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First-principles calculations of thermal, electrical, and thermoelectric transport properties of semiconductors

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Abstract

Transport properties of semiconductors are keys to the performance of many solid-state devices (transistors, data storage, thermoelectric cooling and power generation devices, etc.). Understanding of the transport details can lead to material designs with better performances. In recent years simulation tools based on first-principles calculations have been greatly improved, being able to obtain the fundamental ground state properties of materials (such as band structure and phonon dispersion) accurately. Accordingly, methods have been developed to calculate the transport properties based on an *ab initio* approach. In this review we focus on the thermal, electrical, and thermoelectric transport properties of semiconductors, which represent the basic transport characteristics of the two degrees of freedom in solids – electronic and lattice degrees of freedom. Starting from the coupled electron-phonon Boltzmann transport equations, we illustrate different scattering mechanisms that change the transport features and review the first-principles approaches that solve the transport equations. We then present the first-principles results on the thermal and electrical transport properties of semiconductors. The discussions are grouped based on different scattering mechanisms including phonon-phonon scattering, phonon scattering by equilibrium electrons, carrier scattering by equilibrium phonons, carrier scattering by polar optical phonons, scatterings due to impurities, alloying and doping, and phonon drag effect. We show how the first-principles methods allow one to investigate the transport properties with unprecedented details and also offer new insights to the electron and phonon transport. Current status of the simulation is mentioned when appropriate and some of the future directions are also discussed.

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1. Introduction

The transport property of solids is an old but enlightening problem. Many insightful ideas emerge from such studies, such as the superconductivity, localization, etc., which in some sense reflect the modern development of the condensed matter physics. It is intriguing also because most of our daily “machines” rely on such transport, either of charge, mass, or of energy. Examples include transistors (mobility), phase change medium for data storage (thermal conductivity), thermoelectric cooling and power generation devices (electrons and phonons), and so on. Better materials are constantly sought after for improving the overall performance of the working device. Improvement of existing and discovery of new materials thus require a clear understanding of the underlying physics of the transport properties. Theoretical models combined with experiments have offered great insights into the different mechanisms that affect the transport properties. However some details have been ignored in order for the theories to be tractable. For example, the simple kinetic formula, \( \sigma = \frac{1}{3} cv\Lambda \), describes the electrical conductivity \( \sigma \) in terms of three averaged parameters of the energy carriers: carrier energy \( c \), velocity \( v \) and mean free path \( \Lambda \). In reality, however, these parameters can vary significantly among different carriers and even the same type of carriers. Such information has been difficult to obtain within traditional methods because experiments normally probe transport properties that are averaged over all the carriers, while the theoretical models are intractable without certain approximations thus lack accuracy and in many cases, predictive power. The development of the first-principles computational techniques in recent decades enables one to study materials’ physical properties accurately without adjustable parameters[1]. The merit of the first-principles methods lies in that they are easily justifiable and useful for a wide range of problems. Since the invention[2,3], they have been widely used for studying and also predicting the physical properties of many materials, such as equilibrium crystal structure, electronic band structure, phonon dispersion, etc[4]. We will briefly describe how the first-principles calculations can be utilized for studying the transport properties. We do not discuss how the first-principles methods work but refer readers to some of the books that already exist and the references therein[4,5].

We want to be cautious that “first-principles” is sometimes a misleading name due to the fact that
the underlying theory (which should be referred to as density functional theory, aka DFT) also involves approximations to the quantum mechanical governing equation - Schrodinger equation (including relativistic corrections derived from Dirac equation). For example, only the ground state properties are calculated and the strongly-correlated systems cannot be well described without the introduction of some tuning parameters[4]. In this light, there are also other quantum mechanics-based algorithms which are more accurate for such problems but are also much more time-consuming[6]. In the following the main results are obtained using DFT, which describes many properties of solids very well and currently is sufficient for the transport phenomenon study in commonly-used semiconductors. However, we will also point out that some other formalisms (to more accurately describe the many-body effect of electrons) will be desired along the path of developing first-principles tools for more complex materials.

In this review we focus on the studies of semiconductors, and in particular, their thermal, electrical and thermoelectric properties. Semiconductors, though only a portion of the family of solids, play important roles in various applications such as microelectronics, photovoltaics, optoelectronics and thermoelectrics. Although extensive studies into semiconductors from theoretical models to experiments have revealed many physical mechanisms for describing their transport behaviors, so far it is hard to predict a material’s transport property (for example, mobility) without performing experiments. A computational technique that has the predictive power will thus greatly facilitate the optimization of existing materials and the search of better ones[7]. The first-principles methods have been traditionally applied to study the electronic structure and phonon dispersion, both of which are eigenstates of the system. In a particle picture, each eigenstate corresponds to one or several particles (electrons or phonons) that carry certain energy with some velocity. To describe the transport property such as mobility and thermal conductivity, one must know how these particles evolve under the influence of other states. This process is formalized by the famous Boltzmann transport equation (BTE) and will be the cornerstone of our following discussions. One of the major challenges of solving BTE is to know how the particles are scattered by each other (or by the defect and boundary). The scattering process is determined by the coupling between different eigenstates. The calculation of such coupling strengths has been computationally prohibitive and only becomes accessible recently
thanks to a series of developments for an efficient extraction of the interaction strengths between different eigenstates (electrons and phonons). In the next section, different types of interaction along with their calculation will be discussed under the BTE formalism.

Among all the transport properties, the thermoelectric transport presents a particular challenge, firstly because it is affected by three transport phenomena as shown by the figure of merit $\zeta T = \sigma S^2 T / \kappa$ which characterizes material’s thermoelectric efficiency, where $\sigma, S, \kappa, T$ are the electrical conductivity, Seebeck coefficient, thermal conductivity and absolute temperature, respectively. For semiconductors, the electrical conduction and Seebeck effect are mostly related to the electronic properties while the thermal conduction must consider the phonons. Therefore for the thermoelectric transport as a whole, one must consider both of electrons and phonons, as well as the coupling between them. Secondly, the thermoelectric materials are often complex due to the alloying and doping procedure involved, which always poses a challenge for first-principles simulations. In this review, we briefly summarize the recent efforts towards a first-principles calculation of the thermal, electrical and thermoelectric properties of semiconductors, particularly group IV and III-V semiconductors. In section 2 we first introduce the formalism of the coupled electron-phonon Boltzmann transport equations, which serve as the starting point of our discussions. Under the BTE formalism, basic concepts are explained and the corresponding computational techniques are discussed. The following two sections are devoted to the first-principles results on the transport property: section 3 deals with the thermal transport and section 4 examines the electrical transport. In the final section we briefly discuss the possible future directions that help to improve the first-principles simulations. We anticipate that the development of the computational tools will further extend the calculations to more complex materials.

2. Coupled Electron-phonon Boltzmann Equations

The electron and phonon properties of a solid are normally described by the band structure and phonon dispersion, respectively. For convenience we would like to think of these eigenstates as particles without considering their phase. These particles carry certain charge or energy dictated
by quantum mechanics. This semiclassical picture is not always valid but can be justified if the variation of the external field occurs in a larger scale than the wave packet associated with this particle and the field is relatively weak compared to the periodically varying atomic potential[8]. A particle in an eigenstate with finite group velocity implies that it keeps moving without being deflected under the dynamics of the system, since the Hamiltonian acting on the eigenstate gives itself. In other words, the eigenstate has an infinite lifetime. Practically, this cannot be achieved because there are always perturbations to the system, which causes finite transition probabilities between different states. In a particle picture, the perturbations lead to the scattering of the particle by other particles or impurities, which give rise to the “resistance” for the charge or energy flow. To describe such process, one must have a way to treat the great amount of particles that are moving inside the material with the interactions that lead to their scatterings.

Rigorously, a (6N+1)-dimensional phase space (3N for positions, 3N for momenta and one for time) is used to describe the dynamics of N particles. Such description is adopted by the molecular dynamics technique, and has been used to calculate phonon properties as well as the lattice heat conduction[9,10]. However, the molecular dynamics method lacks the predictive power due to the empirical interatomic potentials involved, while a first-principles calculation with a large system of N particles is formidable. Alternatively, one could use a single-particle distribution function to represent all the particles in the system. This function has only 7 independent variables (3 for positions, 3 for momenta and one for time) and describes how many particles stay in the state characterized by given position and momentum at each time. The crucial step towards describing the transport of the system is to rewrite the dynamics of N particles into a governing equation for the distribution function. This is achieved by the introduction of the BTEs. For the electron and phonon systems in the steady state, the BTEs read as[8,11–13]

\[
\begin{align*}
\mathbf{v}_a(k) \cdot \nabla_r f_a(k) + \frac{F}{\hbar} \cdot \nabla_k f_a(k) &= \left( \frac{\partial f_a(k)}{\partial t} \right)_{\text{coll}} \\
\mathbf{v}_q(q) \cdot \nabla_q n_q(q) &= \left( \frac{\partial n_q(q)}{\partial t} \right)_{\text{coll}}
\end{align*}
\]

(1)

where \( \hbar \) is the reduced Planck constant, \( \mathbf{F} \) is the external force exerting on this particle (electrostatic field in our case, which affects electrons but not phonons) and the velocity vectors...
\( \mathbf{v} \) for electrons (\( \mathbf{v}_\alpha(k) \)) and for phonons (\( \mathbf{v}_\lambda(q) \)) are specified with wave vectors \( k \) (for electrons) and \( q \) (for phonons), as well as band number \( \alpha \) and branch number \( \lambda \). \( f \) and \( n \) represent the distribution functions for electrons and phonons, respectively, with equilibrium state described by Fermi-Dirac and Bose-Einstein statistics:

\[
\begin{align*}
  f^0_\alpha(k) &= \frac{1}{e^{(E^\alpha_k-\mu)\beta}+1} \\
  n^0_\lambda(q) &= \frac{1}{e^{\hbar^2q^2\beta}-1}
\end{align*}
\]  

(2)

The first equation of Eq. (1) is for electrons and will be denoted as electron-BTE while the second equation is for phonons and will be called phonon-BTE. Here we skip the derivation of Eq. (1), which can be found in many textbooks[8,11–13]. We instead want to pay more attention to the meanings of each term and their explicit forms. Briefly speaking, the BTEs describe the balance of the change of the particle number in various available states. There are two major causes for these changes. One is coming from the external field exerting on the particles, with examples including electric field for electrons (second term of electron-BTE) and temperature gradients for both electrons (included in first term of electron-BTE) and phonons (first term of phonon-BTE). The other originates from the scatterings between the particles. For convenience, the terms of the former type in the Boltzmann equation are often called drift terms while those of the latter type are referred to as collision terms[8].

In equilibrium, the distribution functions for electrons and phonons are described by Fermi-Dirac and Bose-Einstein statistics, respectively. As the macroscopic fields start to drive the system, electrons and phonons will move away from the equilibrium, until they are balanced by the collision terms, which rebuild their distribution functions. The degree of the non-equilibrium is thus determined by the strength of the scatterings. For both electrons and phonons, there are various mechanisms that can make transitions from one state to another, such as phonon-phonon interaction, electron-phonon interaction and electron-electron interaction. Besides, due to the impurities involved (for example, by alloying or doping procedure), we can also have impurity scatterings for both electrons and phonons. In general, the collision terms can be described as
where each term represents the scattering probabilities into and out of certain state, added up together according to the Matthiessen’s rule[8]. We neglect the electron-electron scattering because within most temperature ranges it is hardly as effective as other mechanisms in scattering the electrons[11,14]. However we will mention the necessity of considering the electron-electron interaction into the transport calculations as a step beyond current status.

The central problem to the BTEs is to solve the distribution functions. If we know the distribution function, the transport properties can be easily calculated by summing up all the states with corresponding energies and velocities. For this purpose, we need to explicitly describe the collision terms. These scatterings depend on the available states of electrons and phonons, and typically involve the distribution functions in a non-linear way. At moderate external fields, however, the distribution functions only have small deviations from the equilibrium distribution. Consequently, one can expand the distribution function near the equilibrium condition in the BTEs and take the lowest order terms, which results in the linearized BTEs[11]:

\[
\begin{align*}
&\left(\frac{\partial f_a(k)}{\partial t}\right)_{\text{coll}} = \left(\frac{\partial f_a(k)}{\partial t}\right)_{e-ph} + \left(\frac{\partial f_a(k)}{\partial t}\right)_{e-imp} + \\
&\left(\frac{\partial n^0_\lambda(q)}{\partial t}\right)_{\text{coll}} = \left(\frac{\partial n^0_\lambda(q)}{\partial t}\right)_{ph-ph} + \left(\frac{\partial n^0_\lambda(q)}{\partial t}\right)_{ph-e} + \\
&\begin{pmatrix}
\begin{pmatrix}
\frac{\partial f_a(k)}{\partial t}
&\frac{\partial n^0_\lambda(q)}{\partial t}
\end{pmatrix}

e
&\begin{pmatrix}
\frac{\partial f_a(k)}{\partial t}
&\frac{\partial n^0_\lambda(q)}{\partial t}
\end{pmatrix}
\begin{pmatrix}
\frac{\partial f_a(k)}{\partial t}
&\frac{\partial n^0_\lambda(q)}{\partial t}
\end{pmatrix}
\end{pmatrix}
\end{align*}
\]

where \(T\) is the temperature, \(e\) is the electron charge (negative), \(E\) is the electronic energy level and \(\varphi\) is the electrochemical potential (incorporating electrostatic potential \(\phi\) and chemical potential \(\mu\)). Note that we have replaced the distribution functions in the left-hand-side of Eq. (1) with the equilibrium values. By doing so, the first term of electron-BTE is split into two terms including temperature gradient and chemical potential gradient, respectively, the latter of which, when combined with the electrostatic force term in electron-BTE, leads to the second term in the electron-BTE of Eq. (4), by defining \(\varphi = \phi + \mu / e\) [15].

The collision terms, under the linearized BTE formalism, can be generally expressed as
where the collision terms linearly depend on (with the proportionality factors denoted by $A, B, C, D$) the deviations of the distribution functions from the equilibrium:

$$\begin{align*}
\left( \frac{\partial f_{\alpha} (k)}{\partial t} \right)_{\text{coll}} &= A_{k\alpha} \cdot \Delta f_{\alpha} (k) + \sum_{k' \neq k} A_{k'\alpha} \Delta f_{\alpha} (k') + \sum_{q} B_{q\lambda} \Delta n_{\lambda} (q) \\
\left( \frac{\partial n_{\lambda} (q)}{\partial t} \right)_{\text{coll}} &= C_{q\lambda} \cdot \Delta n_{\lambda} (q) + \sum_{q' \neq q} C_{q'\lambda} \Delta n_{\lambda} (q') + \sum_{k\alpha} D_{k\alpha} \Delta f_{\alpha} (k)
\end{align*}$$

(5)

Note that the first-order deviation in the drift terms and the second-order deviation in the collision terms have been ignored because they are higher order contributions to the BTEs. For quasi-ballistic transport, however, these higher order contributions cannot be neglected because the deviation of the distribution function from the equilibrium is relatively large[15]. In Eq. (5) we have also separated the collision terms into two parts. One is directly proportional to the distribution deviation of the state of interest ($k$ or $q$), and the remaining parts rely on the distribution of other states. The linearized BTE has been the starting point for most of the theoretical work on transport properties. There could be situations where the small deviation of the distribution function is not well justified, which however also poses question to the use of the BTE formalism[8]. Overall the linearized BTE has been shown to be applicable to a wide range of conditions and will be the basis of our following discussion.

A linear system as determined by Eq. (4) generally implies that the distribution function for given state depends on those of other states. To obtain the exact results, numerical techniques that solve large linear systems are usually applied, such as direct or iterative methods. One important but not necessary step here is to neglect the dependence of the collision terms for given state on any other state, which means that the terms in the bracket are ignored in Eq. (5). In other words, this assumes that when considering the scatterings of a given state, all other states appear to be at equilibrium. By doing so, the only term left in the collision terms is (take electrons as an example)
where we replace the proportionality factor with the inverse of a characteristic time (the minus sign is for introducing a properly defined positive time constant as will be clear in the explicit forms). Considering the meaning of the collision, this characteristic time represents how fast the non-equilibrium distribution functions return to the equilibrium due to scatterings. For this reason, they have been called the relaxation times[8]. Using Eq. (7), Eq. (4) can then be readily solved yielding

\[
f_a(k) = f_a^0(k) - \tau_{\kappa \alpha} \left[ \mathbf{v}_a(k) \cdot \frac{\partial f_a^0(k)}{\partial T} \nabla T - e\mathbf{v}_a(k) \cdot \frac{\partial f_a^0(k)}{\partial E} \nabla \varphi \right]
\]  

Such approximation largely simplifies the procedure of solving the BTE and has been called the relaxation time approximation (RTA). However, the crude analysis above has totally removed the coupling between the electron and phonon system, which will be incorrect for certain transport phenomena[11]. We will see that some of the terms inside the bracket in Eq. (5), depending on different situations, must be taken into account to capture the necessary physics with as less computational difficulty as possible. In those cases, the collision term proportional to the given state can still be described using the relaxation time and we regard this also as an example of using the RTA. In essence, the RTA specifically considers the relaxation of the non-equilibrium distribution of a given state, or its decay into other available states. In the particle picture, it can be viewed as the averaged time between the two scattering events of a particle at given state. Its product with the particle’s velocity \( v \) gives rise to the concept of mean free path (MFP) \( \Lambda = v \tau \), the averaged distance the particle travels before being scattered. In the following we discuss three major scattering mechanisms that contribute to the collision terms and the corresponding numerical framework that solves them.

### 2.1 Phonon-phonon interaction

A phonon mode describes a collective atomic movement where each atom vibrates according to certain displacement pattern. Each phonon mode is associated with a wave vector and also
frequency, describing the vibration as a wave propagating in the medium. This classification assumes that the change of the total potential energy with respect to the displacements of the atoms can be written in a homogeneous quadratic form [8]

$$H_{pot} = \frac{1}{2} \sum_{\alpha \beta} D_{\alpha \beta} \left( R, R' \right) u_{\alpha} \left( R \right) u_{\beta} \left( R' \right)$$ (9)

where $R \left( R' \right)$ is the equilibrium position of each atom in the crystal, $u$ is the atomic displacement, $\alpha \left( \beta \right)$ represents the three directions in the Cartesian coordinate, and $D$ is the dynamical matrix. We note that the first order derivative of the potential energy with respect to the atomic displacement is the force. The derivatives of the force with respect to the displacement are usually called force constants, which are natural generalizations of spring constants from mechanics. The $n^{th}$ order derivatives of the energy ($n-1^{th}$ order derivative of force) are denoted as $n^{th}$-order force constants. The form assumed by Eq. (9) therefore implies that all the force constants vanish except for the 2$^{nd}$-order force constants (essentially dynamical matrix). This is the harmonic crystal approximation to real solids [8], which justifies the concepts of phonons. It is valid for studying heat capacity since higher-order effects are often negligible when the atomic displacements are small [8]. However, as we have mentioned, if the eigenstates are the true states of the system, a particle in a state with finite group velocity will keep moving without being deflected, implying an infinite phonon thermal conductivity. It is the higher-order effect that limits the intrinsic thermal conduction [8,11]. For this reason, we usually call the 2$^{nd}$-order force constants the harmonic force constants, while referring to the 3$^{rd}$-order force constants and above as anharmonic force constants.

We can understand the anharmonic force constants that lead to thermal resistance as follows. Since phonon modes are only derived from the harmonic force constants, they are not the exact eigenstates of the system when anharmonic force constants exist. However, one can still use these phonons to describe the dynamics of the system by introducing the transition probabilities between different modes. Classically, it can be understood by a model where spring constants can vary with the displacements, and as a result the original eigenmodes will convert to other modes due to the non-linear vibration dynamics. Quantum mechanically, this
is because the higher-order effects introduce the perturbations to the original system, which manifest as the coupling between the eigenmodes that lead to the transition processes (time dependent perturbation theory)[11].

Among the interactions between phonons, the dominant scattering comes from the three-phonon process[11]. There are mainly two different types of them. In one case two phonons merge into one phonon, creating a new particle with higher energy. In another case one phonon transfers its energy into two phonons. Note that in the phonon-phonon scattering process the phonon number is not conserved. There are also higher-order processes such as four phonon scattering. These typically rarely happen and in most situations can be safely ignored.

For a given phonon state, there are several ways to change its distribution function. This state can decay into two phonons, or can be combined with another phonon and merge into a new phonon. These describe the out-scattering terms. Besides, two other phonons can also scatter and merge into this phonon, resulting in an increase of its distribution function, which are called in-scattering terms. Considering all these processes, one can use the Golden Rule to calculate the transition rates of a given state (corresponding to the proportionality factor $C_{q,\delta}$ in Eq. (5)) due to three-phonon scatterings based on the three phonon interaction matrix elements under the linearized BTE formalism[11,16,17]

$$
\frac{1}{\tau_{q,\delta}^{ph-ph}} = \frac{\pi}{\hbar^2 N_q q_{1,\delta}, q_{2,\delta}} \left| V_{q,\delta, q_{1,\delta}, q_{2,\delta}} \right|^2 \left\{ (1 + n_{q_{1,\delta}}^0 + n_{q_{2,\delta}}^0) \delta(\omega_{q,\delta} - \omega_{q_{1,\delta}} - \omega_{q_{2,\delta}}) \delta_{q_{1,\delta} - q_{2,\delta}, G} 
+ (n_{q_{1,\delta}}^0 - n_{q_{2,\delta}}^0) \delta(\omega_{q_{1,\delta}} + \omega_{q_{2,\delta}} - \omega_{q_{1,\delta}}) \delta_{q_{1,\delta} + q_{2,\delta}, G} 
- \delta(\omega_{q_{1,\delta}} + \omega_{q_{1,\delta}} - \omega_{q_{2,\delta}}) \delta_{q_{1,\delta} + q_{2,\delta}, G} \right\} \right\} \right\} \right\} \right\} \right\} \right\} \right\}
$$

where the three phonon interaction matrix is explicitly related to the anharmonic force constants via (in the following we only consider up to the 3rd-order, or cubic, force constant):

$$
V_{q,\delta, q_{1,\delta}, q_{2,\delta}} = \left( \frac{\hbar}{2} \right)^{3/2} \sum_{R, r_1, r_2} \Psi_{0x, r_1, r_2}^{\alpha, \beta, \gamma} \times \left[ \frac{e^{i(q_{1,\delta} + q_{2,\delta})}}{M_{1, r_1, r_2} \omega_{q_{1,\delta}} \omega_{q_{2,\delta}}} \right]^{1/2} \right\} \right\} \right\} \right\} \right\} \right\} \right\}
$$

In the above expressions, $N_q$ is the number of the discrete points in the reciprocal space mesh ($\mathbf{q}_1$ and $\mathbf{q}_2$ go over the same mesh), $n_{q,\delta}^0$ is the Bose-Einstein distribution of
phonons (Eq. (2)). \( \omega_{\mathbf{q}, \mathbf{\tau}} \) is the phonon frequency, \( e_{\mathbf{q}, \mathbf{\tau}} \) describes the displacement pattern for the phonon eigenmodes and \( M_{\mathbf{\tau}} \) is the atomic mass on the sublattice site \( \mathbf{\tau} \). The cubic force constant \( \Psi^{eff}_{0\mathbf{\tau}, \mathbf{R}_1\mathbf{\tau}_1, \mathbf{R}_2\mathbf{\tau}_2} \) is defined as the third order derivative of the total energy with respect to the displacements of three atomic sites \( (0\mathbf{\tau}, \mathbf{R}_1\mathbf{\tau}_1, \mathbf{R}_2\mathbf{\tau}_2) \) along the corresponding directions \( (\alpha, \beta, \gamma) \), for which \( (\mathbf{R}\mathbf{\tau}) \) represents the atom at the sublattice site \( \mathbf{\tau} \) in the unit cell \( \mathbf{R} \) away from the origin and \( \alpha_i \) are in Cartesian coordinates. For phonons, we have intentionally ignored the non-equilibrium distribution of other states when evaluating the scattering of a given state (see Eq. (5)). As a result, the collision term due to phonon-phonon interaction in Eq. (3) can be written as

\[
\left( \frac{\partial \mathcal{H}_\mathbf{\alpha}(\mathbf{k})}{\partial \mathbf{t}} \right)_{\text{ph-ph}} = -\frac{\Delta n_{\mathbf{q}}(\mathbf{q})}{\tau_{\text{ph-ph}}} \tag{12}
\]

with the relaxation times given by Eq. (10). During these scatterings, energy and crystal momentum conservation need to be satisfied, which impose the conditions on which processes are allowed, as shown by the delta functions in Eq. (10). The crystal momentum conservation requires that the three phonon wave vectors involved in the scattering add up to zero or up to an integer number of reciprocal lattice vectors. Phonon wave vectors are usually defined in the first Brillouin zone (FBZ). As shown by Fig. 1, when two of the phonon wave vectors are small the third wave vector is just their summation and the total crystal momentum is not changed. This is called normal process. On the other hand, when two of the phonon wave vectors are large, their summation will fall outside the FBZ and reciprocal lattice vectors have to be involved to bring it back. Such case is called Umklapp process and causes the change of total crystal momentum. It is these Umklapp processes that scatter phonons back to the equilibrium (reduce the net crystal momentum when they are set up by the driving force) and essentially create the thermal resistance[18].
In fact the phonon-phonon interaction is the main contribution to the thermal resistance for the semiconductors in most temperature ranges. Using RTA, we can write the phonon collision term as

$$
\left( \frac{\partial n_{ij}(\mathbf{q})}{\partial t} \right)_{\text{coll}} = -\frac{\Delta n_{ij}(\mathbf{q})}{\tau_{q^2}}
$$

(13)

where the relaxation time includes phonon-phonon scatterings and also other terms due to impurities or electrons when necessary. Combined with Eq. (1), the phonon distribution function can be readily solved:

$$
n_{ij}(\mathbf{q}) = n_{ij}^0(\mathbf{q}) - \tau_{q^2} \dot{V}_{q^2}(\mathbf{q}) \cdot \frac{\partial n_{ij}^0(\mathbf{q})}{\partial T} \nabla T
$$

(14)

The thermal conductivity as the ratio of the heat current and the temperature gradient is then obtained

$$
\kappa_{ph} = \frac{1}{3\Omega N_q} \sum_{\mathbf{q}_k} \abs{\mathbf{V}_{\mathbf{q}_k}}^2 \tau_{\mathbf{q}_k} \hbar \omega_{\mathbf{q}_k} \frac{\partial n_{\mathbf{q}_k}}{\partial T}
$$

(15)

where $\Omega$ is the unit cell volume and $N_q$ is the number of the discrete points in the reciprocal space. It is clear from the discussion above that the key ingredients for describing the phonon-phonon interaction are the cubic force constants in Eq. (11). Harmonic force constants involve two independent atomic variables and are routinely calculated to obtain the phonon dispersion by the use of the density functional perturbation theory (DFPT)[19]. Anharmonic force constants, however, involve three independent atomic variables and are much harder to obtain due to their larger number involved. There are mainly two ways to
obtain the anharmonic force constants. One is based on a real space approach and the other
starts from the reciprocal space. The real space approach starts by creating many different
atomic configurations in a supercell (one that includes many primitive cells) where one or
more atoms are displaced from the equilibrium positions along certain directions[20]. The
first-principles results for the forces on all the atoms are recorded. These forces can be written
in an expansion expression in terms of the force constants and the displacements (here only
up to third order contributions are explicitly shown):

\[
F^\alpha_{R\tau} = -\sum_{R_i\tau_1} \Phi^{\alpha,\beta}_{R\tau, R_i\tau_1} u^\beta_{R_i\tau_1} - \frac{1}{2} \sum_{R_i\tau_1, R_j\tau_2} \Psi^{\alpha,\beta,\gamma}_{R\tau, R_i\tau_1, R_j\tau_2} u^\beta_{R_i\tau_1} u^\gamma_{R_j\tau_2}
\]

(16)

where \( \Phi^{\alpha,\beta}_{R\tau, R_i\tau_1} \) is the harmonic force constant and \( \Psi^{\alpha,\beta,\gamma}_{R\tau, R_i\tau_1, R_j\tau_2} \) is the cubic force constant,
\( F^\alpha_{R\tau} \) describes the force along \( \alpha \) direction exerted on the atom at \( (R\tau) \) and \( u^\alpha_{R\tau} \) is its
corresponding displacement. The different real space approaches differ in that how the force
constants are fitted from the above equation. It has been shown by Esfarjani et al [20] that
using the symmetry (permutation, translation, rotation and Huang symmetry) the number of
force constants that are required to fit can be largely reduced. They have shown the extraction
of force constants up to 4\(^{th}\)-order that compare reasonably well with the exact results when an
artificial interaction potential is used. Other methods such as those based on compressive
sensing or slave mode expansion are also employed to recognize the most important
anharmonic terms so as to further reduce the number of anharmonic force constants that one
has to deal with and simplify the analysis [21–23]. The advantage of the real space approach
is that one can treat the DFT package as a black box and only produce his own
post-processing code, thus reducing the programming complexity. However, it has been
known from the phonon dispersion calculation that, the real space approach is not as accurate
as the reciprocal space method (DFPT) at a given wave vector \( q \) because the former one
introduces errors when truncating the force constants beyond the size of the supercell while
the latter rigorously treats the infinite size of the crystal[19]. The power of DFPT relies on the
“2n+1” theorem[19,24], which states that in order to know the 2n+1\(^{th}\)-order derivatives of the
total energy only the information of the n\(^{th}\)-order derivatives of the wavefunctions is required.
For phonon dispersion calculation, the first order derivatives of the wavefunctions are already obtained, which can be used to calculate up to 3rd-order force constants. This formalism directly calculates the phonon modes at given \( \mathbf{q} \) and do not rely on supercells. However, its programming complexity is greater than the real space method, and up to now the calculations that used this approach are only restricted to single-element materials such as Si, diamond and graphene[17,25–30]. We want to note that a further development on the reciprocal space method towards more complicated materials can provide an evaluation of the accuracy of the first-principles method as we move from well-known materials to unexplored ones.

2.2 electron-phonon interaction

The BTEs as shown by Eq. (1) will be decoupled without the interaction between the electron and phonon systems[11]. This interaction behaves as a perturbation to each of them from the other system, and cause scatterings for both electrons and phonons. Electronic band structures are derived assuming atoms are at their equilibrium positions. In practice, the atoms vibrate according to the phonon modes and are seldom at equilibrium positions. The phonon modes change the environment (potential energy) the electrons experience and therefore perturb the electron’s movements. In a particle picture, such coupling means that electrons can collide with phonons, making transition from one state to another. These processes need to satisfy the energy and crystal momentum conservation. For the electron-phonon interaction, Born-Oppenheimer approximation is often assumed, which states that as the atoms vibrate, electrons respond so fast that they almost see a static atomic configuration and therefore finds the lowest energy corresponding to that configuration. This simplifies the problem because electrons and phonons still have their own eigenstates, and are only coupled to each other via the coupling matrix element. In an intuitive way, Bloch proposes an analytic form for the electron phonon coupling as follows[31]

\[
\delta V = \sum_{\tau, \mathbf{R}} \mathbf{u}_{\tau, \mathbf{R}} \cdot \frac{\partial V}{\partial \mathbf{u}_{\tau, \mathbf{R}}}
\]  

(17)
where $u$ is the atom displacement and $\frac{\partial V}{\partial u_{r,r'}}$ is the perturbed potential of electrons due to the atomic movements. In this expression we regard $u_{r,r'}$ as operator acting on the phonon eigenstates while $\frac{\partial V}{\partial u_{r,r'}}$ as operator acting on the electron states. A simple explanation of this equation is that, as the atoms in the system move around, the electrons will adjust themselves to find the lowest energy for that atom configuration and therefore the potential energy also changes. It has been shown by Ziman that Eq. (17) can be derived following a more rigorous quantum mechanical treatment\[32\]. The resulting matrix element is

$$\langle n_{q,\ell} \pm 1|u|n_{q,\ell}\rangle \cdot \langle k'\beta|\partial_{q,\ell}V|k\alpha\rangle$$

(18)

where we have transformed the change of the energy due to the displacement of one single atom $\frac{\partial V}{\partial u_{r,r'}}$ to the potential change $\partial_{q,\ell}V$ due to a collective atomic motion corresponding to a phonon mode $(q, \lambda)$. $|k\alpha\rangle$ and $|k'\beta\rangle$ are different electron states that are coupled through the perturbation, and $|n_{q,\ell}\rangle$ represents the phonon state (Note that it is characterized by the number of phonons and the final state can only differ by one). The scatterings can be grouped into two main processes. In one case the electron is scattered into another state by absorbing one phonon and thus increases its energy. Alternatively, the scattering can emit one phonon and the electron energy is lowered. These two processes are distinguished by the first term in Eq. (18) which, depending on if the phonon is absorbed or emitted, gives different proportionality factors to the magnitude of the coupling\[11\].

The second term in Eq. (18), which includes electron states and perturbed potential, is the key element that is required to calculate the scattering rates and thus the transport properties. For convenience, we define the electron-phonon interaction matrix element as

$$g_{q\ell\ell'}(k,k',q) = \left(\frac{\hbar}{2m_\ell\omega_{q,\ell}}\right)^{1/2} \cdot \langle k'\beta|\partial_{q,\ell}V|k\alpha\rangle$$

(19)

which only differs from Eq. (18) by a prefactor inversely proportional to the square root of
the phonon frequency. In Eq. (19) $m_0$ is the electron mass. Under the linearized BTE formalism, it can be shown by using the Golden Rule that the collision terms (Eq. (5)) due to the electron-phonon interaction take the following form[33]

$$\left\{ \frac{\partial F_{\text{ph}}(k)}{\partial t} \right\}_{\text{e-ph}} = \sum_{k'_\alpha q'_\lambda} F_{k\alpha}(k'\beta, q'\lambda) \cdot \Delta f_{k\alpha} + \sum_{k'_\beta q'_\lambda} F_{k'\beta}(k\alpha, q\lambda) \cdot \Delta f_{k'\beta} + \sum_{k'_\lambda q'_\alpha} F_{k'\lambda}(k\alpha', k'\beta) \cdot \Delta n_{k'\lambda}$$

$$\left\{ \frac{\partial n_{\text{ph}}(q)}{\partial t} \right\}_{\text{e-ph}} = \sum_{k\alpha k'\beta} G_{k\alpha}(k'\beta, q\lambda) \cdot \Delta f_{k\alpha} + G_{k'\beta}(k\alpha, q'\lambda) \cdot \Delta f_{k'\beta} - \sum_{k\alpha k'\lambda} G_{k\lambda}(k\alpha', k'\beta) \cdot \Delta n_{k'\lambda}$$

where the coefficients $F$ and $G$ only depend on the equilibrium distribution functions:

$$F_{k\alpha}(k'\beta, q\lambda) = \left[ (n_{k\alpha}^0 + f_{k'\beta}^0) \Pi_+ + (n_{k\alpha}^0 + 1 - f_{k'\beta}^0) \Pi_- \right]$$

$$F_{k'\beta}(k\alpha, q\lambda) = \left[ (n_{k\alpha}^0 + 1 - f_{k'\beta}^0) \Pi_+ + (n_{k\alpha}^0 + f_{k'\beta}^0) \Pi_- \right]$$

$$F_{k'\lambda}(k\alpha', k'\beta) = \left[ (f_{k'\beta}^0 - f_{k'\lambda}^0) \Pi_+ + (f_{k'\beta}^0 - f_{k'\lambda}^0) \Pi_- \right]$$

$$G_{k\alpha}(k'\beta, q\lambda) = \left[ - (n_{k\alpha}^0 + f_{k'\beta}^0) \Pi_+ + (n_{k\alpha}^0 + 1 - f_{k'\beta}^0) \Pi_- \right]$$

$$G_{k'\beta}(k\alpha, q\lambda) = \left[ (n_{k\alpha}^0 + 1 - f_{k'\beta}^0) \Pi_+ - (n_{k\alpha}^0 + f_{k'\beta}^0) \Pi_- \right]$$

$$G_{k'\lambda}(k\alpha', k'\beta) = \left[ (f_{k'\beta}^0 - f_{k'\lambda}^0) \Pi_+ - (f_{k'\beta}^0 - f_{k'\lambda}^0) \Pi_- \right]$$

with $\Pi_+$

$$\Pi_+ = \frac{2\pi}{\hbar} g_{\alpha\beta\lambda}(k, k', q) \cdot \delta \left( E_{k'\beta} - E_{k\alpha} - \hbar \omega_{k\lambda} \right) \cdot \delta_{k-k-k'\lambda \alpha \beta \lambda}$$

$$\Pi_- = \frac{2\pi}{\hbar} g_{\alpha\beta\lambda}(k, k', q) \cdot \delta \left( E_{k'\beta} - E_{k\alpha} + \hbar \omega_{k\lambda} \right) \cdot \delta_{k-k-k'\lambda \alpha \beta \lambda}$$

denoting phonon absorption and emission processes, respectively. During these scattering processes, the energy and crystal momentum conservation need to be satisfied, as clear in Eq. (22). Similar to the phonon interaction case, the electron phonon scattering processes can also be classified into normal and Umklapp types. Normal processes occur for electrons and phonons with relatively small wave vectors while Umklapp processes happen for those with large wave vectors[11].

For the phonon-phonon interaction, we have ignored the dependence of the collision terms on the states except the state of interest, implying that when considering the scattering of one phonon state, other states are assumed to be at equilibrium. For the electron-phonon
interaction, we will however show that the dependence of the electron scattering on other phonon states cannot be neglected in some situations. To simplify Eq. (20), we see that there are terms directly proportional to the state of interest (just as the case for phonon-phonon interaction), which can be rewritten by defining new relaxation times due to the electron-phonon coupling

$$\frac{1}{\tau_{k\alpha}} = \sum_{k'\beta, q} F_{k\alpha}(k', \beta, q) \quad (23)$$

The first relaxation time \( \tau_{k\alpha} \) describes the scattering of electrons by equilibrium phonons, and is the dominant scattering mechanism for the mobility of lightly-doped semiconductors near room temperature or above[11,13]. For normal electrical property (electrical conductivity and Seebeck coefficient) calculation, one uses the RTA and assumes \( \Delta f_{k\alpha} = 0 \) and \( \Delta n_{q\alpha} = 0 \) in Eq. (20), and therefore the collision term can be written in the form as Eq. (7), where the relaxation time includes the electron-phonon relaxation time defined by Eq. (23) and also the relaxation time due to impurities when necessary[13]. The treatment of the impurity scattering will be given below. The distribution function is then readily solved giving Eq. (8). The electrical conductivity is obtained by summing up all the electron states and dividing the total electrical conduction by the electrochemical potential gradient (assume temperature is uniform):

$$\sigma = \left( \frac{e}{3\Omega N_k} \sum v_{s,k\alpha} \Delta f_{k\alpha} \right) (\nabla \varphi) = \frac{e^2}{3\Omega N_k} \sum v_{k\alpha}^2 \tau_{k\alpha} \left( -\frac{\partial f_{k\alpha}^0}{\partial E} \right) \quad (24)$$

where \( N_k \) is the number of the discrete points for electrons in the reciprocal space. Note that we have used the deviation of the distribution since at equilibrium the electrical conduction will be zero. The Seebeck coefficient measures the induced voltage difference across a sample in response to a given temperature gradient. In this case we keep the term in \( \Delta f_{k\alpha} \) that is proportional to the temperature gradient and obtain
We call this Seebeck coefficient as diffusive because it originates from the diffusion of electrons. This is also distinguished from another contribution to the Seebeck effect which will be clear later. We can also obtain the electronic contribution to the thermal conductivity

\[ \kappa_e = \kappa_0 - T\sigma S^2 \]  

where

\[ \kappa_0 = \frac{1}{3\Omega N k T} \sum_{k_a} (E - \mu)^2 \mathbf{v}_{k_a}^2 \tau_{k_a} \left( -\frac{\partial f_{k_a}^0}{\partial E} \right) \]  

The second relaxation time \( \tau_{q\beta}^{e-ph} \) in Eq. (23) describes the scatterings of phonons by equilibrium electrons[34]. As we have mentioned, typically this is not as significant as the phonon-phonon scattering. However, if we carefully examine its dependence on the electron state, we will see that this type of scattering strongly increases as the Fermi level increases (or increased carrier concentration). This implied that, in heavily-doped semiconductors, the phonon scattering by electrons might play a non-negligible role. We will clarify this point more when discussing the first-principles results.

In evaluating the electrical property, we have assumed \( \Delta f_{k'\beta} = 0 \) in Eq. (20). We should note that this is neglected essentially because the terms containing \( \Delta f_{k'\beta} \) sum up to approximately zero. In metals and for elastic scattering with impurities, this approximation is not valid and therefore an extra correction term \( (1 - \cos \theta) \) is often added to the electron-phonon relaxation time to take into account the terms containing \( \Delta f_{k'\beta} \), leading to the so-called momentum relaxation time[13,35]

\[ \frac{1}{\tau_{e-ph,momentum}^{k\alpha}} = \sum_{k'\beta,q\lambda} F_{k\alpha}(k'\beta, q\lambda)(1 - \cos \theta) \]  

where \( \theta \) is the angle between the two velocity directions of the initial and final electron state. In semiconductors, however, it has been proven, based on deformation potential models,
that for nearly isotropic scattering, the neglect of $\Delta f_{k'\beta}$ will not cause much difference[13]. We will also show that without considering terms containing $\Delta f_{k'\beta}$, good agreement for the electrical properties in silicon with experiments can be achieved[36–38]. In essence, the momentum relaxation time is only an approximation to the full expression in Eq. (20), and further work should be done to evaluate the accuracy associated with the use of the two relaxation times for calculating the transport property of semiconductors within the first-principles approach.

Another important simplification we have made when evaluating the electrical property is $\Delta n_{q\ell} = 0$, for the reason that the frequent phonon-phonon Umklapp scatterings normally restrict the phonon distribution function close to the equilibrium[39]. This assumption makes non-equilibrium phonons have no effect on the electron system. However, below the Debye temperature, phonons have larger deviations from the equilibrium because the phonon scatterings are largely suppressed by the decreased temperature. These non-equilibrium phonons (non-zero $\Delta n_{q\ell}$) in the electron system (Eq. (20)) then contribute to an extra scattering for the electron’s collision terms. Under a given temperature gradient, we use the RTA for the phonon system and write $\Delta n_{q\ell} = \tau_{q\ell} \cdot v_{q\ell} \cdot \nabla T \cdot \frac{\partial n_{q\ell}}{\partial T}$. The electron-BTE can then be solved giving the electron distribution function (here we only consider the non-equilibrium part that results from the non-equilibrium phonons):

$$\Delta f_{k\alpha} = \tau_{k\alpha} \cdot \nabla T \cdot \sum_{k'\beta, q\ell} \left[ F_{q\ell}(k\alpha, k'\beta) \cdot \tau_{q\ell} \cdot v_{q\ell} \cdot \frac{\partial n_{q\ell}}{\partial T} \right]$$ (29)

Considering the resulting electrical current per temperature gradient, we arrive at an extra contribution to the Seebeck coefficient due to the non-equilibrium phonons

$$S_{\text{ph}} = \frac{2e}{3\Omega N_k N_q k_B T^2} \sum_{q\ell} \left[ \hbar \omega_{q\ell} \cdot v_{q\ell} \cdot \left( \sum_{k\alpha, k'\beta} \left( \tau_{k\alpha} v_{k\alpha} - \tau_{k'\beta} v_{k'\beta} \right) f_{k\alpha}^0 (1 - f_{k'\beta}^0) n_{q\ell}^0 \rho_{\ell} \right) \right]$$ (30)

This effect describes an extra current generated when there is a phonon heat flow due to the non-equilibrium phonons, as if the electrons were “dragged” by phonons. Therefore it has
been dubbed the phonon drag effect\cite{33,39,40}. We will have more discussions on this effect in section 4. Here we want to mention that the above picture describing the phonon drag effect is based on the Seebeck effect, where a temperature gradient induces a phonon heat flow, which delivers its momenta to the electron system and creates the electrical current. Because of the Kelvin relation $\Pi = TS$\cite{15,41,42}, an extra contribution to the Seebeck coefficient also implies an extra Peltier coefficient. This can be understood by the first two terms of the collision terms for phonon in Eq. (20), which transfer the momenta of the electron system to phonons when there is an electrical current and thus non-zero $\Delta f_{\alpha \mathbf{k} \alpha}$. We will not examine this second case but point out that the derivation based on the Peltier picture will lead to the same result as shown by Eq. (30).

All of the above transport properties are contained in Eq. (20), of which the key element is the electron-phonon interaction matrix element given by Eq. (19). In evaluating the electron-phonon coupling matrix, besides the electron wavefunctions and phonon modes, the change of the electron potential energy with respect to the atomic displacements is required. One way is to use the deformation potential model\cite{11,13}, which connects the magnitude of the perturbed potential to the change of the band edge energy with respect to the lattice dilation. By using the deformation potential, the large amount of the coupling matrix elements are not needed, thereby greatly saving the computational cost. However the use of this parameter ignores the dependence of the electron-phonon coupling on the electron states, and is only valid for band edge states and long wavelength phonons, which cannot be accurate for materials in general. A more accurate way is to use DFPT, because when calculating the phonon dispersion, DFPT already obtains the perturbed potential and therefore provides the electron-phonon coupling matrix as a by-product. However, different from the use of real space force constants in calculating the phonon-phonon coupling matrix, direct electron-phonon coupling matrix calculations using DFPT are time-consuming. It will thus be favorable to develop methods to reduce the DFPT calculation and uses formulas similar to Eq. (11) to obtain the electron-phonon coupling matrix for any state, since a fine mesh is necessary for the convergence of the transition rates. Such an approach has recently been developed\cite{43,44}, which utilizes the maximally-localized Wannier functions\cite{45} to transform
the DFPT results from the reciprocal space to the real space. In the real space, the electron states can be described by orbital-like states while the perturbed potential is centered around the displaced atom, both of which are localized in real space[43]. These essentially serve as the tight-binding basis and can be used to construct the electron and phonon states as well as their coupling matrix for any given states. In fact, this whole process can be understood as a generalization of the tight-binding model. Efficient calculation of electron-phonon coupling matrix can then be done with DFPT results on a coarse mesh only. This method has allowed the electron-phonon coupling constant to be examined for many materials, especially superconductors[46–48]. We refer readers to the literature for more details on the Wannier states[45] and the extraction of electron phonon matrix elements[43,44]. We also want to note that, as pointed out by the original work[43], the Wannier interpolation for the electron-phonon coupling matrix will be less accurate if there are long range interactions in the system, such as the electric field accompanied by the longitudinal optical phonon near the zone center in polar semiconductors. For these materials, these long range forces need to be separately treated, as what has been done for the LO-TO splitting in the phonon dispersion calculation using DFPT[19,49,49]. Recently these long-range effects have been incorporated into the Wannier interpolation scheme by using a rigid ion model (assigning effective charges to each atom and consider their Coulomb interactions with electrons) [50] to approximate the electron-phonon matrix element for long-wavelength LO phonons [51,52]. This development thus further extends the efficient computation of electron-phonon couplings towards polar semiconductors.

### 2.3 Impurity scattering

In semiconductor engineering, external methods that modify the physical properties of the materials are indispensable, with examples including doping and alloying. For the thermoelectric property in particular, doping changes the carrier concentration and is used to find the optimal carrier concentration for achieving a higher efficiency. On the other hand the alloying provides a way to modify the band structure for higher Seebeck coefficient[53] or to create more phonon scatterings for lower thermal conductivity[54]. In general, these material
processing techniques break the translational symmetry of the periodic crystal, leading to further scatterings of electrons and phonons. In most cases they are treated as perturbations to the pristine crystal, therefore the scattering rates can be studied by a perturbative approach. Generally we regard them as impurity scattering because they introduce irregularities into the host material and have similar treatment for the scattering rate calculation.

First we consider neutral impurities of a single type embedded in a medium. The key idea of the perturbative approach is to replace the real material with a pristine one, therefore justifying the use of the eigenstate[55]. In a more general sense, when considering a disordered crystal, one can replace that with an ordered virtual crystal that has an averaged property (structure, force constant, electron potential, etc.), which is also known as the virtual crystal approximation (VCA)[55,56]. The disorder is then treated as a perturbation to the system. For relatively long wavelength phonons, this approximation is reasonable because these phonons mainly see an averaged potential[55]. It is questionable when the wavelength of phonons is comparable to the characteristic length of the disorder. It has been shown by using a full-order scattering theory that, the VCA fails for high frequency phonons and predicts incorrect phonon density of state[57]. However, such high frequency phonons are shown to have large scattering rates[57], which make VCA still a good approximation in evaluating the heat transport of alloys.

The perturbation due to the introduction of the impurities leads to scatterings between the original states. Using the Golden Rule, we can write the scattering rates of phonons due to the impurity scattering as[58]

$$\frac{1}{\tau_{q\lambda}^{\text{ph-imp}}} = f \sum_{q\lambda'} \frac{\pi \Omega}{2 \omega_{q\lambda}^2} \frac{1}{V_s} \left| \langle q\lambda | T | q\lambda' \rangle \right|^2 \delta \left( \omega_{q\lambda} - \omega_{q\lambda'} \right)$$  \hspace{1cm} (31)

where $f$ is the volume fraction of the impurity, $V_s$ is the volume of the scatterer, $|q\lambda\rangle$ describes the phonon eigenmode normalized to the unit cell volume $\Omega$ and $T$ is the matrix that describes the perturbation induced by a single impurity that couples different phonon modes. Note that in Eq. (31) we have assumed dilute limit for which the interaction between impurities is ignored. In most cases, the T-matrix is approximated by the perturbation matrix,
known as the Born approximation, which can be shown to give rise to a scattering rate proportional to the frequency to the fourth power\cite{59,60} (Rayleigh scattering). Similarly, for electrons we can write down the scattering rates based on the potential difference $\Delta V$ between the pristine material and the one with impurity:

$$\frac{1}{\tau_{\text{imp}}} = \frac{c}{\Omega N_k} \sum_{k\beta} \frac{2\pi}{\hbar} |\langle k'\beta | \Delta V | k\alpha \rangle|^2 (1 - \cos \theta)$$

(32)

where $c$ is the density of the impurity, $|k\alpha\rangle$ describes the electron eigenstates and $\theta$ is the angle between the group velocity vectors of the initial and final state. This perturbative approach has been widely applied to study the influence of neutral defects on material’s physical property. The calculation is usually performed in a supercell where an atom is replaced by the defect. By comparing the results with those obtained in the same supercell of pristine crystal, defect-induced perturbation can be extracted and used for understanding how the defect modifies material’s property. A common example is the calculation of formation energy of different substitutes in some host materials\cite{61}. We will see that this method has been combined with the first-principles calculation to study the defect scattering of electrons and phonons. This perturbative thinking can also be applied to the transport studies of alloys using the VCA (instead of constructing a supercell) as we have mentioned. We will see that good agreements have been achieved for the electrical and thermal conductivity of SiGe alloy\cite{62,63}, as well as the thermal transport in some other alloys\cite{64–66}, thereby justifying the applicability of the VCA. It should be noted that, the perturbed potential (either for electrons or phonons) in the above situations are relatively short-ranged and therefore a supercell in which the perturbed potential decays to zero is possible within the DFT capability. In fact, the recovering of bulk potential at distances away from the defect has been used to set the correct energy reference shift in some calculations\cite{62}.

However, when the impurities introduced into the material are not charge-neutral (for example ionized dopants), they will create long-range Coulomb interaction, which decays much slower and makes the supercell approach inappropriate due to the interaction between the dopant and its artificial images entailed by the periodic boundary condition. Besides, when modeling the charged impurity in a supercell, the net charge density would lead to the
divergence of the total energy. The way to overcome this difficulty is to introduce a uniform jellium charge background, making the whole system charge-neutral. This trick however introduces the spurious interaction between the electron and the background charge. Though the spurious interactions decay to zero as the supercell size increases to infinity, within the current DFT capability such a large supercell is almost impossible.

For comparison between dopant elements in the same column, a supercell can still be used in spite of the long-range Coulomb potential[67]. This is because the elements have same charge and therefore the incorrect treatment on the Coulomb interactions will not affect their comparison. To quantitatively evaluate the electron scattering by the charged impurities, one must find ways to extend the screened Coulomb potential beyond the supercell. Rurali et al[68] have found in the silicon nanowire transmission calculation that the perturbed potential induced by a single charged impurity converges quickly around the dopant, and its values in the outer region of the dopant, though varying as the system size increases, can be reproduced by a simple screened electrostatic model. Therefore the potential corresponding to a long nanowire is constructed by connecting the converged potential in the inner region and the screened electrostatic potential in the outer region. Similarly, a screened Thomas-Fermi potential was used by Restrepo et al to extend the potential obtained from the first-principles calculation beyond the supercell, and the perturbed potential in the whole space is then used to calculate the electron impurity scattering rates in silicon[36]. However, we find that there has not been a consensus on which physical model is better suited for extending the screened Coulomb potential beyond the supercell and how accurate these methods are in different situations. Regarding the thermodynamic information (e.g. formation energy) of the point defects in solids, there have been great developments on the correction schemes for the spurious interactions due to the long-ranged forces[69]. It will be desirable to see how similar ideas can be used to facilitate the calculation of electron scattering rate due to the charged impurities, which has the control over the mobility of heavily-doped semiconductors.

3. Thermal Transport in Semiconductors
From this section on, we discuss the first-principles results on the transport properties of semiconductors. We first examine the thermal transport properties. Some previous review papers have discussed the first-principles calculations of the thermal conductivity, mainly for illustrating the power and accuracy of the first-principles methods [70,71], by comparing with the experimental data. In the following our discussions will specifically focus on the column IV and III-V semiconductors. To give a unified picture for both phonons and electrons, we will discuss their relaxation times and mean free paths from pure crystals to alloys. We show that the first-principles calculation not only enables us to investigate the thermal transport properties in a variety of materials with unprecedented details, but also provides new insights into some of the old topics, for which traditional perspectives may not be entirely correct.

3.1 Phonon relaxation times

Among all the semiconductors, silicon is probably the best-studied one. It can serve as a testbed for the first-principles computational tools and also provides insightful details into the phonon transport properties in general. However, a first-principles prediction of its thermal conductivity has been challenging, because of the requirement of obtaining accurate harmonic and anharmonic force constants that lead to phonon dispersion and phonon scattering rates. Force constants extracted from the empirical interatomic potential have been shown to give qualitative agreement with experiments but the accuracy is far from being satisfactory[72]. The thermal conductivity result also largely depends on the empirical potential chosen to use, therefore questioning such an approach as a quantitative tool for evaluating the thermal transport. By using the BTE formalism with force constants extracted from the DFT calculations, Broido et al were able to first show that the first-principles approach can predict the thermal conductivity of silicon and germanium within 5% of the experimental value at room temperature without any adjustable parameter[17], where a reciprocal space method is used to extract force constants. The real space approach, developed by Esfarjani et al[16,20], further allows an easy extraction of force constants in more complex materials, and is soon applied by many groups to other materials, including III-V semiconductors[73–75], II-VI binary compounds[76], typical thermoelectric materials[64–66,77–85] and 2D materials[86–93]. We should note that not all of these work
used the RTA. In fact, in materials where normal processes are more frequent compared to Umklapp processes like graphene, it was shown that an exact solution of the full BTE is necessary to obtain quantitative agreement with experiments[87,94], which is also used in the original work on silicon[17]. However for silicon, germanium and III-V semiconductors, Umklapp scatterings are relatively strong and as a result the use of RTA only introduces small errors compared to an exact calculation[16,74].

The first-principles approach allows us to look into the details of each phonon mode in contributing to the thermal transport. In Si and Ge, it has been found that more than 90% of the heat is carried by acoustic phonons at room temperature[17]. While the optical phonons carry much less heat than acoustic phonons, they provide significant scattering channels for the acoustic phonons. For Si and Ge, it was found that more than 50% of the scattering processes of the acoustic phonons involve at least one optical phonon[26]. Therefore, neglecting the optical branch (in the sense that their scatterings with acoustic phonons are also neglected) will largely overestimate the thermal conductivity. Similar results have also been obtained in GaAs, where over 90% of the heat is carried by acoustic phonons over a temperature range from 100K to 400K[75]. These results thus quantified our qualitative understanding based on the theoretical models.

From the first-principles method, we can further examine the mode-dependent relaxation times. It has been found that in Si and Ge the normal process scattering rate scales with $\omega^2$ while the Umklapp process scattering rate scales with $\omega^{3-d}[16,26]$. Figures 2a and 2b show the frequency-dependence of the relaxation times in Si. Normal scattering usually dominates for relatively low frequency phonons and the squared frequency dependence agrees with the commonly-used form, while the Umklapp process starts to dominate at higher frequencies, giving rise to a higher-order frequency dependence. However such higher order frequency dependence has been neglected in many recent room temperature RTA calculations. The first-principles results thus highlight the importance of using the correct scaling behavior of the relaxation time, which directly affects the estimation of the thermal conductivity[26]. The relaxation times of GaAs are also shown in Fig. 2, exhibiting similar frequency scaling behavior[75]. The majority of the relaxation times at high symmetry points at 300K are
within the range of 3~10 ps, while TO modes at L and X points have longer relaxation times of ~30 ps, as shown in Fig. 3. The quantitative determination of these relaxation times therefore provides critical information for hot-phonon effects in many optoelectronic devices.

Figure 2. Phonon relaxation times from first-principles with respect to the phonon frequency for (a) Umklapp processes in Si, (b) normal processes in Si, (c) Umklapp processes in GaAs and (d) normal processes in GaAs. Note that for the low frequency phonons, the normal process relaxation time has been fitted with $\omega^2$ while the Umklapp process relaxation time is fitted using $\omega^3$. (Si results reprinted with permission from Esfarjani et al [16]. Copyright (2015) American Physical Society; GaAs results reprinted with permission from Luo et al [75]. Copyright (2015) Institute of Physics)

Figure 3. Phonon scattering rates in GaAs as a function of temperature for (a) LO phonon at $\Gamma$-point, (b) TO phonon at $\Gamma$-point, (c) TO phonon at L-point and (d) TO phonon at X-point. Shown in the figure is the
total scattering rate as well as its decomposition into different types of three-phonon scattering processes (op: optical phonon; ac: acoustic phonon). The obtained first-principles results (lines) are compared to previous first-principles calculations (Ref. (A) from Ref. [95] and Ref. (C) from Ref. [96], only performed at Γ-point) as well as experimental data (Ref. (B) from Ref. [97,98]). (Reprinted figure with permission from Luo et al [75]. Copyright (2015) Institute of Physics)

Such first-principles calculation has also been applied to study the heat conduction in superlattice[99], where one obtains the force constants from a virtual crystal calculation with averaged masses and the masses of different atoms are included via Eq. (11). For an infinite GaAs/AlAs superlattice, it has been found that by treating the interface roughness as a random mixing of Ga and Al atoms in a narrow range, the low frequency phonons dominate the heat conduction and are mainly affected by phonon-phonon scatterings, while higher frequency phonons are also scattered due to the interfacial disorder[99], as shown in Fig. 4.

![Figure 4](image.png)

**Figure 4.** Phonon relaxation times from first-principles with respect to the phonon frequency for an infinite GaAs/AlAs superlattice. The plot shows the intrinsic phonon scattering rates at two different temperatures (100K and 300K) as well as the temperature-independent interfacial scattering. It is clear that for low frequency phonons (less than 1THz) the anharmonic scattering dominates while for higher frequency phonons the interfacial scattering starts to play an important role. (Reprinted figure with permission from Luckyanova et al [99]. Copyright (2015) American Association for the Advancement of Science)

The origin of some thermal transport features can also be understood by studying the scattering channels for each phonon branch. Work has been done to systematically examine the temperature dependent thermal conductivity of III-V semiconductors using first-principles[74], including all the binary compounds determined by (Al, Ga, In)-(P, As, Sb). The effects of different elements on the phonon properties can then be conveniently compared. It was found that the gap between the acoustic and optical branch will modify the
scattering processes of the acoustic phonons (due to the requirement of the energy conservation), and therefore consistently affect the thermal conductivity when the V-element moves downwards in the periodic table. By taking into account the phonon-isotope scattering (the treatment essentially uses Eq. (31) but with Born approximation, which reduces to the Tamura model[60]), good agreements with the experiments can be achieved in these materials[74]. A further study into GaN reveals that the isotope scattering is relatively strong compared to the intrinsic phonon-phonon scattering, and an isotopic enrichment should increase the thermal conductivity of GaN by ~65% at room temperature [73]. This large “isotope effect” was claimed to result from (1) a large gap between the acoustic and optical branch and (2) a large phonon energy scale, both of which reduce the phonon-phonon scattering rates and make the isotope scattering relatively more significant [73]. These calculations offered new insights into the isotope effect in semiconducting materials and have implications for engineering the thermal transport properties.

3.2 Phonon mean free path distribution

The first-principles method can also provide details into the mean free path distribution of phonons, which is important information for nanoscale applications due to the size effect. To clarify the contribution to the heat conduction from different phonon modes, a thermal conductivity accumulation function has been introduced by Dames and Chen[100], which is defined as the contribution to the thermal conductivity from all phonons with mean free paths below certain value. Although a kinetic formula $\kappa = \frac{1}{3} c_v v \Lambda$ typically uses a single mean free path, the real crystal can have a mean free path distribution spanning across several orders of magnitude from submicron to hundreds of microns. For example, an analysis based on the kinetic formula predicts a mean free path of 41 nm in Si [101]. The spectral phonon property analysis based on the molecular dynamics simulation, however, shows that the phonon mean free paths in Si span from 30 nm to 100 \( \mu \)m at room temperature[102]. This emphasizes the importance of recognizing the broad mean free path distribution when considering the nanostructures where long-mean-free-path phonons will be more frequently
scattered by interfaces and boundaries, for which a diffusive transport model will break down and ballistic effects must be considered.

Compared to the molecular dynamics simulation, we regard the first-principles calculation as a more accurate way to look into the spectral phonon properties of materials. It has been shown that, at room temperature more than 50% of the heat is carried by phonons with mean free paths longer than 1 μm in Si [16]. This fraction will become larger at lower temperatures. It should also be noted that in Si the mean free path of most phonons span almost four orders of magnitude, from 10 nm to 100 μm at room temperature (Fig. 5a). Similarly in GaAs, the mean free paths span over three orders of magnitude from 10 nm to more than 30 μm[75]. It was also found that more than 50% of the thermal transport comes from phonons with mean free paths longer than 350 nm at room temperature, and at a lower temperature (95K) 50% of the heat is carried by phonons with mean free paths longer than 4 microns (Fig. 5b). In an infinite GaAs/AlAs superlattice, phonons with mean free paths longer than 216 nm were found to contribute 87% (at 100K) and 71% (at 300K) to the total heat conduction[99] (Fig. 5c). We want to briefly mention that, the recent development in the phonon mean free path spectroscopy has opened up the possibility of reconstructing such mean free path accumulated thermal conductivity information from the experimental data based on the quasi-ballistic heat transport[103–105]. Good agreement has been achieved in Si between the reconstructed accumulation curve and the first-principles results[106–108]. Clearly such understanding of the mean free path distribution is crucial for modeling and engineering the nanostructures, and the first-principles calculation provides an accurate evaluation for these information.
Figure 5. Accumulated contribution to thermal conductivity from first-principles with respect to the phonon mean free path for (a) Si, (b) GaAs and (c) GaAs/AlAs superlattice. The results for Si are evaluated at 277K while the other two materials are evaluated at both room temperature and cryogenic temperatures. The “sawtooth” features for long MFP phonons are due to the finite mesh size used. (Si results reprinted with permission from Esfarjani et al [16]. Copyright (2015) American Physical Society; GaAs results reprinted with permission from Luo et al [75]. Copyright (2015) Institute of Physics; GaAs/AlAs superlattice results reprinted with permission from Luckyanova et al [99]. Copyright (2015) American Association for the Advancement of Science)

3.3 Effect of alloying and doping

First-principles calculation on an alloy system was first performed for SiGe alloy using the virtual crystal approximation by Garg et al [63]. It was uncovered that more than 50% of the heat conduction is carried by phonons with mean free path longer than 1 μm, as shown in Fig. 6. Besides, the large reduction of the thermal conductivity with only a small fraction of Ge added into Si is well captured by the first-principles calculation and agrees reasonably well with the experiment at room temperature. Compared to the pristine crystals, the mass disorder more strongly scatter high frequency phonons and make low-frequency (long-wavelength) phonons more significant in the thermal transport. Similar materials, including PbTe$_x$Se$_{1-x}$, Mg$_2$Si$_x$Sn$_{1-x}$ and BiSb alloy, have also been studied using the same VCA [64–66]. These results provide a deeper understanding of the thermal transport in alloy system and also guidelines for engineering the thermal performance of an alloy structure.

Figure 6. Accumulated thermal conductivity from first-principles with respect to phonon mean free path in Si$_{0.5}$Ge$_{0.5}$ alloy. (Reprinted figure with permission from Garg et al [63]. Copyright (2015) American Physical Society)
Doping procedure introduces ionized dopants and electrons into the host material, which also increase the scattering rates for phonons. It has long been thought that the reduction of the thermal conductivity of semiconductors is mainly due to the impurity scatterings from the dopants. The electron scattering of phonons, though formalized and discussed in metal systems [34,109,110], was regarded as insignificant for semiconductors. With the first-principles approach, one can now incorporate the phonon scattering rate due to electrons (the second line in Eq. (23)) into the intrinsic phonon-phonon scattering rate (Eq. (10)) using the Matthiessen’s rule \( \frac{1}{\tau_{q,i}} = \frac{1}{\tau_{q,i}^{ph-ph}} + \frac{1}{\tau_{q,i}^{e-ph}} \). The effect of electron scattering of phonons can then be evaluated at different temperatures and doping levels. Surprisingly, it has been found in Si that the thermal conductivity starts to decrease as the carrier concentration goes beyond \( 10^{19} \text{cm}^{-3} \) and the reduction reaches 45% in p-type silicon at around \( 10^{21} \text{cm}^{-3} \) [111], as shown by Fig. 7a. The major scatterings due to electron-phonon coupling are found in low frequency acoustic phonons and optical phonons, as shown in Fig. 7b. In fact, based on the deformation potential model, it can be found that the phonon scattering rates due to electrons linearly scales with phonon frequency [111] while as we have discussed above the intrinsic phonon-phonon scattering is dominated by the normal process for low frequency phonons, with a frequency dependence as frequency to the second power. The latter therefore drops faster than the former and as the carrier concentration increases low-frequency phonons will be more strongly scattered by the electrons. This finding has great implications for thermoelectric applications, which typically involve heavily-doped semiconductors, and also provide a modern understanding into the effect of the electron-phonon coupling on the thermal transport.
Figure 7. (a) Reduction of lattice thermal conductivity due to the phonon scattering by electrons for both n-type (blue curve) and p-type (red curve) silicon at 300K, and (b) the comparison between phonon-phonon scattering rates (red dots) and phonon-electron scattering rates (blue dots) for n-type silicon. The carrier concentration for part (b) is assumed to be $10^{21}$ cm$^{-3}$. (Reprinted figures with permission from Liao et al [111]. Copyright (2015) American Physical Society)

4. Electrical Transport of Semiconductors

While the first-principles calculation for the phonon thermal transport has been largely improved in the past decade, most of the evaluations of the electrical transport properties still require the use of parametrized models such as constant relaxation time approximation or deformation potential approximation[112]. These parametrized models for the electron scattering process have helped to understand the electron transport properties in certain thermoelectric materials and give qualitative agreement with the experiment [113–117]. However, such methods lack the accuracy if the band structure becomes complicated, in which case the electron phonon coupling no longer satisfies a simple parametrized model. Therefore it is necessary to have an accurate description for the electron phonon scattering through the first-principles approach. Only till very recently has the first-principles method been applied to study the electrical properties including mobility and Seebeck coefficient in silicon[36,37]. For the first-principles calculation of electrical properties using parametrized models for electron scattering, readers can refer to Ref. [112]. Here only the work that has applied the full calculation of the electron-phonon scattering rates will be discussed in detail. As a widely-studied material, silicon still presents a challenge to the electrical property calculation and can offer new perspectives to the thermoelectric application with
detailed study into its electrical transport as we will show. In the following we will discuss this material and some other efforts on III-V semiconductors, and also explain the difficulties involved in these calculations.

4.1 Electron relaxation times, mean free paths and mobility

Lying in the center of the electrical property calculation is the electron relaxation time. For semiconductors there are two main mechanisms that lead to the scatterings of electrons: electron-phonon scattering and electron-impurity scattering. In lightly-doped silicon, the electron-phonon scattering dominates over most of the temperature range. As the doping level increases, the electrons near the band edge are strongly scattered by the ionized dopants, resulting in a decreased mobility. Traditionally, the scattering has been treated by making approximations for the interaction matrix. For example, the electron-phonon interaction matrix element is readily described by the use of deformation potential while the electron-impurity interaction is approximated using screened Coulomb potential model[13]. By considering both electron-phonon and electron-impurity scattering in solving the BTE formula, good agreements have been achieved for the carrier concentration dependent mobility compared to the experiment[113]. However, the use of the parametrized model neglects details of the electron transport such as the dependence of the electron-phonon coupling on the electron energy and wave vector, and therefore is not satisfactory as a quantitative predictive tool. The first-principles approach provides such information with no adjustable parameters, and is more accurate and suitable for studying the contribution to the electrical transport from each electron state. The first-principles method was first applied to study the intervalley scatterings between certain electron pockets in Ge, GaAs and GaP [118,119]. The results obtained agree well with the experiments, justifying the use of first-principles approach in evaluating the electron-phonon scattering rate. These results also have implications for high-field electrical transport and the relaxation of hot carriers. However, to accurately predict the electrical transport property, we must recognize the difference of the relaxation times between different electron states. Such an analysis based on first-principles calculation was given by Restrepo et al[36], who showed that the electron
scattering rates in silicon on a large energy scale (several electron volts) follow the shape of the electron density of state. This general feature comes from the fact that at energy levels away from the band edge, the scattering rates are mainly limited by the available states that one electron can be scattered into, and therefore higher density of state provides more scattering channels and reduce the scattering time. In their original work, only the energy dependence of the scattering rate is explicitly given[36]. A further detailed study into Si and Ge shows that the electron-phonon coupling strength varies significantly along different crystallographic directions, which causes wave-vector dependence of the scattering rates as well[120], as shown in Figs. 8a and 8b. We note that such a full examination of the electron-phonon scattering over the whole Brillouin zone has enabled the analysis of the phonon-assisted optical absorption as well as hot carrier relaxation process in silicon[14,121], and is also crucial for a first-principles prediction of the electrical property. Hot carrier relaxation has also been examined in GaAs with the electron scattering times being studied over the whole Brillouin zone[122] (Fig. 8c). It was suggested that the polar optical phonon scattering is not as strong as what had been previously believed and does not dominate over the acoustic phonon scattering for hot electrons.

Figure 8. Electron-phonon scattering rates compared with the electron density of state for (a) Si, (b) Ge and (c) GaAs. It is shown that the scattering rate profile in a larger energy scale closely follows the density of state profile. The wave-vector-dependence of the scattering rates can be seen by the scatter of the data at the same energy level. The above calculations assume pure crystals and therefore the Fermi level lies inside the band gap. (Si and Ge results reprinted with permission from Tandon et al [120]. Copyright (2015) American Institute of Physics; GaAs results reprinted with permission from Bernardi et al [122]. Copyright (2015) United States National Academy of Sciences)

These calculations based on the first-principles approach have offered insights into the regimes where old models cannot reach or reliably analyze, such as the relaxation of hot
electrons with a wide distribution of lifetimes and the transition probabilities between different pockets examined independently. For the transport property such as low-field mobility and Seebeck coefficient, the electrons near the band edge (within a few $k_BT$ from the band edge) are the dominant players, because only these electrons are notably thermally excited due to the Fermi-Dirac statistics. Their mode-dependent scattering rates are more sensitively affected by the environment such as impurities, and require a closer examination. Figure 9a shows the electron scattering rates in silicon close to the band edge[38]. The “kink” at an energy around 0.06eV above the conduction band edge is due to the phonon emission process (note that the electron needs to have some energy above the conduction band edge to emit a phonon). It is shown that a fine mesh and an adaptive Gaussian broadening parameter for approximating the delta functions when evaluating the energy conservation are required to resolve this “kink” feature [38]. As the doping level increases, the band edge electrons will be scattered more strongly by ionized dopants, causing a U-shape scattering rate profile as shown in Fig. 9b. Band edge electron relaxation times have also been calculated in GaAs, where “kinks” in the scattering rate profile can be associated with different electron pockets (Fig. 9c), and a GW calculation (including electron many-body effect) was used to accurately describe the band structure[122].

In addition to the relaxation times, electron mean free path is another important piece of...
information for nanoscale modeling and engineering. Figure 10a shows the electron mean free path as a function of the electron energy. At lower doping concentrations, there is a maximal electron mean free path with respect to the electron energy. This is because at higher energies electrons suffer more scatterings (Fig. 9) and have smaller relaxation times, while near the band edge the group velocity tends to be zero. The competence between the relaxation time and the group velocity gives rise to the maximal electron mean free path. Figure 10b further shows the accumulated contribution to the electrical conductivity with respect to the electron mean free path[37]. Compared to the phonon mean free path distribution, we see that electron mean free paths span over a smaller range than those of phonons, from 10 nm to 100 nm for lightly-doped silicon and from 1 nm to 10 nm for heavily-doped silicon. The sharp cut-off of the accumulation curve for the electrical conductivity is due to the maximal electron mean free path as we have discussed above. The clear difference between the phonon mean free path and electron mean free path can be used to quantitatively examine the effect of nanostructuring techniques in reducing the phonon thermal conductivity while maintaining the electrical transport properties[37]. However such detailed mean free path information based on first-principles calculations has not been reported for other semiconducting materials. We anticipate that along the path more materials can be studied with detailed examination into their electron scattering mechanisms and mean free path distributions.

Figure 10. (a) Electron mean free path as a function of the electron energy and (b) accumulated contribution to the electrical conductivity with respect to the electron mean free path compared to the accumulated thermal conductivity contribution (with respect to phonon mean free path) in silicon from 100K to 400K. In part (b), two different doping levels (lightly-doped and heavily-doped) are shown. (Reprinted figures with permission from Qiu et al [37]. Copyright (2015) Institute of Physics)
Given the relaxation time of each electron state, mobility can be calculated based on the BTE solution (Eq. (24)). An accurate prediction of the transport property will involve integration over the whole Brillouin zone, thereby requiring a large amount of relaxation time data, which in turn depend on the scattering processes between many electron and phonon states. As a result, scattering rates between certain pockets are not sufficient in describing the transport property and the first-principles calculation of mobility has always been challenging due to the large mesh entailed. Restrepo et al first reported the mobility calculation for silicon including both electron-phonon and electron-impurity scattering[36]. The phonon-limited mobility (neglect the impurity scattering) for n-type Si is calculated to be 1970 cm$^2$/Vs at low carrier concentration, which agrees reasonably well with the experimental mobility (~1700 cm$^2$/Vs [123]) of lightly-doped silicon but slightly larger. The mesh density was not reported in their work but a more accurate calculation will inevitably require a dense mesh. A linear interpolation scheme [38] that obtains the electron-phonon matrix elements on a fine mesh has been shown to give rise to a mobility value (~1860 cm$^2$/Vs) closer to the experimental result. The linear interpolation works well in this case essentially because the electron-phonon coupling matrices do not vary much in the first Brillouin zone for Si. As more complicated materials are considered, however, we consider the Wannier interpolation [43] as a more accurate approach for extracting the electron-phonon coupling matrix.

4.2 Seebeck coefficient

On the other hand, the Seebeck coefficient requires a more careful look. This is because in lightly-doped silicon the phonon drag effect has been found to be significantly strong at low temperatures[39,124]. Therefore when considering the Seebeck coefficient we should add up both the diffusive contribution (Eq. (25)) and the phonon drag contribution (Eq. (30)). We note that in the expression for the phonon drag Seebeck coefficient both relaxation times for electrons and phonons are required, thereby making the calculation even more challenging than the mobility calculation. The calculation using first-principles-extracted deformation potential was able to describe the diffusive Seebeck coefficient[113], which is obtained by removing an empirical phonon drag contribution from the measured data. Recently Mahan et
al combined the first-principles phonon relaxation times with the deformation potential model to describe both the diffusive and phonon drag contribution to the Seebeck coefficient, and has seen a good agreement with experiments across a wide temperature range [125]. However, it was found that the phonon drag from the simulation increases faster with decreased temperature than the experimental value, which could result from the use of the deformation potential model. In a later study using a full first-principles calculation with both electron and phonon relaxation times extracted from DFT calculations, the Seebeck coefficient including the phonon drag was predicted with excellent agreements compared to the experiment [126], as shown in Fig. 11a. This emphasizes the use of a first-principles approach for better description of coupled electron-phonon transport. These calculations [125,126] quantitatively investigated the temperature dependence of the phonon drag effect: although it dominates at low temperatures, it has influences even beyond the room temperature. It was shown that at room temperature in lightly-doped n-type (p-type) silicon the phonon drag contribute to 30% (40%) of the total Seebeck coefficient [126]. As we have mentioned, phonon drag effect can be understood as the momentum transfer from non-equilibrium phonons with long mean free paths to the electron system. In such processes, the phonons are the key players and one important information will be how much are the contributions from different phonons. Figure 11b shows the accumulated contribution to phonon drag with respect to the phonon mean free path. It can be seen that the phonons that contribute to the phonon drag mostly have long mean free paths (small wave vectors) compared to those thermal phonons that carry heat. This is essentially because of the momentum and energy conservation for the electron-phonon scattering process, which restricts the wave vector of phonons to be small so that they can scatter band edge electrons (normal process). Clearly, we see from Fig. 11b that there is a spectral difference between the contributions to the thermal conductivity and to the phonon drag from phonons. Such quantitative information can be used to engineer the material’s property by modifying the phonon spectrum. For example, one can enhance the thermoelectric performance by filtering out short-mean-free-path phonons, in which case the phonon drag will be retained but the thermal conductivity can be reduced, a strategy recently proposed in Ref. [126].
Figure 11. (a) First-principles calculation of the temperature dependent Seebeck coefficient in pure Si and (b) the accumulated contribution to the phonon drag. In part (a) the total Seebeck coefficient is also decomposed into the diffusion part and the phonon drag part, the latter of which dramatically increases as the temperature decreases. The results agree well with the experimental data[124]. In part (b) the accumulated curves are shown for three temperatures (solid lines for 300K, dashed lines for 200K and dotted lines for 100K). The first-principles results are from Ref. [126].

Furthermore, by considering the phonon scattering by electrons in the phonon relaxation time calculation, the reduction of the phonon drag effect at higher carrier concentrations, known as the saturation effect[39], can also be captured by the fully first-principles approach[126], with a good agreement with the experiment shown in Fig. 12. This reduction comes from the increased scatterings of phonons with long wavelengths by electrons. As shown above, phonons that contribute to phonon drag mostly have small wave vectors and thus low frequencies. For these low frequency phonons, the electron scattering becomes appreciable as we have discussed in section 3. The impurity scattering is in fact insufficient to strongly affect these phonons despite of their effect on the mobility, because it scales as $\omega^4$ and decreases quickly as the phonon frequency becomes small. It was further uncovered that[126], though the phonon drag is largely reduced due to the saturation effect, it is not negligible for a heavily-doped silicon. In fact its contribution is comparable to the diffusive Seebeck coefficient at $10^{19}$ cm$^{-3}$ even at room temperature (Fig. 12). This result challenges the previous belief that the phonon drag effect vanishes in heavily-doped samples, and provides new perspectives into the electron-phonon coupled transport of the decades-old material - silicon.
The total Seebeck coefficient (black curves) is also decomposed into the diffusion part (green curves) and phonon drag part (red curves). The phonon drag at low carrier concentration does not depend on the doping level. The dotted lines assume that this value is used throughout the whole doping range. Clearly for high doping concentrations the dotted line overestimate the Seebeck coefficient, and the difference is due to the scattering of phonons by electrons, leading to a reduction of the phonon drag part. The first-principles results are from Ref. [126].

We have seen that the use of the first-principles method has provided us insightful details into the electrical transport properties in silicon. However, up to now there are only few materials whose mobility or Seebeck coefficient has been calculated using a parameter-free first-principles approach, with examples including silicon, MoS$_2$ and phosphorene [36–38,126,127]. Here we deem the use of the deformation potential model (even with deformation potential extracted from the first-principles) as an complementary but not a predictive method, because it neglects the electron-state-dependence of the electron-phonon coupling, which is hardly justified for general materials, especially considering the fact that the material search has been going towards more complex ones. Large discrepancy has also been seen between different studies using similar deformation potentials extracted from DFT calculations for the same material. We however believe that a detailed study into the validity of the deformation potential by comparing that with first-principles results will help to elucidate its applicability in the wide range of many new and unexplored materials.

One difficulty associated with the first-principles electrical property calculation of other materials is that, efficient interpolation of the electron-phonon matrix elements from a coarse onto a dense mesh, necessary for the integration to obtain the relaxation times, requires the force constant (or the perturbed potential) to be short-ranged in real space[43]. However in
materials with more than two elements, accompanied with the longitudinal optical (LO) phonon near the zone center there will be a polarization field, which induces an electric field that can cause scatterings for electrons. This scattering due to the long-range electric field induced by polar LO phonon is usually called polar optical phonon scattering. Therefore, the interpolation scheme as we have discussed before will be less accurate if the transport property of the material is governed by such scatterings. For a finite phonon wave vector, DFPT exactly solves the eigen-equation and therefore automatically includes such long-range effect. This merit of DFPT is also used in the study of hot electron relaxations in GaAs[122]. The interpolation method for the electron-phonon coupling matrix, however, needs to be modified to correctly capture the long-range effect. Work has been down to use the first-principles band structure combined with scattering time models to evaluate the electrical property of other materials. For example the mobility of SrTiO$_3$ was modeled using the Fröhlich model for the interaction matrix[128]. Such approach however does not differentiate the importance of different scattering mechanisms in affecting the transport property. Recently Sjakste et al and Verdi et al incorporated the polar optical phonon scattering into the Wannier interpolation scheme by using a rigid ion model to treat the long-range effect, where each atom carries certain effective charge due to their ionic character and the polar scattering mainly comes from the Coulomb interaction between them and the electrons [51,52]. The interpolated electron-phonon matrix elements agree well with those obtained from direct DFPT calculations [51,52]. We believe the inclusion of the polar scattering will facilitate the study of the electrical transport of more complicated semiconductors and provide more possibilities for engineering material’s electrical property.

4.3 Effect of alloying

Using a perturbative approach with information extracted from DFT calculations, Murphy-Armando and Fahy examined the alloying scattering for electrons in SiGe alloy[62]. They quantified the contribution to the electron scattering rates from intervalley scattering and intravalley scatterings, and found that they are comparable with each other, contrary to some previous modeling results claiming that intervalley scattering is negligible. They have
also obtained good agreements with the experimental value by incorporating the deformational potential model for describing the electron-phonon scattering. Particularly, the first-principles results correctly captured the abrupt change of the mobility in Si$_{1-x}$Ge$_x$ alloy at $x \sim 0.85$, which is due to the $\Delta - L$ band crossing[62]. We note that the perturbed potential $\Delta V$ in this case is the potential difference between the pristine Si and pristine Ge, while the electron wavefunctions are obtained in the virtual crystal with averaged properties. Care must be taken to set the reference for the potential energy[62].

The same perturbative approach has been applied to compare the effect of different substitutes on the electron scattering within the same column[67]. As we have mentioned in section 2, the supercell method will inevitably introduce spurious interactions due to the image atoms and background compensating charges, therefore questioning a direct evaluation of the perturbed potential. However, in the same column the element has the same number of valence electrons and the errors thus introduced will be the same. The difference mainly comes from the scattering due to strain-induced disorder. We note that to compare elements in different columns, an accurate description of the perturbed potential will be necessary. One way to achieve this has been illustrated by Restrepo et al in their calculation of electron-impurity scattering rate[36]. Other techniques borrowed from the treatment of calculating formation energy [69] can also be possibly used to develop an efficient method for describing electron scattering by charged impurities.

Besides the alloying scattering for electrons, the alloy structure also modifies the phonon modes and therefore the coupling between electrons and phonons. In the lowest-order approximation, phonon eigenmodes as well as the perturbed potential can be obtained in the virtual crystal. The electron-phonon scattering rates are then calculated based on these virtual crystal perturbation combined with the electron modes of the virtual crystal, assuming the electronic disorder is weak[129]. However the disorder effect on the electron-phonon coupling has been ignored in such approach. We would expect that a larger supercell calculation will help to understand how the electron phonon coupling is modified by the alloy structures, with detailed studies into the electron relaxation times and mean free path distribution, providing crucial information for evaluating the electron size effect at nanoscale.
5. Summary

We have reviewed the recent development of the first-principles approach to obtain the transport properties (in particular, electrical conductivity, Seebeck coefficient and thermal conductivity) of materials, which provides us a better understanding of the transport features and leads to rational designs of material properties. The coupled electron-phonon Boltzmann transport equation has been introduced, discussed and used for relating the transport property to the eigenmodes (electrons, phonons) and their coupling with each other. The relaxation time is introduced as a measure of the decay of non-equilibrium distribution back into equilibrium state, and is the key variable in calculating the transport property under the so-called relaxation time approximation. Relaxation times derive from the scatterings between the states, which are governed by the coupling due to perturbations introduced to the original system. For phonon-phonon scattering, the perturbations come from the anharmonic interatomic forces. For electron-phonon scattering, the atomic displacements perturb the environments seen by the electrons. Besides, impurities naturally introduce perturbations into the original pristine crystal. All of these can be analyzed using the same framework and we have seen that the crucial ingredient in describing these scattering processes and thus relaxation times is the coupling matrix that determines the transition probabilities from one state to another.

Methods to extract the electron-phonon coupling matrix based on first-principles calculations are briefly discussed. For the thermal conductivity calculation, the important information is the anharmonic force constant while for the electrical transport properties the electron-phonon coupling matrix is the vital component. An efficient extraction of such information is needed, due to the requirement for the convergence of the relaxation time calculation involving the states across the Brillouin zone.

We have seen that the calculation of thermal transport has been greatly improved in recent years, from simple semiconductors to more complex systems. Some practical systems like alloys can now be calculated with good agreements with experiments. Further direction could be the calculations of even more complex materials like perovskites and organic semiconductors. These complex materials are difficult for a first-principles calculation mainly because the computational
time scales approximately as $N^3$ ($N$ is the total number of atoms in the unit cell) due to the non-locality inherent in quantum mechanics [4]. An order-N method (computational time scales as $N$) that uses certain features of the system to reduce the computational load has seen great advantage over traditional method[4], but has not been introduced into the first-principles calculation of the thermal conductivity. Another possible strategy for accelerating the calculation is to use effective force constants to represent some atomic cluster in a large unit cell. For example, the inorganic-organic perovskite semiconductor CH₃NH₃PbI₃ which recently gains the popularity due to its potential in solar cells [130] has an ABO₃ structure with site-A replaced by an organic group. If the organic group can be represented by a meta-atom with effective force constants (between this meta-atom and other atoms), the total number of atoms can be greatly reduced. This strategy, if possible, can facilitate the first-principles thermal transport study into many inorganic-organic hybrid materials.

In comparison with the first-principles calculation for thermal transport, calculations of electrical transport in semiconductors have only received attention recently and up to now only few materials (silicon, phosphorene, MoS₂, etc.) are calculated within a fully first-principles approach for the electrical transport properties [36–38,126,127]. Other methods usually use parameterized models such as the deformation potential model, fitted from experiments or first-principles calculations. The first-principles method can provide insightful details into the relaxation time profile and mean free path distribution of the electrons, which are of great significance for modeling and engineering nanoscale devices. Besides, unprecedented details in the transport property have been seen by using the first-principles approach. For example, the phonon drag effect was seen to play a non-negligible role in the Seebeck coefficient even in heavily-doped samples at room temperature. Along this path, we think some of the challenges include the consideration of the spin-orbit coupling for heavy elements and alloy (and doping) effects on the electrical transport. Since the band structure greatly affects the scattering rate profile as we have seen in GaAs[122], more advanced first-principles method such as GW calculation (a method to more accurately describe the electron-electron interaction with G stands for Green’s function and W stands for the screened Coulomb interaction) to take into account the electron many-body effects on the band structure will be necessary for a better description of the electronic properties.
In addition, one can develop the theory to recast the transport property calculation based on BTEs using the matrix elements between different eigenstates into one that only uses the matrix elements in the Wannier basis, the latter of which requires less computational work but contains all the essential information, a strategy already mentioned in Ref. [43]. The Wannier basis (for electron and phonons) may also serve as good starting points for the study of transport problems in disordered materials (such as polymer or amorphous structure), where the periodic boundary condition breaks down and the dynamics of the system is more often described using spatially-localized modes. Another challenging question is how to concurrently solve the coupled electron-phonon BTEs, putting the effect of electron on phonons and that of phonon on electrons on an equal footing. For this, one has to use iterative solver to deal with the BTEs, which goes beyond the relaxation time model. This further step will provide us more information for the coupled electron phonon transport. It is also possible that unique physical behaviors may emerge from such coupled transport problem. In general, the advancement of the first-principles computational technique, as we believe, extends our toolbox for studying the transport properties in a quantitative way and will finally open up the venue towards a high-throughput material search based on *ab initio* predictions.
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