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ab initio Electronic Transport Model with Explicit Solution to the Linearized Boltzmann Transport Equation

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Abstract

Accurate models of carrier transport are essential for describing the electronic properties of semiconductor materials. To the best of our knowledge, the current models following the framework of the Boltzmann transport equation (BTE) either rely heavily on experimental data (i.e., semi-empirical), or utilize simplifying assumptions, such as the constant relaxation time approximation (BTE-cRTA). While these models offer valuable physical insights and accurate calculations of transport properties in some cases, they often lack sufficient accuracy – particularly in capturing the correct trends with temperature and carrier concentration. We present here a transport model for calculating low-field electrical drift mobility and Seebeck coefficient of n-type semiconductors, by explicitly considering relevant physical phenomena (i.e. elastic and inelastic scattering mechanisms). We first rewrite expressions for the rates of elastic scattering mechanisms, in terms of ab*initio* properties, such as the band structure, density of states, and polar optical phonon frequency. We then solve the linear BTE to obtain the perturbation to the electron distribution – resulting from the dominant scattering mechanisms – and use this to calculate the overall mobility and Seebeck coefficient. Therefore, we have developed an ab initio Model for calculating mobility and Seebeck coefficient using Boltzmann Transport (aMoBT) equation. Using aMoBT, we accurately calculate electrical transport properties of the compound n-type semiconductors, GaAs and InN, over various ranges of temperature and carrier concentration. aMoBT is fully predictive and provides high accuracy when compared to experimental measurements on both GaAs and InN, and vastly outperforms both semi-empirical models and the BTE-cRTA. Therefore, we assert that this approach represents a first step towards a fully ab initio carrier transport model that is valid in all compound semiconductors.

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Keywords: band transport model, mobility, Seebeck coefficient, electron scattering, ionized impurity, acoustic phonon, polar optical phonon, band structure, density of states, III-V semiconductors

I. INTRODUCTION

Accurate models of carrier transport are essential for describing the electronic properties of semiconductor materials, which are particularly important for clean energy applications ranging from photovoltaics to thermoelectrics to photoelectrocatalysts. There has been an increased focus on using compound semiconductors, including those that are degenerately and heavily doped, for these applications. To better understand existing materials and discover new ones, a fully predictive model that correlates electronic structure to properties is essential. Unfortunately, to the best of our knowledge, no model, based on *ab initio* calculations, currently exists to fully capture the elastic and inelastic scattering effects of charge carriers; as a result, errors arise when utilizing the current models. While an *ab initio* model will certainly improve our understanding of the carrier transport mechanism(s) in existing semiconductors, it can also aid in the search for high-performing materials by improving the accuracy of high-throughput computations^{1,2}.

There currently exist two main categories of models, based on the Boltzmann transport equation (BTE), for calculating the conductivity and Seebeck coefficient of semiconductors that are governed by band conduction. The first category of BTE-based models are commonly known as single parabolic band models, even though the treatment of the conduction band may not be explicitly parabolic. These models can be described as "semi-empirical", since experimentally measured parameters, such as the electron or hole effective mass, band gap, dielectric constant and polar optical (PO) phonon frequency, are used in closed-form expressions for the various scattering rates. Note that the overall mobility due to elastic scattering is calculated by averaging, according to Matthiessen's rule, the mobilities due to each scattering contribution. The main adjustable parameter in these models is the effective mass, which can be varied to fit the calculated transport properties to the experimental measurements. While such models often impressively capture the changes in properties over various ranges of temperature and carrier concentration, they are restricted to the materials for which experimental data are available; therefore, the predictability of such models are very limited.

There are numerous examples of models in this category^{3–7}, such as that by Ehrenreich⁶, who modeled the GaAs band structure and PO-phonon scattering by reviewing the experimental data⁶, and that by Sankey et al.⁵, who considered the effects of resonance, ionized

impurity, and polar optical phonon scattering in GaAs. In these models, all of the scattering mechanisms are commonly treated using the relaxation time approximation (RTA); here, the relaxation time is written as a power law function of energy – thus, the details of elastic and inelastic scattering (e.g., PO phonon) captured by the *ab initio* band structure are disregarded. Scattering rates, particularly inelastic ones, have already been shown to not follow such power law distributions^{3,8}, so the basic assumptions fail. Even in cases where the BTE is explicitly solved for PO phonon and the perturbation to the electronic distribution is obtained without the RTA assumption, the results are still heavily dependent on available experimental data. As an example, Miller et al.⁸ used the latter approach to calculate the mobility and Seebeck coefficient of InN samples, which had been grown by molecular beam epitaxy (MBE) and plasma assisted MBE so that all exhibited heteroepitaxial growth with linear charged dislocations; thus, these dislocations were found to be the limiting scattering mechanism.

The second category of BTE-based models relies on the ab initio band structure of the material, rather than specific experimentally measured parameters, but generally utilizes the relaxation time approximation to the BTE (BTE-RTA) as a simplification. Restrepo et al. each calculated the mobility of n-doped silicon at different electron concentrations in BTE-RTA and ab initio framework where electron-phonon interactions are treated as elastic with the electron distribution unchanged from the equilibrium Fermi-Dirac. On the other hand, the constant relaxation time approximation (BTE-cRTA) simplifies the equation even more, which enables closed form expressions for both conductivity divided by relaxation time and Seebeck coefficient. The advantage of these models is the ability to calculate properties of new materials, for which experimental data is unavailable. This type of model works well for some materials for which the relaxation time is fairly constant, as evidenced by the work of Madsen and Singh¹⁰. However, inelastic scattering mechanisms change the electron energy and directly affect the distribution. Lumping all the elastic and inelastic scattering mechanisms into a single constant and assuming an equilibrium Fermi-Dirac distribution in BTE-cRTA framework greatly damages the predictive ability of such models; as an example, transport properties in some cases are very far from experimental measurements. Furthermore, the relaxation time constant is usually determined by fitting the calculated conductivity to experimental data. It should be noted that the calculation of this constant is not necessary when calculating the Seebeck coefficient. This is due to the simplifying assumptions that the relaxation time is both a constant and direction independent¹⁰ which does not always hold. Therefore, BTE-cRTA suffers not only from inaccuracy in predicting the changes of properties with temperature or carrier concentration in many materials, but also from lack of pure predictability since it still relies on experimental data for the computation of the relaxation time.

Instead, we propose that accurate calculations of electronic transport properties, within the Boltzmann transport framework, are possible by combining relevant treatment of the elastic and inelastic scattering mechanisms with *ab initio* calculations of the electronic and phonon band structures. Ultimately, an *ab initio* theory for carrier transport will need to qualitatively and quantitatively predict trends in material properties, such as conductivity and Seebeck coefficient, as a function of temperature or carrier concentration. Validation of the theory against experimentally measured properties will thus give insight into which scattering effects are dominant.

In this paper, we present a band transport model for calculating low-field electrical drift mobility and Seebeck coefficient of n-type semiconductors which we refer to as aMoBT: ab initio Model for calculating mobility and Seebeck coefficient using Boltzmann Transport (aMoBT) equation. We then validate aMoBT by calculating the properties of two III-V semiconductors, GaAs and InN, with different carrier concentrations over various temperatures, and comparing them to experimental values as well as those calculated using the other transport models described above. We choose these materials because the ab initio band structure of GaAs is similar to those used in the earlier semi-empirical models at it can be reasonably well described with a single band model, whereas the ab initio band structure of InN and the limiting scattering mechanisms are quite different; thus, these two materials allow us to bracket the range of expected behavior of our proposed model.

II. CARRIER TRANSPORT MODEL

A. Solution to the Boltzmann Transport Equation

In order to calculate the mobility and Seebeck coefficient, we solve the Boltzmann transport equation (BTE) using Rode's iterative method^{3,8,11–17} (Appendix A 2) to obtain the electron distribution in response to a small driving force (e.g. a small electric field or a

small temperature gradient). It is important to note that we do not use the relaxation time approximation (RTA) in solving the BTE, so neither a variable nor a constant relaxation time appears in this expression. Due to the assumption of a small driving force, we aim to calculate only the linear response to the perturbation; thus, the general form of the electron distribution remains the at equilibrium Fermi-Dirac distribution. We can then write:

$$f(\mathbf{k}) = f_0[\varepsilon(k)] + xg(k) \tag{1}$$

where f is the actual distribution of the electrons, including both elastic and inelastic scattering mechanisms, f_0 is the equilibrium Fermi-Dirac distribution, x is the cosine of the angle between the small driving force and \mathbf{k} , g(k) is the perturbation to the distribution caused by the small driving force and finally $k = |\mathbf{k}|$. For the sake of simplicity, we express the conduction band as the average energy of the electrons as a function of distance, k, from the conduction band minimum (CBM) which is often at the center of the Brillouin Zone (i.e. Γ point); furthermore, we assume that the small driving force is aligned with k (i.e., x=1). Although this is similar in spirit to the isotropic band assumption, we take the anisotropy into account by averaging the energy values of the ab initio calculated band structure, ε (\mathbf{k}), as a function of k rather than explicitly including \mathbf{k} in every direction. Alternatively, if we wish to consider the directional transport properties, we can include the calculated band structure only in that specific direction. Here, we will focus on calculating and reporting the overall average mobility and Seebeck coefficient.

Our goal is to calculate the perturbation to the distribution³, g(k). In the reformulated Boltzmann transport equation shown in Equation 2, there are scattering-in, $S_i(g)$, and scattering-out, S_o , terms for inelastic scattering mechanisms. However, these terms also depend, in turn, on the electronic distribution as well as elastic scattering rates, ν_{el} . Therefore, the BTE must be solved self-consistently to obtain g(k):

$$g(k) = \frac{S_i[g(k)] - v(k)\left(\frac{\partial f}{\partial z}\right) - \frac{eE}{\hbar}\left(\frac{\partial f}{\partial k}\right)}{S_o(k) + \nu_{el}(k)}$$
(2)

where E is the low electric field and v(k) is the electron group velocity. The derivation of the BTE in the form shown in Equation 2 can be found in the literature³. The inelastic scattering mechanism that tends to dominate at room temperature is polar optical (PO) phonon scattering, for which we have provided the description of the $S_i(g)$ and S_o terms in Equations A9 and A10. The influence of inelastic scattering mechanisms on g, and therefore the overall mobility, are captured through the terms $S_i(g)$ and S_o in Equation 2, while elastic scattering mechanisms affect the overall mobility by the term ν_{el} . This term is the sum of all elastic scattering rates inside the material, it can be evaluated according to Matthiessen's rule:

$$\nu_{el}(k) = \nu_{ii}(k) + \nu_{pe}(k) + \nu_{de}(k) + \nu_{dis}(k)$$
(3)

where the subscripts el, ii, pe, de, and dis stand for elastic, ionized impurity, piezoelectric, deformation potential and dislocation scattering rates, respectively. Therefore, the effect of relevant elastic and inelastic scattering mechanisms are taken into account by explicitly solving the BTE (Equation 2) to obtain g(k).

When calculating various properties, several terms in Equation 2 will be set to zero. For a Seebeck coefficient, S, calculation, the applied electric driving force, $-\left(\frac{eE}{\hbar}\right)\left(\frac{\partial f}{\partial \mathbf{k}}\right)$, is set to zero. Only the thermal driving force, $v\left(\frac{\partial f}{\partial z}\right)$, in Equation 2 is taken into consideration when calculating the perturbation to the electron distribution³. Assuming a uniform electron concentration over the space at which a small temperature difference exists, the Seebeck coefficient is³:

$$S = \frac{k_B}{e} \left[\frac{\varepsilon_F}{k_B T} - \frac{\int k^2 f (1 - f) \left(\frac{\varepsilon}{k_B T}\right) dk}{\int k^2 f (1 - f) dk} \right] - \frac{\frac{J}{\sigma}}{\frac{\partial T}{\partial z}}$$
(4)

For a mobility calculation, the applied thermal driving force in Equation 2 is set to zero, so that only the contribution of the electric driving force is included. The mobility is:

$$\mu = \frac{1}{3} \frac{\int v(k) \left(\frac{k}{\pi}\right)^2 \left(\frac{g}{E}\right) dk}{\int \left(\frac{k}{\pi}\right)^2 f dk}$$
 (5)

Note that in Equation 5, the free electron density of states, $\left(\frac{k}{\pi}\right)^2$, has been used, which would limit its applicability in compound semiconductors. Thus, the replacement of this term by its *ab initio*-calculated counterpart would greatly improve the accuracy of the resulting mobility. Furthermore, the scalar group velocity, $v\left(k\right)$, is used since the energy is averaged as a function of distance from the Γ point. In general, we use the band structure, density of state, electron group velocity, conduction band wavefunction admixture and PO phonon frequency in calculating the mobility and Seebeck coefficient. Therefore, all of the required inputs to Equation 5 are calculated *ab initio*, which greatly enhances the predictability of the model. In other words, the main difference between our proposed carrier transport model and previous semi-empirical models^{3-6,8,11-14,18-20} is the use of *ab initio* parameters instead of

experimentally measured electron effective mass, band gap, etc., which eliminates the need for theories such as $\mathbf{k} \cdot \mathbf{p}$ to describe the nonparabolicity or anisotropy of the conduction band. Instead, for calculating the overlap integral, we express the conduction band wavefunction as a linear combination of s-type and p-type basis functions, with coefficients of a and c, respectively³. These coefficients can be directly calculated ab initio without the need to assume an s-like conduction band wavefunction (i.e., no assumption of a parabolic band).

The rates of the elastic scattering mechanisms, ν_{el} , are calculated from the electron group velocities, v, and density of states, D_S ; thus, the mobility may be calculated directly from the electronic band structure. The original form of these equations from $\mathbf{k} \cdot \mathbf{p}$ theory, and the modified equations that we propose, are listed in Table I. Note that in every equation, $\frac{\hbar k}{md(k)}$, which, in semi-empirical models, is the group velocity fitted to experiment by the band gap and effective mass of the semiconductor (included in d(k), see Table I), has been replaced by its *ab initio* counterpart, or v(k), which is calculated directly from the band structure.

As an example, the DFT-calculated density of states (DOS) can be plugged into Equation A7 to obtain the inverse charge screening length, β , in ionized impurity scattering. Furthermore, the numerator and denominator of the integrand in Equation 5 both contain the density of states of a free electron gas, $\left(\frac{k}{\pi}\right)^2$. Since this can also be calculated *ab initio* for the specific system of interest, D_S can instead be substituted in the equation for calculating the mobility and reformulated in terms of the energies, ε :

$$\mu = \frac{1}{3E} \frac{\int v(\varepsilon) D_S(\varepsilon) g(\varepsilon) d\varepsilon}{\int D_S(\varepsilon) f(\varepsilon) d\varepsilon}$$
(6)

where, again, v(k) is the electron group velocity and g is the perturbation to the electron distribution, which is calculated iteratively using Equation 2, and can be expressed both as a function of k or $\varepsilon(k)$ (i.e., the band structure).

Once the mobilities of the electrons and holes are known, the electrical conductivity can be readily calculated:

$$\sigma = ne\mu_e + pe\mu_h \tag{7}$$

where n and p are the concentration of electrons and holes, respectively, e is the absolute value of the charge of an electron and μ_e and μ_h are the mobility of electrons and holes respectively.

TABLE I. The original equations^{3,8}, based on $\mathbf{k} \cdot \mathbf{p}$ theory for elastic scattering rates and overall drift mobility, and proposed modifications, based on *ab initio* parameters, introduced in this work.

$\mathbf{k}\cdot\mathbf{p}$ theory with empirical parameters	$ab\ initio$
a $\nu_{ii}(k) = \frac{e^4 N m d(k)}{8\pi \epsilon_0^2 \hbar^3 k^3} \left[D(k) \ln \left(1 + \frac{4k^2}{\beta^2} \right) - B(k) \right]$	$\nu_{ii}\left(k\right) = \frac{e^{4}N}{8\pi\epsilon_{0}^{2}\hbar^{2}k^{2}v(k)} \left[D\left(k\right)\ln\left(1 + \frac{4k^{2}}{\beta^{2}}\right) - B\left(k\right)\right]$
$\beta^{2} = \frac{e^{2}}{\epsilon_{0}k_{B}T} \int \left(\frac{k}{\pi}\right)^{2} f(1-f) dk$	$\beta^{2} = \frac{e^{2}}{\epsilon_{0}k_{B}T} \int D_{S}(\varepsilon) f(1-f) d\varepsilon$
$ \nu_{pe}(k) = \frac{e^2 k_B T P^2 m d(k)}{6\pi \hbar^3 \epsilon_0 k} \left[3 - 6c^2(k) + 4c^4(k) \right] $	$ u_{pe}\left(k\right) = \frac{e^{2}k_{B}TP^{2}}{6\pi\hbar^{2}\epsilon_{0}v(k)}(3 - 6c^{2}\left(k\right) + 4c^{4}\left(k\right)) $
b $c^{2}(k) = 1 - \frac{1+\alpha}{2\alpha}, \ \alpha^{2}(k) = 1 + \frac{2\hbar^{2}k^{2}}{m\varepsilon_{g}}(\frac{m}{m^{*}} - 1)$	$c\left(k\right)$: obtained directly from wavefunctions
$\nu_{de}(k) = \frac{e^{2}k_{B}TE_{D}^{2}mkd(k)}{3\pi\hbar^{3}c_{el}} \left[3 - 8c^{2}(k) + 6c^{4}(k) \right]$	$\nu_{de}(k) = \frac{e^2 k_B T E_D^2 k^2}{3\pi \hbar^2 c_{el} v(k)} \left[3 - 8c^2(k) + 6c^4(k) \right]$
$\nu_{dis}\left(k\right) = \frac{N_{dis}e^{4}md(k)}{\hbar^{3}\epsilon_{0}^{2}c_{l}^{2}} \frac{1}{\left(1 + \frac{4k^{2}}{\beta^{2}}\right)^{3/2}\beta^{4}} , \frac{1}{d(k)} = 1 + \frac{m/m^{*} - 1}{\alpha}$	$\nu_{dis}\left(k\right) = \frac{N_{dis}e^{4}k}{\hbar^{2}\epsilon_{0}^{2}c_{l}^{2}v(k)} \frac{1}{\left(1 + \frac{4k^{2}}{\beta^{2}}\right)^{3/2}\beta^{4}}$
$\mu_{overall} = \frac{\hbar}{3m} \frac{\int k^3(g(k)/Ed(k))dk}{\int k^2fdk}$	$\mu_{overall} = rac{1}{3E} rac{\int v(arepsilon) D_S(arepsilon) g(arepsilon) darepsilon}{\int D_S(arepsilon) f(arepsilon) darepsilon}$
$g(k) = f(k) - f_0(k)$	$g\left(\varepsilon\right) = f\left(\varepsilon\right) - f_0\left(\varepsilon\right)$

The subscripts stand for: ii (ionized impurity), pe (piezoelectric acoustic phonon), de (deformation), and dis (charged dislocation scattering). The parameters are: m (electron mass), m^* (effective mass), ϵ_0 (low-frequency dielectric constant), ϵ_g (band gap), v(k) (electron group velocity), $D_S(\varepsilon)$ (ab initio calculated density of states), c(k) (contribution of p-type orbitals to the conduction band), β (inverse ionized impurity charge screening length), E_D (deformation potential), c_l (lattice constant), E (small electric field), and c_{el} (spherically averaged elastic constant). E(k) and E(k) are just collection of the parameters: E(k) and E(k) are just collection of the parameters: E(k) and E(k) and E(k) are just collection of the

The c(k) parameter is the contribution of the p orbital to the wavefunction of the band. In the $\mathbf{k} \cdot \mathbf{p}$ formulation, it has a closed-form expression that includes the band gap and experimental effective mass. In the ab initio formulation, this wavefunction admixture can be calculated by projecting the wavefunctions onto spherical harmonics that are nonzero within the sphere around each ion; this procedure is already implemented in the Vienna ab initio Simulation Package (VASP)^{21–24}.

It should be noted that there are fundamental differences between the model that we have presented here and those relying on the relaxation time approximation (RTA), and particularly, BTE-cRTA. Rather than simplification of the collision term in the BTE (Equation A2) through the RTA (Equation A3), we fully involve this term by considering both elastic and inelastic scattering mechanisms. It is noteworthy that the BTE-cRTA formulation only implicitly takes into account elastic and inelastic scattering mechanisms, by fitting

the overall relaxation time to experimental data with no explicit consideration of changes in electron distribution from each type of scattering mechanism. Furthermore, unlike the semi-empirical models that were described above, we use *ab initio* parameters; thus, higher predictability and little to no dependence on experimental data is achieved.

B. ab initio Parameters

The main input that is needed for the transport model is the crystal structure of the semiconductor material, from which *ab initio* parameters, such as the (optimized) lattice constant, PO phonon frequency, dielectric and piezoelectric constants, deformation potential and effective mass, can be computed.

We also need to know the Fermi level to compute scattering rates in Table I. In order to obtain the Fermi level, we calculate the carrier concentration and match it to the given concentration (input), n, according to Equation 8:

$$n = \frac{1}{V} \int_{\varepsilon_c}^{+\infty} g(\varepsilon) f(\varepsilon) d\varepsilon \tag{8}$$

Since both of the III-V semiconductors considered here are n-type, the concentration of hole carriers is negligible. The concentration of ionized impurities, N_{ii} (see Table I), is the sum of the concentration of all ionized centers regardless of the sign of their charge, since they are scatterer centers in both cases²⁵:

$$N_{ii} = N_A + N_D + \frac{N_{dis}}{c_l} \tag{9}$$

where N_D and N_A are concentration of donors and acceptors, respectively. N_{ii} can then be calculated at a given electron concentration, n, by iteratively solving the charge balance equation⁸:

$$n + N_A = N_D + \frac{N_{dis}}{c_l} \tag{10}$$

where the density of dislocations, N_{dis} , is only relevant for InN and is considered to be zero for GaAs. In both GaAs and InN, temperatures lower than 20 K need not be considered due to the deionization of shallow donors at lower temperatures, as observed experimentally²⁶. In the case of InN, electronic scattering from existing linear charged dislocations thus becomes important. The density of the dislocations, N_{dis} , can be determined from TEM images, in the units of (cm⁻²). We can thus obtain the overall density in bulk, by assuming that these

linear dislocations are uniformly developed along the c-axis. This is reflected in dividing the dislocation density by the lattice constant, c_l , in Equation 10. By doing that, we are assuming that there is one unit of positive charge (donor) per unit cell. For InN samples, according to Miller et al.⁸, one can assume full ionization of the donors, and therefore, a compensation level of one (i.e., $\frac{N_D+N_A}{n}=1$ or $N_D>>N_A$). Also, the assumption of donor or acceptor charged dislocations yields similar results⁸; therefore, we assume donor dislocations dominate here. It should be noted that we compare the calculated N_{dis} with the corresponding experimental data if available; otherwise, the limit for electronic properties at different values of N_{dis} can be calculated without the need for experimental data.

On the other hand, in a pure, epitaxially-grown, high-mobility GaAs sample with an electron concentration of $n = 3.0 \times 10^{13}$, no dislocations exist (i.e. $N_{dis} = 0$). The concentrations of donors and acceptors have been separately reported^{3,26}, so this provides validation of the accuracy of aMoBT, without needing to solve for N_{ii} . However, in the general case where the electron concentration is unknown, we can plot the mobility and Seebeck coefficient at different compensation ratios to define the limit of the transport properties, as shown in Figure 5. Therefore, it is important to note that only when comparing with experimental mobilities/Seebeck coefficients do we use experimentally measured electron concentrations; otherwise, we may calculate *ab initio* mobility or Seebeck coefficient, for example, at various electron concentrations, without any reliance on experimental data (e.g., as shown in Figure 5).

We use Brooks-Herring theory for singly-charged ionized impurity scattering²⁵, as shown in Table I. This is supported by the fact that in GaAs, oxygen impurities, O_{As}^{+1} , have been confirmed to be dominant and singly charged²⁷, while in InN, nitrogen (donor) vacancies, V_N^{+1} , are dominant and singly charged²⁸. It should be noted that the Brooks-Herring formulation is more accurate at low carrier concentrations, since at high concentrations, despite the inherent assumption of the theory, not all electrons are screened by the charge of an ionized center. More information on the Brooks-Herring ionized impurity model is available in Appendix A 2.

In order to calculate the low- and high-frequency dielectric constants, we use density functional perturbation theory (DFPT), as implemented in VASP, to determine Born effective charges, dielectric and piezoelectric tensors, including local field effects in DFT, as well as the force-constant matrices and internal strain tensors. We then subtract the ionic contribution to the static dielectric tensor to obtain the high-frequency dielectric constant^{29,30}. Furthermore, the inelastic scattering effect is strongly dependent on the longitudinal polar optical phonon frequencies, ω_{po} . These frequencies can be calculated using the Phonopy code³¹, where we identify the highest energy peak in the optical phonon density of states. It should be noted that at and around the Γ point, the phonon frequency is almost constant (Figure 8).

To calculate *ab initio* the deformation potential, E_D , we strain the system and calculate the energy of the conduction band of InN and GaAs unit cells at different volumes. Then, we approximate the deformation potential using the following equation:

$$E_D = -V \left(\frac{\partial E_{CBM}}{\partial V} \right)_T \Big|_{at \ V = V_0} \tag{11}$$

where V is the volume, E_{CBM} is the energy of the CBM and V_0 is the volume of the relaxed structure (i.e., zero pressure)^{32,33}. It should be noted that since the absolute value of E_{CBM} is a function of the volume itself, we use the difference between the energy of the first conduction band and the first valence (core) band. Furthermore, the elastic and piezoelectric constants have been already calculated *ab initio* for GaAs and InN, and are available in the literature. For GaAs, we use the values calculated by Beya-Wakata et al.³⁴, and for InN we use the values calculated by Sarasamak et al.³⁵, to obtain the piezoelectric coefficient and elastic constant used in the equations for piezoelectric scattering in Table I.

As a comparison, the electrical conductivity and Seebeck coefficient are also computed using the widely-used BTE-cRTA formulation. We choose the BoltzTraP package¹⁰, which uses Fourier interpolation of the calculated bands, and differentiate the band energies to find the group velocities of the electrons. Other than the need to fit the relaxation time to experimental measurements of the conductivity, the BoltzTraP/BTE-cRTA implementation represents an otherwise parameter-free model that can be adapted to different semiconductor materials.

III. COMPUTATIONAL METHODOLOGY

For each semiconductor material, the geometry of the unit cell is optimized, and the density of states and band structure are calculated. In the case of zinc blende GaAs and wurtzite InN, the unit cells are optimized using Kohn-Sham density functional theory (KS-

TABLE II. Structure of GaAs and InN calculated with DFT, using the GGA-PBE exchange-correlation functional. Changes in the lattice constants compared to experimental values^{42,43} upon optimization are reported below.

Compound	Space Group	$ \mathbf{a} \; (\mathring{\mathrm{A}})$	$\%$ change in $ \mathbf{a} $	$ \mathbf{c} $ (Å)	$\%$ change in $ \mathbf{c} $
GaAs	F-43m	5.75	2.17%	-	-
InN	$P6_3mc$	3.533	0.56%	5.693	0.8%

DFT)^{36,37}, as implemented in VASP. The generalized gradient approximation of Perdew, Burke, and Ernzerhof (GGA-PBE)^{38,39} is used to express the exchange-correlation potential, and Projector Augmented Wave (PAW) potentials^{40,41} are used to represent the valence wavefunctions. Information regarding the structure of these two systems and their changes upon geometry optimization have been summarized in Table II. The initial structures are obtained from the literature^{42,43}.

We then compute the electronic band structure of these materials. The energy cutoff for the plane wave basis set is set to 500 eV. The band structure is computed in line mode along seven high-symmetry k-points in the IBZ, with 20 k-points between each pair of highsymmetry points. The self-consistent density of states (DOS) calculation is performed using a $20 \times 20 \times 20$ k-point mesh, for both GaAs and InN. The non-self consistent energy calculations are performed in a special k-point mesh around the Γ point, at which the conduction band minimum (CBM) occurs in both direct band gap GaAs and InN. This k-point mesh contains a total of 10,234 points in the Irreducible Brillouin Zone (IBZ), with mesh spacing of 0.001, 0.01, or 0.1 fractional units, to completely account for band anisotropy while remaining dense enough around the Γ point to obtain accurate group velocity and effective mass values. For BTE-cRTA calculations, we used a 21×21×21 k-point mesh equivalent to 4831kpoints in IBZ to minimize the band crossing error¹⁰. We did not observe any changes in the calculated properties in larger k-point meshes. To determine the effect of presumably more accurate band structure calculations on the band curvature, effective mass, and group velocity, we have also employed the GW method. Only 941 k-points in IBZ have been used for GW calculations since it is more computationally demanding. Using more k-points does not change the calculated effective mass. The GW0 band structure calculations are performed using the maximally-localized Wannier functions (MLWFs) interpolation, as implemented in VASP and the Wannier code⁴⁴. It is very important to also show the feasibility of the ab initio model with band structures calculated using DFT+U, since the GW and hybrid functional (e.g., HSE⁴⁵⁻⁴⁷) methods are computationally demanding for complex materials with larger unit cells than GaAs. On the other hand, for InN, all methods and functionals attempted, including LDA, GGA, HSE and GW0, resulted in a band structure with zero band gap and a falsely predicted p-like conduction band. Only GGA+U⁴⁸, with U values obtained from the literature^{49,50} ($U_d = 6$ for In and $U_p = 1.5$ for N), produced a correct band structure and with a more s-like conduction band particularly around Γ point, which is consistent with the self-interaction corrected band structure reported by Furthmüller et al. 51. In the process of choosing the U value for GGA+U band structure calculations on GaAs, however, the values (U = 8 eV for both d orbitals of Ga and As) recommended by Persson and Mirbt⁵², with an emphasis on correctly obtaining the band gap and effective mass values, result in GaAs falsely becoming an indirect semiconductor, with the conduction band minima located at the L and X k-points rather than the Γ point⁵². Therefore, we have also employed effective U values of 7 eV (Ga) and 6 eV (As), for which a direct band structure is obtained. We have calculated mobilities obtained from both of these band structures and compared them with the ones obtained by the GW band structure. In order to calculate the group velocities, v(k), and the overall average effective mass, we have fitted a sixth degree polynomial to the calculated conduction band (i.e., average energy as a function of distance from Γ point or $\varepsilon(k)$ with $R^2 > 0.99$:

$$v\left(k\right) = \frac{1}{\hbar} \frac{\partial \varepsilon}{\partial k} \tag{12}$$

$$m^* = \left(\frac{1}{\hbar^2} \frac{\partial^2 \varepsilon}{\partial k^2}\right)^{-1} \bigg|_{at \ k=0}$$
 (13)

It should be noted that we do not use the value of effective mass in the proposed carrier transport model. Rather, we calculate it solely to compare with experiment and evaluate the effect of the shape of the conduction band (i.e., group velocities) calculated by various methods, such as GGA, GGA+U, and GW. Fitting polynomials to the numerically calculated conduction band and density of states results in smooth plots of mobility and Seebeck coefficient, as presented here, while preserving the values that are calculated *ab initio* with $R^2 > 0.99$ in all segments fitted. We fit these polynomials at different segments of the band structure and carefully choose only the ones that result in the maximum R^2 and minimum

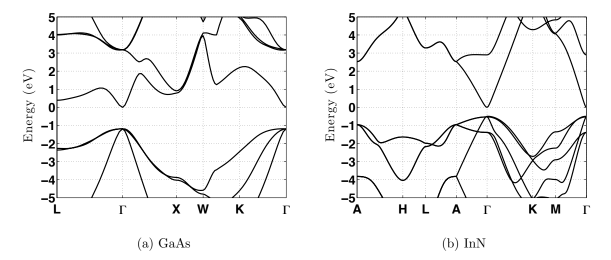


FIG. 1. Band structure of cubic GaAs and wurtzite InN, normalized so that the Fermi level is set to zero at the conduction band minimum.

discontinuity where the polynomials meet. This results in very smooth calculated group velocities, and, subsequently, other transport properties.

IV. RESULTS AND DISCUSSIONS

A. ab initio Calculated Parameter Inputs to the Transport Model

The computed band structures of GaAs and InN are shown in Figure 1. We have calculated a GW0 band structure, which starts from the wavefunctions previously computed using the GGA-PBE functional, as shown in Figure 1a.

The band structures used in previous semi-empirical models^{3,8} express the energy of the conduction band as a function of the distance from the Γ point. Instead, we calculate the *ab initio* band structure in a three-dimensional grid around the CBM, and then average the energy values of the k-points that share the same distance from the Γ point (Figure 2). For both GaAs and InN, the *ab initio* and $\mathbf{k} \cdot \mathbf{p}$ band structures agree well at small k-points; however, they diverge at larger k-points. This directly impacts the group velocity of the electrons and, ultimately, the transport properties – particularly at higher temperatures where higher energy electrons have nonzero occupation.

We have also calculated a GGA+U⁴⁸ band structure, with U values taken from the published literature^{49,50}, as shown in Figure 1. For InN, GGA+U correctly yields an s-like

TABLE III. Inputs to the transport model, as calculated *ab initio* compared to experimentally measured values. The bolded numbers are used in our transport property calculations; note that not all appear in the final expressions listed in Table I.

	GaAs		InN	
Parameter ^a	$ab\ initio$	Exp.	$ab\ initio$	Exp.
$c_l \text{ (nm)}$	0.562	0.575 ³	0.565	0.569 ³
ω_{po} (THz)	8.16	8.73^{-3}	$\boldsymbol{17.83}^{\;3}$	17.65
ϵ_0	12.18	$12.91\ ^3$	11.42	10.3^{-8}
ϵ_{∞}	10.32	$10.91\ ^3$	$\boldsymbol{6.24}$	6.7 8
E_D (eV)	6.04	8.6^{-3}	4.46	3.6^{-3}
m^*	$0.053\text{-}0.066~^{\rm b}$	$0.06360.082\ ^{53,54}$	0.062, 0.071 (GW)	$0.05 \text{-} 0.08 ^{55-60}$
$\varepsilon_g \text{ (eV)}$	0.96, 1.19 (GW)	1.424^{-6}	0.50	$0.675 \text{-} 0.7^{\ 55,57,61}$

^a The parameters are: c_l (lattice constant), ω_{po} (PO phonon frequency), ϵ_0 (low-frequency dielectric constant), ϵ_{∞} (high-frequency dielectric constant), E_D (deformation potential), m^* (effective mass), ε_g (band gap).

conduction band and a band gap of 0.5 eV, which is comparable to the self-interaction corrected band gap of 0.58 eV reported by Furthmüller et al.⁵¹ and the experimental values of $0.675\text{-}0.7 \text{ eV}^{55,57,61}$ (Table III). We include DFT+U calculations only to show the feasibility of these less-expensive methods, in the case of more complex semiconductor materials for which a GW calculations is too expensive. Also, DFT usually suffers from vastly underestimating the effective mass^{1,2}, and the introduction of the fitting parameter U may reduce the predictability of the *ab initio* model as a whole. Therefore, we stress that all reported transport properties are calculated here using the parameter-free GW band structures, unless otherwise stated.

Although we do not directly use the value of the electron effective mass in the transport property expressions, we see that the calculated effective mass of 0.062 for InN is consistent with the previously calculated effective mass (0.066) using an empirical pseudopotential⁶², and well within the range (0.05-0.08) measured experimentally⁵⁵⁻⁶⁰.

^b The GaAs effective masses are calculated as 0.053 (GGA+U, this work), 0.066 (GGA+U, with published U^{52}), and 0.063 (GW0).

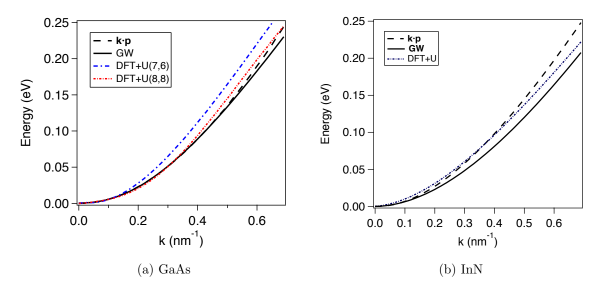


FIG. 2. The conduction bands expressed in terms of the average energy as a function of distance from the CBM (i.e., center of Brilloun zone, or Γ point), as calculated from semi-empirical expressions (in $\mathbf{k} \cdot \mathbf{p}$ formulation) versus *ab initio*. The difference at higher k values has a significant impact on transport properties, especially at high temperatures. The values of U for the d orbitals of Gallium and Arsenic, respectively, are in parentheses, while those for InN are taken from the published literature^{49,50}.

We also show the calculated phonon band structure and density of states of these two compounds in Figure 8. For GaAs, the calculated PO-phonon frequency of 8.16 THz is shown in Figure 8a. For InN, the calculated optical phonon frequency of 17.83 THz is close to the 17.65 THz value reported by Bungaro et al. 55,63 . We have listed all the parameters that are used in our transport model in Table III. We have calculated all of these parameters, as bolded in Table III, ab initio to demonstrate the feasibility of a fully predictive model for transport properties. The only exceptions are the elastic and piezoelectric constants, which are necessary to calculate the piezoelectric coefficient, P, in Table I. As described earlier, we have instead used the previously calculated values from published DFT studies for these constants 34,35 .

TABLE IV. Carrier concentrations of various experimentally fabricated and characterized GaAs samples. For the "pure" sample, data is available roughly between 5-1000 K. For the real samples, mobility data is also tabulated at different temperatures.

Sample	Concentration, $n \text{ (cm}^{-3})$	Donor, N_D	Acceptor, N_A	Reference
pure	3×10^{13}	5.2×10^{13}	2.2×10^{13}	3
a	2.7×10^{13}	4.8×10^{13}	2.1×10^{13}	26
c	7.7×10^{14}	1.1×10^{15}	3.3×10^{14}	26
e	3.1×10^{15}	4.7×10^{15}	1.6×10^{15}	26

B. Model Validation on GaAs

In order to evaluate the accuracy of aMoBT, we first calculate the mobility of three experimentally synthesized and characterized GaAs samples, as described by Stillman et al.²⁶. We also perform this analysis over a wide temperature range for high purity GaAs samples with very low electron concentrations, as labeled as "pure" in Table IV.

As shown in Figure 3a, the most accurate GW band structure results in the best agreement with experimental data. The DFT+U band structure, however, does provide us with limits of the mobility over different temperatures. When calculating the mobility and Seebeck coefficient, we calculate the Fermi level by first calculating the electron concentration through Equation 8, and then matching it to a given concentration. The calculated properties are very sensitive to the calculated Fermi level. Therefore, for comparison, we have included the results using both the ab initio DOS used in Equation 8, and the free electron DOS. As shown in Figure 3a, the ab initio model for DOS performs better for lower electron concentrations and lower temperatures, while the free electron DOS is more suitable for higher temperatures, and, particularly, at higher electron concentrations. We acknowledge that because of the log scale in Figure 3a, seeing the quantitative agreement is difficult. Therefore, we report the calculated relative error compared to the experiment for the best cases for each sample – from the ab initio DOS for sample a and from the free electron DOS for samples c and e. The minimum, maximum and the relative error in calculating the mobility of sample a are 2.25% (at 195 K), 29.42% (at 29 K), and 13.33%, respectively. These numbers are 1.02% (at 167K), 15.01% (at 49K), and 7.97% for sample **c** and 0.22% (at 195K), 7.90% (at 40K), and 4.04% for sample e. Overall, the agreement is poorer at higher electron concentrations and lower temperatures; this is attributed to the inaccuracy of the Brooks-Herring ionized impurity scattering model at high electron concentrations, as briefly described in Section II. Furthermore, the model has also been validated with the data on crystalline samples with very high purity. The calculated electron mobilities, assuming the limit that only one scattering mechanism exists at a time, along with the overall mobility, are shown in Figure 3b. The reasonable agreement between the calculated and experimental mobilities provides independent validation of the transport model. The minimum, maximum and average relative error of calculated mobility are 0.46% (at 394K), 23.55% (at 175K) and 9.53% respectively for temperatures above 20 K. The mobility is mainly limited by ionized impurity scattering at low temperatures, piezoelectric scattering at intermediate temperatures, and polar optical phonon scattering at higher temperatures (> 60 K); all of these are consistent with the previous results shown by semi-empirical models^{3,4,6} yet no experimental parameter has been used here in predicting the correct changes with the temperature and the carrier concentration.

Once we have the calculated mobility, at a given electron concentration, we can calculate the electrical conductivity of GaAs by Equation 7. For now, we assume that the carrier concentration remains constant with temperature over the range of interest. We then compare to the experimental conductivity and those values calculated using the BTE-cRTA framework, under the scenarios listed in Figure 4. As shown, not only does BTE-cRTA fail to correctly predict the trend for conductivity with temperature, but also quantitatively differs from the experimental values.

Finally, we calculate the Seebeck coefficients of the GaAs samples (assumed to be at 300 K), and compare them to the values reported previously by Rode and Knight⁴ (Figure 5). Since the data are for various samples with different electron concentration and compensation ratios, we choose various values of $N_{ii}/n = (N_D + N_A)/n$. As shown, a range of Seebeck coefficients are calculated at each electron concentration, which includes the experimentally measured points. It should be noted that not knowing beforehand the compensation and concentration of donors and acceptors, as well as their charge states, limits the overall predictability of aMoBT. However, even given these limitations, the close fit between *ab initio* and experimental properties provides independent validation of the viability of aMoBT. For further evaluation, we have calculated the Seebeck coefficient, assuming Pisarenko behavior

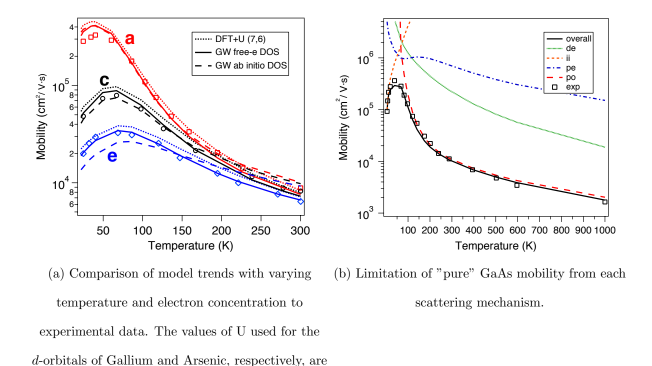


FIG. 3. The calculated and experimental^{3,64} mobility data for GaAs at various electron concentrations and temperatures. More details on the experimental data, including donor and acceptor concentrations, are available in Table IV.

in parentheses.

and compared it to aMoBT in Figure 5. We use Equation 14 with two fitting parameters: effective mass, m^* and r. It should be noted that in the case where the best agreement with experiment, through Pisarenko behavior, is only achievable by choosing either $m^* = 0.11$ or r = 0.35, both of these values are far from experimental measurements and thus lack physical meaning. Furthermore, in Figure 5 BTE-cRTA correctly predicts the change of Seebeck coefficient with carrier concentration fully *ab initio* without the need to calculate the relaxation time constant. However, the predicted values are far from the experimental results and those calculated by aMoBT. This is attributed to the treatment of $\tau(\epsilon)$ as a single constant, τ , which affect both conductivity and Seebeck coefficient when integrated over energy.

$$S \simeq \frac{k_B}{e} \left[\frac{5}{2} + r + \ln \frac{2 \left(2\pi m^* k_B T \right)^{3/2}}{h^3 n} \right]$$
 (14)

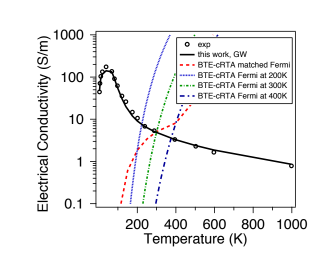


FIG. 4. Electrical conductivity of GaAs calculated using aMoBT (solid line) and the BTE-cRTA framework, and compared to experimental³ data. The Fermi level is calculated by matching the calculated carrier concentration to $n=3\times 10^{13}$. This has been done either at the mentioned temperature and kept constant over the whole temperature range, or in the case of "matched Fermi", at each temperature, the Fermi level is adjusted to the given n. The relaxation time, τ , is determined by fitting the calculated conductivity to the corresponding experimental value at 300 K. The calculated value for τ is $5\times 10^{-13}s$.

C. Model Validation on InN

In order to further evaluate the accuracy of aMoBT and its applicability to more complicated semiconductors, we also calculate the mobility and Seebeck coefficient (Figure 6) of three experimentally synthesized and characterized InN samples by Miller et al.⁸. These calculations are more challenging due to the reported presence of linear charged dislocations in the crystal structure^{8,65–67}, due to the processing conditions employed. For each sample at a given carrier concentration, as shown in Table V, we change the concentration of dislocations, N_{dis} , until the calculated mobility values match the experimental measurements. The fitted N_{dis} (Table V) is within the range of measured concentrations from transmission electron microscopy analysis (TEM)⁸, which confirms that the limiting mechanism is indeed scattering from dislocation lines.

As shown in Figure 6a and 6b, while there is an excellent agreement between the calculated and experimental mobility, the calculated Seebeck coefficients for samples B and C exhibit more pronounced changes with temperature than the experimental Seebeck coeffi-

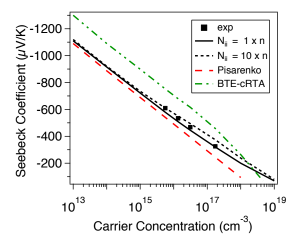


FIG. 5. Calculated (by aMoBT) and experimental⁴ GaAs Seebeck coefficient, at different ratios of the ionized impurity concentration, N_{ii} , to the electron concentration, n. For the Pisarenko plot, using Equation 14, we have used values of $r = -\frac{1}{2}$ for acoustic phonons, and $m^* = 0.063$. The Fermi levels for BTE-cRTA calculations were obtained by integrating the density of state to obtain the corresponding carrier concentrations; thus the same band structure and Fermi levels have been used to obtain the black and green lines.

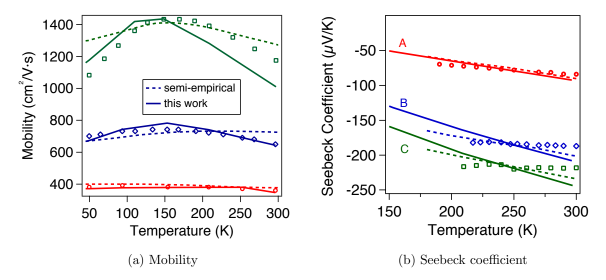


FIG. 6. Calculated (by aMoBT) and experimental transport properties of the three InN samples listed in Table V. The dashed line is calculated by the semi-empirical model used by Miller et al.⁸, while the solid line is calculated by the proposed *ab initio* transport model (aMoBT).

TABLE V. Measured⁸ and calculated InN dislocation density, corresponding to the mobility and Seebeck coefficient reported in Figures 6a and 6b.

		$N_{dis} \ (\mathrm{cm}^{-2})$		
Sample	Experimental	Semi-empirical ⁸	This work	
A	$\approx 1 \times 10^{11}$	1.5×10^{11}	8.20×10^{10}	
В	$2-5\times10^{10}$	1.5×10^{10}	1.18×10^{10}	
С	$\approx 1 \times 10^9 - 5 \times 10^{10}$	4.1×10^9	3.47×10^9	

cients. The mobility of the samples is found to be limited by charged dislocations, particularly at low temperatures. The next limiting mechanism is polar optical phonon scattering, which is more important at higher temperatures while ionized impurity scattering is more important at lower temperatures. This can be seen in Figure 7, which shows the mobility of sample B if it were limited by each type of scattering mechanism, as well as the overall mobility. These findings are in agreement with the semi-empirical transport model⁸, except that all parameters are obtained from *ab initio* calculations that require knowledge only of the crystal structure of the material. Comparing the transport properties calculated from using model with those calculated using semi-empirical models (including experimentally measured band gap and effective mass (See Table III under "Exp.") in Figure 6 shows that although quantitative agreement with experiment is slightly better with the semi-empirical model, Seebeck coefficient calculations on samples B and C, and the mobility of the sample at high temperature, show much better accuracy with the *ab initio* model presented here.

Finally, we should once again acknowledge the assumptions and limitations of the current model when applied to the other types of semiconductors. Most importantly, the formulation presented in this work is for low-field transport (particularly drift mobility and Seebeck coefficient), in which the changes to the electron distribution are merely a linear perturbation to the equilibrium Fermi-Dirac distribution; thus, the applicability of the current model for high-field transport or heavily doped and polar semiconductors where the linear BTE formulation fails⁶⁸, is very limited. Furthermore, we have averaged the energy around CBM and expressed the energy values in the band structure as a function of the absolute value of \mathbf{k} , or simply, the distance from Γ point in the reciprocal space. Therefore, the reported mobility values are averaged and the effect of band structure anisotropy is not fully

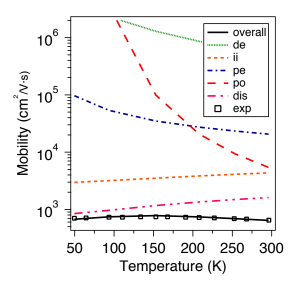


FIG. 7. Calculated and experimental⁸ values for InN mobility at $n = 9 \times 10^{17}$ cm⁻³ (sample B in Table V). Each line represents the mobility if limited only by the corresponding mechanism.

captured. It is possible, however, to include the band structure of the material only in the specific orientation of interest to account for anisotropy. Currently, the model is limited to a single conduction band. Although the single band ab initio model can be used for prediction of many direct band gap semiconductors, it will only result in an overestimation of transport properties of semiconductors with more complex band structure. This is due to the fact that currently, interband scatterings between several bands that are participating in transport are neglected. In future, we will solve coupled-BTE and take into account two and more participating bands which enables calculation of both electron and hole mobilities in more materials. Finally, although the usage of the Hubbard U parameter in the band structure calculation might limit the predictability of the model in calculating overall transport properties, this can be properly addressed by using more accurate methods of band structure calculations as reported here. We include DFT+U calculations here only to show the feasibility of working with the model when GW or other less commonly used methods are not technically or otherwise feasible.

V. CONCLUSIONS

We have presented an *ab initio* transport model for calculating the electrical mobility and Seebeck coefficient of n-type semiconductors using Boltzmann transport (aMoBT) equation.

By using the inputs from density functional theory calculations, and considering all relevant physical phenomena (i.e., elastic and inelastic scattering mechanisms), we have successfully calculated highly-accurate transport properties of GaAs and InN over various ranges of temperature and carrier concentration. aMoBT provides both qualitative and quantitative improvements in accuracy compared to the widely-used semi-empirical and constant relaxation time approximation model solutions to the Boltzmann transport equation. Future work will focus on extending this model to p-type semiconductors.

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Appendix A: Boltzmann Transport Equation

The Boltzmann transport equation (BTE) describes the non-equilibrium behavior of charge carriers (e.g., electrons or holes) by statistically averaging over all possible quantum states. For the electron distribution, f, this is represented by the BTE:

$$\frac{df\left(\mathbf{k}, T, t\right)}{dt} = \left(\frac{\partial f\left(\mathbf{k}, T, t\right)}{\partial t}\right)_{s} - \frac{d\mathbf{k}}{dt} \cdot \nabla_{k} f\left(\mathbf{k}, T, t\right) - \mathbf{v}\left(\mathbf{k}\right) \cdot \nabla_{r} f\left(\mathbf{k}, T, t\right) \tag{A1}$$

where f is a function of state \mathbf{k} , temperature T, and time t, and $\mathbf{v}(\mathbf{k})$ are the electron group velocities. The three terms on the right-hand side of Equation A1 refer, respectively, to the temporal rate of change of f due to all scattering processes, rate of change of f due to external forces, and diffusion from the carrier density gradient.

If the external forces consist only of a low electric field, **E**, and no magnetic field, **B**, such that $\frac{d\mathbf{k}}{dt} = \frac{e\mathbf{E}}{\hbar}$, then the low-field BTE becomes:

$$\frac{df(\mathbf{k}, T, t)}{dt} + \mathbf{v}(\mathbf{k}) \cdot \nabla_r f(\mathbf{k}, T) + \frac{e\mathbf{E}}{\hbar} \cdot \nabla_k f(\mathbf{k}, T) = \left(\frac{\partial f(\mathbf{k}, T, t)}{\partial t}\right)_s \tag{A2}$$

1. Constant Relaxation Time Approximation

Furthermore, f can be described as a first-order (linear) perturbation, $g(\mathbf{k})$, from the (equilibrium) Fermi-Dirac distribution, f_0 , due to scattering:

$$\left(\frac{\partial f\left(\mathbf{k}, T, t\right)}{\partial t}\right)_{s} = -\frac{f\left(\mathbf{k}\right) - f_{0}\left(\mathbf{k}\right)}{\tau} = -\frac{g\left(\mathbf{k}\right)}{\tau} \tag{A3}$$

$$f_0\left[\varepsilon\left(\mathbf{k}\right)\right] = \frac{1}{e^{\left[\varepsilon\left(\mathbf{k}\right) - \varepsilon_F\right]/k_B T} + 1} \tag{A4}$$

where the dependence of ε on **k** is given by the electronic band structure, and the various scattering terms and time dependence are lumped into the electronic relaxation time, τ .

If τ is a constant, then this major simplification results in the BTE-cRTA. This assumption simplifies the theory to an extent that closed form expressions for conductivity and Seebeck coefficient can be obtained¹⁰. In this approach, the details of all elastic and inelastic scattering mechanisms are lumped into the relaxation time constant, τ . While popular, this approach suffers from the following disadvantages: 1. τ is obtained by fitting to the experimental data for the conductivity of the material, which limits the predictability of the model, and 2. Due to oversimplification of the transport mechanism, it may result in incorrect values and even incorrect trends with temperature or carrier concentration, as illustrated

in Figure 4. Therefore, by explicitly including all possible electronic scattering mechanisms, one can determine which mechanisms are physically relevant for a given semiconductor.

2. Explicit Solution of Linear BTE

To go beyond the relaxation time approximation, both elastic scattering mechanisms, for which the kinetic energy of the electrons remains constant, and inelastic scattering mechanisms, for which there is a change in the electron distribution, should be taken into account. If the system is governed only by elastic scattering mechanisms, the relaxation time, τ , is equal to the inverse of the overall elastic scattering rates, which is the sum of all individual rates. Evidently, τ is not constant but does depend on energy; however, it does not necessarily follow a power law dependence (e.g., in InN⁸). However inelastic scattering mechanisms also limit the mobility, and therefore, the conductivity, of the semiconductor; as an example, polar optical (PO) phonon scattering is the main electron-phonon interaction that limits mobility at high temperatures in GaAs. Thus, we need to first calculate the perturbation, g, to the electron distribution due to elastic and inelastic scattering mechanisms, and then integrate g over all states to obtain the mobility. Details on this approach are given below.

The most relevant elastic scattering mechanism for compound semiconductors is expected to be ionized impurity scattering at low temperatures. Ionized impurity scattering occurs when a charged center is introduced inside the bulk material. As a result of Coulombic interactions between the electron and ion, electrons scatter to different states (i.e., become distracted). The ionized impurity scattering rate, ν_{ii} (i.e., a component of the overall ν), may be expressed using Brooks-Herring theory²⁵:

$$\nu_{ii} = \frac{e^4 N}{8\pi \epsilon_0^2 \hbar^2 k^2 v} \left[D \ln \left(1 + \frac{4k^2}{\beta^2} \right) - B \right] \tag{A5}$$

where the charge screening potential, ϕ , is obtained by solving Poisson's equation:

$$\phi = \frac{q}{4\pi\epsilon_0 r} \exp\left(-\beta r\right) \tag{A6}$$

and inverse screening length, β , is given by:

$$\beta^{2} = \frac{e^{2}}{\epsilon_{0}k_{B}T} \int D_{S}(\varepsilon) f(1-f) d\varepsilon$$
 (A7)

where f is the electron distribution and ϵ_0 is the low-frequency dielectric constant. Details on the α , D and B parameters are given in the literature³.

At high temperatures, after an inelastic (e.g., electron phonon) scattering event, where the electron scatters from momentum state k to k', the energy of an electron changes, and hence, the electron distribution also changes. (Note that the distribution may also be perturbed by external forces, such as an electric field or temperature gradient.) Thus, f becomes a function of k, so it must be mapped via the electronic band structure, $\varepsilon(k)$. This effect can be shown as a deviation from Fermi-Dirac behavior (Equation A3). After some mathematical manipulation, for which details can be found in the literature³, the BTE can be reformulated as:

$$g = \frac{S_i(g') - \nu \left(\frac{\partial f}{\partial z}\right) - \frac{eF}{\hbar} \frac{\partial f}{\partial k}}{S_o + \nu_{el}} \tag{A8}$$

$$S_{i}(g') = \int dk' X g(k') \left[s_{inel}(k', k) \left[1 - f(k) \right] + s_{inel}(k, k') f(k) \right]$$
(A9)

$$S_{o} = \int dk' \left[s_{inel}(k, k') \left[1 - f(k') \right] + s_{inel}(k', k) f(k') \right]$$
(A10)

Detailed integrated expressions for the scattering in, S_i , and scattering out, S_o , terms are available in the literature³. The reformulated BTE can then be solved iteratively, using Rode's method^{3,8,11–14} since $S_i(g')$ and f themselves are functions of g. First, the Fermi-Dirac distribution can be plugged into the right-hand side of Equation A8 to obtain the first guess, g_1 , which in turn is used to obtain a new electron distribution to solve for the next guess, g_2 ; this process continues until g converges to a unique value. Typically, five iterations are required for the perturbation to converge for polar optical phonon scattering in GaAs and InN. More details on Equations A8-A9 are available in the literature³.

3. Phonon Dispersion

Polar optical phonon scattering originates from interactions between electrons and high-frequency optical phonons. They provide the dominant inelastic electron scattering mechanism near (and above) room temperature in compound semiconductors. This is attributed to the high energies of optical phonons being comparable to k_BT at high temperatures. The scattering rates themselves are strongly dependent on the polar optical phonon frequencies. ω_{po} . These frequencies can be calculated using the Phonopy code³¹ which solves for dynamical matrix from the force constants calculated using density functional perturbation theory (DFPT), as implemented in VASP. The phonon band structures for GaAs and InN are shown in Figure 8.

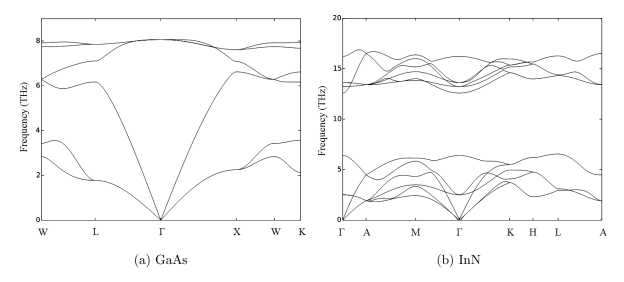


FIG. 8. Phonon band structures of InN and GaAs calculated by Phonopy³¹.

Appendix B: Sensitivity analysis

1. Sensitivity to the calculated dielectric constants

We have done sensitivity analysis for the calculated mobility of GaAs pure sample at different dielectric constants. As shown in Figure 9, the result is sensitive to dielectric constants at low and high temperatures but much less sensitive at temperatures in 100-200K range. Inaccurate calculation of dielectric constant in this case can result in up to -41% (at 40 K) calculated value of mobility compared with measured values in case of -20% from the base value for dielectric constant and it can go up to +43% (at 5K) for +20% from the base value. The base values are the ones reported in Table III calculated from *ab intitio* assuming the relaxed structure. This shows the importance of accurate calculation of these constants at least with 5-10% accuracy.

2. Sensitivity to the lattice constants

We also applied $\pm 3\%$ strain to the lattice constant of the relaxed GaAs and recalculated the band structure, DOS and optical phonon frequencies to see how sensitive is the calculated mobility with respect to the crystal structure. We assume that everything else is kept constant according to the base case (see Table III). According to Figure 10, the calculated mobility is extremely sensitive to the crystal structure. This is mainly due to the impact that

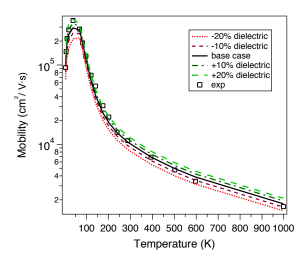


FIG. 9. Sensitivity analysis of the mobility of GaAs pure sample (see Table IV). We changed here only the static, ε_s , and high frequency, ε_{∞} , dielectric constant from -20% to +20% of the base case values reported in Table III. The results are sensitive to dielectric constant at low and high temperatures.

the structure has on the band shape (i.e. group velocity of the electrons) since the mobility at any temperature is affected. For example, the GW band structure of -3% strained GaAs gives the effective mass of 0.026 while that of +3% strained GaAs gives the effect of 0.10. Both of these values are well outside of the range of the reported experiemntal values 0.064-0.082 (see Table III). Also, these strained structures are extremely unlikely to be relaxed with any functional since their built in pressure with GGA-PBE functionals are already 10.66 kB and -74.77 kB while the relaxed structure that we have calculated and reported in Table II has a built in pressure of only -0.3 kB. Nevertheless, Figure 10 shows the importance of accurate calculation of the crystal structure and subsequently the band structure (i.e. group velocities).

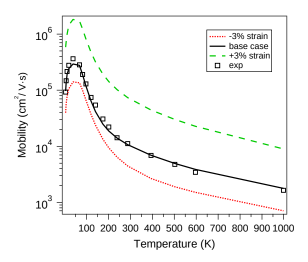


FIG. 10. Sensitivity analysis of the mobility of GaAs pure sample (see Table IV) calculated by aMoBT. We changed here the crystal structure and subsequently the newly calculated optical phonon frequencies. The calculated mobility is sensitive to the strain at all temperatures.

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