb'}, \tag{22}
\]

and

\[
\langle \psi_{bb} | \frac{\partial H(x, k)}{\partial k} | \psi_{bb'} \rangle = v_{bb'}(k). \tag{23}
\]

Combining all this terms together, the equation of motion for the Wigner function \( f_{bb'}(x, k, t) \) is:

\[
\frac{\partial f_{bb'}(x, k, t)}{\partial t} + i \left[ \mathcal{E}(k) + D(k) \cdot E, f(x, k, t) \right]_{bb'} + \frac{1}{2} \left\{ v(k), \frac{\partial f(x, k, t)}{\partial x} \right\}_{bb'} - eE \cdot \frac{\partial f_{bb'}(x, k, t)}{\partial k} = 0. \tag{24}
\]

This is the equation of motion for the Hamiltonian \( H_0 + e x \cdot E \), which, however, doesn’t take into account for the effect of electronic collisions, in particular electron-phonon scattering. This effect is added as a perturbation, and we define the electron-phonon collision matrix as \[18 \ 19 \ 20 \ 21 \ 22\]:

\[
\frac{\partial f_{bb'}(x, k, t)}{\partial t} \bigg|_{coll} = (1 - \delta_{bb'}) \frac{\Gamma_b(k)}{2} f_{bb'}(x, k, t) + \delta_{bb'} \frac{1}{V} \sum_{k' \nu} A_{kk',b} f_{bb'}(x, k', t). \tag{25}
\]

Here, the diagonal terms of \( f \) are modified by the scattering matrix \( A \), which is built as the electron-phonon collision matrix of the Boltzmann transport equation. The off-diagonal terms instead are built \[10 \ 18 \ 41\], from the electron-phonon linewidths \( \Gamma_b(k) = A_{kk,b} \). The electron-phonon scattering
matrix is computed as [37]:

\[
A_{kk',bb'} = \delta_{kk'} \delta_{bb'} \frac{2\pi}{N_q} \sum_{m,vq} |g_{mmv}(k,q)|^2 \left[ \left( 1 - \tilde{f}_m(k+q) + \bar{n}_\nu(q) \right) \delta(\epsilon_n(k) - \epsilon_m(k+q) - \omega_\nu(q)) \right. \\
+ \left( \tilde{f}_m(k+q) + \bar{n}_\nu(q) \right) \delta(\epsilon_n(k) - \epsilon_m(k+q) + \omega_\nu(q)) \\
+ \frac{2\pi}{N_q} \sum_{m,vq} |g_{mmv}(k,q)|^2 \left[ \tilde{f}_n(k)(1 - \tilde{f}_m(k+q)) \bar{n}_\nu(q) \delta(\epsilon_n(k) - \epsilon_m(k+q) - \omega_\nu(q)) \right. \\
+ \left. \tilde{f}_m(k+q)(1 - \tilde{f}_n(k)) \bar{n}_\nu(q) \delta(\epsilon_n(k) - \epsilon_m(k+q) + \omega_\nu(q)) \right],
\]

where \( \omega_\nu(q) \) is the phonon frequency at wavevector \( q \) and branch index \( \nu \), \( \bar{n}_\nu(q) \) is the Bose–Einstein distribution function, and \( |g_{mmv}(k,q)|^2 \) is the strength of the electron-phonon interaction. All these quantities can be computed using density-functional (perturbation) theory.

Finally, the WTE becomes

\[
\frac{\partial f_{bb'}(x,k,t)}{\partial t} + i \left[ E(k) + D(k) \cdot E, f(x,k,t) \right]_{bb'} + \frac{1}{2} \left\{ v(k), \frac{\partial f(x,k,t)}{\partial x} \right\}_{bb'} - eE \cdot \frac{\partial f_{bb'}(x,k,t)}{\partial k} = \frac{\partial f_{bb'}(x,k,t)}{\partial t}_{\text{coll}},
\]

where \( f \) is the Fermi–Dirac distribution, and \( f^E \) is the unknown quantity to be found from the WTE. Note that \( f^E \) is a vector to be found for every direction of the electric field.

We split the solution of the WTE in two parts, diagonal \((b = b')\) and off-diagonal \((b \neq b')\) contributions. The diagonal components of the WTE are

\[
eE \cdot \frac{\partial f_{bb}(k)}{\partial k} = \sum_{k',v'} A_{kk',bb'} f_{v'v'}(k'),
\]

which is equivalent to the BTE problem. The equation can thus be solved using standard techniques developed for the BTE (see e.g. refs. [12]). We verified that exact solutions of the BTE don’t modify results significantly, and therefore we adopt the relaxation time approximation, and approximate the diagonal terms as

\[
eE \cdot \frac{\partial f_{bb}(k)}{\partial k} = \Gamma_b(k) f_{bb}(k).
\]

Using the linearized expression for the Wigner function (and neglecting terms quadratic in \( E \)), the equation is readily solved by

\[
f^{E,bb}_{bb}(k) = \frac{e}{\Gamma_b(k)} \frac{\partial f_b(k)}{\partial k}.
\]
The off-diagonal terms evolve according to

$$i \hbar \left[ \mathcal{E}(k) + D(k) \cdot E, f(k) \right]_{bb'} - e E \cdot \frac{\partial f_{bb'}(k)}{\partial k} = -\frac{\Gamma_b(k) + \Gamma_{b'}(k)}{2} f_{bb'}(k). \quad (32)$$

Using the linearized expression, we obtain:

$$i \left( \epsilon_b(k) - \epsilon_{b'}(k) \right) f_{bb'}^E(k) + i \left( \epsilon_{b'}(k) - \epsilon_b(k) \right) d_{bb'}(k) = -\frac{\Gamma_b(k) + \Gamma_{b'}(k)}{2} f_{bb'}^E(k). \quad (33)$$

The equation is readily solved finding

$$f_{bb'}^E(k) = \frac{\bar{f}_b(k) - \bar{f}_{b'}(k)}{\epsilon_b(k) - \epsilon_{b'}(k)} \frac{2e v_{bb'}(k)}{2i \left( \epsilon_b(k) - \epsilon_{b'}(k) \right) + \left( \Gamma_b(k) + \Gamma_{b'}(k) \right)}.$$

Similarly, we can solve the WTE for the response to a thermal gradient, similarly to what discussed in Ref. [18]. We can now set the electric field to zero and linearize the Wigner function as:

$$f_{bb'}(k) = \bar{f}_b(k) \delta_{bb'} + f_{bb'}^T(k) \cdot \nabla T. \quad (35)$$

The diagonal components of the WTE are then

$$v(k) \frac{\partial \bar{f}_b(k)}{\partial T} = -\sum_{k',b'} A_{kk',b} f_{bb'}^T(k'), \quad (36)$$

which can be solved in the relaxation time approximation as discussed above for the electrical conductivity. The off-diagonal components are given by

$$i \left( \epsilon_b(k) - \epsilon_{b'}(k) \right) f_{bb'}^T(k) + \frac{1}{2} \left( \frac{\partial \bar{f}_b(k)}{\partial T} + \frac{\partial \bar{f}_{b'}(k)}{\partial T} \right) v_{bb'}(k) = -\frac{\Gamma_b(k) + \Gamma_{b'}(k)}{2} f_{bb'}^T(k), \quad (37)$$

which can again be solved trivially in terms of $f_{bb'}^T(k)$.

### 1.3 Transport coefficients

Having computed the Wigner function from the WTE, the expectation value of an operator $A$ can be computed in the phase-space representation as

$$\langle A(t) \rangle = \frac{g_s}{V N_k} \sum_{kk',b} \int f_{bb'}(x, k, t) A_{bb'}(k) dx,$$

where the factor $g_s = 2$ takes into account for the spin-degeneracy (we are only considering non-magnetic systems), and $N_k$ is a normalization for the number of wavevectors. We can apply this formula to estimate the current of a steady-state homogeneous system, i.e. when $f_{bb'}(x, k, t)$ doesn’t depend on space and time. The charge current $j$ can be computed as:

$$j = \frac{e g_s}{V N_k} \sum_{kk'} f_{bb'}(k) v_{bb'}(k) = \frac{e g_s}{2VN_k} \sum_{kk'} \left\{ v(k), f(k) \right\}_{bb'}, \quad (39)$$

where we used an anticommutator to symmetrize results. The electrical conductivity easily follows, since $j = \sigma E$. 

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More precisely, we can define the Onsager transport coefficients as:

\[ J = L_{EE} E + L_{ET} \nabla T, \]
\[ Q = L_{TE} E + L_{TT} \nabla T, \]  
(40)  
(41)

where \( Q \) is the heat flux, and the response coefficients can be computed as:

\[ L_{EE}^{ij} = \frac{e g_s}{V N_k} \sum_{kb} \frac{1}{2} \left\{ v^i(k), f^E_j(k) \right\}_{bb}, \]
\[ L_{ET}^{ij} = \frac{e g_s}{V N_k} \sum_{kb} \frac{1}{2} \left\{ v^i(k), f^T_j(k) \right\}_{bb}, \]
\[ L_{TE}^{ij} = \frac{g_s}{V N_k} \sum_{kb} (\epsilon_b(k) - \mu) \frac{1}{2} \left\{ v^i(k), f^E_j(k) \right\}_{bb}, \]
\[ L_{TT}^{ij} = \frac{g_s}{V N_k} \sum_{kb} (\epsilon_b(k) - \mu) \frac{1}{2} \left\{ v^i(k), f^T_j(k) \right\}_{bb}. \]  
(42)  
(43)  
(44)  
(45)

As customary in electronic transport theory, we recognize the electrical conductivity as \( \sigma = L_{EE} \) and the Seebeck coefficient as \( S = -L_{EE}^{-1} L_{ET}. \)

The expression for the electrical conductivity is readily computed. In fact, we can write the electrical conductivity as

\[ \sigma^{ij} = \sigma^{BTE,ij} + \Delta \sigma^{ij}, \]  
(46)

where the first term takes into account for the diagonal (BTE-like) components of the WTE and the second one for the off-diagonal ones.

Substituting the solution of the WTE in the definition of electrical conductivity, we find that the off-diagonal contribution is

\[ \Delta \sigma^{ij} = \frac{e g_s}{2 V N_k} \sum_{kbb', b \neq b'} (v^i_{bb'} (k) f^E_{jbb'} (k) + f^E_{jbb'} (k) v^i_{bb'} (k)) \]
\[ = 2 g_s e^2 \frac{V N_k}{kkbb', b \neq b'} v^i_{bb'} (k) v^j_{bb'} (k) \frac{\bar{f}_b (k) - \bar{f}_b (k)}{\epsilon_b (k) - \epsilon_b (k)} \frac{\Gamma_b (k) + \Gamma_b (k)}{4 (\epsilon_b (k) - \epsilon_b (k))^2 + (\Gamma_b (k) + \Gamma_b (k))^2}. \]  
(47)  
(48)

Note that \( \Delta \sigma^{ij} \) is a positive quantity and therefore always increases the estimate of conductivity with respect to the BTE (since \( \Gamma_{k,b} > 0 \) and \( \bar{f} \) is a decreasing monotonic function of \( \epsilon \)). The diagonal contribution to the electrical conductivity is readily computed within the relaxation time approximation, and it can be shown to be

\[ \sigma^{BTE,ij} = \frac{g_s}{V N_k} \sum_{kb} v^i_{bb} (k) f^E_{jbb} (k) \approx \frac{g_s e^2}{V N_k} \sum_{kb} \frac{\partial \bar{f}_b (k)}{\partial \epsilon} v^i_{bb} (k) v^j_{bb} (k) \frac{1}{\Gamma_b (k)}. \]  
(49)

### 1.4 Methods

We use density functional theory as implemented in the plane-wave software suite Quantum-ESPRESSO \[27, 28\]. To compute the ground state, we use ultrasoft pseudopotentials from the GBRV library \[43\], with the PBEsol functional. We use an energy cutoff of 80 Ry, and integrate the Brillouin zone with a \( 8 \times 8 \times 8 \) mesh of k-points. We build the trigonal unit cell using experimental
Figure 4: Computational estimates of the electrical conductivity (panel a) and Seebeck coefficient (panel b) of Bi$_2$Se$_3$ as a function of temperature, for different values of hole doping concentration. Solid lines are estimated using the Wigner transport equation, while dotted lines are semiclassical estimates obtained solving the Boltzmann transport equation. The hole doping case is qualitatively symmetric to the electron doping one.
estimates of the crystal structure [44], i.e. with a lattice parameter of 9.839 Å, and an angle $\alpha$ such that $\cos \alpha = 0.91068$. The Wannier functions are computed using $p$ orbitals on both Bi and Se atoms as initial guesses.

Phonon properties, and the electron-phonon matrix elements are computed with density-functional perturbation theory [29] on a coarse grid of $4 \times 4 \times 4$ q-points. Electron-phonon matrix elements are subsequently interpolated on a finer grid using a multi-dimensional linear interpolation [12], while electronic energies and velocities are interpolated using Wannier90 [30]. Transport properties have been implemented in an in-house software. The Dirac-delta ensuring energy conservation during an electron-phonon scattering event has been approximated using an adaptive-smearing scheme [45]. Transport properties have been converged with respect to the k-points mesh used to integrate the Brillouin zone.

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