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Ab initio study of electron-phonon interaction in phosphorene

Bolin Liao,1 Jiawei Zhou,1 Bo Qiu,1 Mildred S. Dresselhaus,2,3,* and Gang Chen1,†

1Department of Mechanical Engineering, Massachusetts Institute of Technology, Cambridge, Massachusetts, 02139, USA
2Department of Electrical Engineering and Computer Science, Massachusetts Institute of Technology, Cambridge, Massachusetts, 02139, USA
3Department of Physics, Massachusetts Institute of Technology, Cambridge, Massachusetts, 02139, USA

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The monolayer of black phosphorus, or “phosphorene,” has recently emerged as a two-dimensional semiconductor with intriguing highly anisotropic transport properties. Existing calculations of its intrinsic phonon-limited electronic transport properties so far rely on the deformation potential approximation, which is in general not directly applicable to anisotropic materials since the deformation along one specific direction can scatter electrons traveling in all directions. We perform a first-principles calculation of the electron-phonon interaction in phosphorene based on density functional perturbation theory and Wannier interpolation. Our calculation reveals that (1) the high anisotropy provides extra phase space for electron-phonon scattering, and (2) optical phonons have appreciable contributions. Both effects cannot be captured by the deformation potential calculations. Our simulation predicts carrier mobilities ~170 cm²/V s for both electrons and holes at 300 K, and a thermoelectric figure of merit $zT$ of up to 0.14 in $p$-type impurity-free phosphorene at 500 K.

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I. INTRODUCTION

Twenty years ago, Hicks and Dresselhaus predicted that low-dimensional conductors could have better thermoelectric performance compared with their three-dimensional bulk counterparts, mainly owing to the quantum confinement effect [1,2]. In particular, the electronic density of states in low-dimensional systems usually exhibits sharp changes with respect to the carrier energy, which is of significant benefit for improving the Seebeck coefficient [3]. In the past two decades, researchers have widely utilized the much-advanced nanotechnology to boost the thermoelectric performance using various approaches [4,5]. Experimentally, artificial low-dimensional structures, such as quantum dots [6], quantum wells [7], superlattices [8,9], a two-dimensional electron gas [10], and nanowires [11,12], have all been studied for enhanced thermoelectric properties. These structures, however, are difficult to scale up due to the complexity of the fabrication process and the high cost.

Immediately after the first successful isolation of graphene [13], a stable monolayer of carbon, the look into natural low-dimensional conductors for good thermoelectrics started, in the hope that these materials are easier to obtain at a lower cost [14–16]. Unfortunately, graphene itself turns out to be a not-so-good thermoelectric material, due to (1) the high electron-hole symmetry and the absence of a band gap, which results in large detrimental bipolar conduction; (2) the linear dispersion for low-energy excitations, which leads to a smooth quadratic density of states without the preferred sharp features; and (3) ultrahigh lattice thermal conductivity [17,18], in part coming from the large contribution of the less-scattered flexural phonon mode [19] (there have been theoretical works suggesting that the classical size effect in nanostructured graphene can largely reduce its lattice thermal conductivity [20,21]). Along these lines, natural two-dimensional materials with a sizable band gap, quadratic low-energy dispersion, and suppressed flexural phonon modes have been sought as better candidates for thermoelectrics, in addition to the generally preferred high carrier mobility. Some subsequently synthesized or isolated two-dimensional materials usually fit some of the above criteria, but for most cases also possess serious drawbacks. For example, a monolayer of transition-metal dichalcogenide MoS₂ comes with a band gap while being limited by its relatively low carrier mobility [22,23]; silicene and germanene, monolayers of silicon and germanium atoms arranged in honeycomb lattices, possess similar low-energy electronic structures as that of graphene, with very small band gaps (a few meV), which only arise from spin-orbit coupling [24,25].

Recently, another member of the two-dimensional-material family, single layers of black phosphorus dubbed “phosphorene,” has emerged and attracted intense research interest [26–34]. In a phosphorene layer, phosphorus atoms are arranged in a puckered honeycomb lattice [35] with low symmetry and high anisotropy. This hingelike puckered structure leads to intriguing mechanical properties, such as a negative Poisson ratio [36]. The resulting electronic structure is also highly anisotropic, with a fundamental band gap of 2 eV [37] that can be potentially tuned either by changing the number of layers [26], controlling the edge termination and the width of a ribbon [38], or imposing a strain [39,40]. The low-energy dispersion is quadratic with very different effective masses along armchair and zigzag directions [37] for both electrons and holes. This anisotropic electronic structure is useful for thermoelectric materials, since in the direction with a smaller effective mass, the carrier mobility and thus the electrical conductivity can be high, while the larger effective mass along the other direction contributes to an overall large density of states that improves the Seebeck coefficient. Moreover, few-layer black phosphorus has been experimentally found to exhibit high carrier mobility, especially for holes [26–28,41,42], while theoretical calculations on single-layer
phosphorene have suggested even higher values [37,43], and the possible tunability via applying a strain [44,45]. These features have stimulated lots of research efforts in evaluating the potential thermoelectric performance of phosphorene. Lv et al. calculated the thermoelectric power factor of phosphorene [46], and further showed that strain helps improve the thermoelectric performance of phosphorene via inducing band convergence [47]. Qin et al. simulated the lattice thermal conductivity of phosphorene from first principles [48], showed that the thermal transport was also highly anisotropic, and revealed that the much-reduced lattice thermal conductivity compared with graphene could be largely attributed to the suppressed flexural mode. Jain et al. refined the lattice thermal conductivity calculation using the full iterative solution of the Boltzmann transport equation [49]. Fei et al. pointed out that the directions with higher electrical and thermal conductivity, respectively, in phosphorene are orthogonal to each other, which leads to a promising thermoelectric figure of merit along the armchair direction, exceeding 2 at 500 K [37].

Although the aforementioned works have studied the thermoelectric properties of phosphorene in some detail, the treatment of electron-phonon interaction in phosphorene has been limited to the constant relaxation time approximation for calculating Seebeck coefficient, and the deformation potential approximation for calculating the electrical conductivity. In particular, the existing deformation potential calculations [37,43] obtained separate deformation potentials for different transport directions by deforming the lattice along that direction. The validity of such an approach is questionable because the deformation along one direction will scatter electrons going in all directions. In this paper, we study the electron-phonon interaction in phosphorene fully from first principles. We find that the deformation potential calculations tend to overestimate the carrier mobility, and we fully assess the potential of phosphorene as a thermoelectric material based on our simulation results.

II. METHODS

We first carry out the standard density functional theory calculation of the electronic structure of phosphorene after obtaining a fully relaxed crystal structure using the QUANTUM ESPRESSO package [50] and a norm-conserving pseudopotential with the Perdew-Wang exchange-correlation functional within the local density approximation [51]. We use 200 × 200 k mesh and the triangular integration method [52] to generate an accurate electronic density of states. The phonon dispersion and the electron-phonon scattering matrix elements are calculated within density functional perturbation theory [53], initially on a coarse 10 × 10 × q mesh, and then along with the electronic structure on a coarse 10 × 10 k mesh, are interpolated using the EPW package [54,55] to a dense 300 × 300 k mesh covering half of the Brillouin zone centered around the Γ point and 300 × 300 q mesh in the full Brillouin zone using maximally localized Wannier functions [56] for calculating the electronic relaxation time due to electron-phonon interaction, which is given by Fermi’s “golden rule” [57] as

\[
\frac{1}{\tau_{ep}(\mathbf{k})} = \frac{2\pi}{\hbar} \sum_{\mathbf{k}'} |\langle \mathbf{k}' | \partial_{\mathbf{q}} V | \mathbf{k} \rangle|^2 \left( (f_{\mathbf{k}'} + n_{\mathbf{q}}) \delta(E_{\mathbf{k}} - E_{\mathbf{k}'} + \hbar\omega_{\mathbf{q}}) \delta_{\mathbf{k} + \mathbf{q}, \mathbf{k}'} + (1 + n_{\mathbf{k}'} - f_{\mathbf{k}}) \delta(E_{\mathbf{k}} - E_{\mathbf{k}'} - \hbar\omega_{\mathbf{q}}) \delta_{\mathbf{k} - \mathbf{q}, \mathbf{k}'} \right) \left( 1 - \frac{\mathbf{v}_{\mathbf{k}'} \cdot \mathbf{v}_{\mathbf{k}}}{|\mathbf{v}_{\mathbf{k}'}||\mathbf{v}_{\mathbf{k}}|} \right),
\]

where \( \mathbf{k}, \mathbf{k}', \text{ and } \mathbf{q} \) are wave vectors of the initial and final electronic states and the participating phonon state; \( E_{\mathbf{k}}, E_{\mathbf{k}'}, \text{ and } \hbar\omega_{\mathbf{q}} \) are their energies; \( f_{\mathbf{k}', f_{\mathbf{k}}, n_{\mathbf{k}}}, \text{ and } n_{\mathbf{q}} \) are their equilibrium distribution functions; \( \langle \mathbf{k}' | \partial_{\mathbf{q}} V | \mathbf{k} \rangle \) is the electron-phonon matrix element; \( \mathbf{G} \) is a reciprocal lattice vector; and \( \mathbf{v}_{\mathbf{k}} \) and \( \mathbf{v}_{\mathbf{k}'} \) are the group velocities of the initial and final states. The factor \( (1 - \frac{\mathbf{v}_{\mathbf{k}'} \cdot \mathbf{v}_{\mathbf{k}}}{|\mathbf{v}_{\mathbf{k}'}||\mathbf{v}_{\mathbf{k}}|}) \) takes into account the momentum loss in the scattering processes, and the thus defined relaxation time is usually named “the momentum relaxation time” and used for calculating transport properties [58,59]. The summation in Eq. (1) is performed using the triangular method [52] to improve the convergence, and eliminate the need of choosing the Gaussian broadening parameter when doing the summation using Gaussian functions to approximate the delta functions. The calculated electronic relaxation times are then plugged into the standard formulas of the transport properties based on the Boltzmann transport equation [60]. We use the following formulas for calculating transport properties (along a certain direction, denoted as \( x \)):

\[
\sigma_{xx} = \frac{1}{NA} \sum_{\mathbf{k}} -e^2 v_x^2 \tau_{ep} \frac{\partial f_{\mathbf{k}}}{\partial E_{\mathbf{k}}},
\]

where \( \sigma_{xx} \) is the electrical conductivity, \( S_{xx} \) is the Seebeck coefficient, \( \kappa_{xx} \) is the zero-current electronic thermal conductivity, \( N \) is the total number of \( k \) points in the full Brillouin zone, \( A \) is the area of a unit cell, \( e \) is the electron charge, \( v_x \) is the group velocity along the \( x \) direction, \( \tau_{ep} \) is the electron-phonon relaxation time from Eq. (1), and \( T \) is temperature. The figure of merit \( zT \) is defined as \( (S^2\sigma T)/\kappa \). The whole process is parameter-free and has been applied to study the thermoelectric transport properties of silicon by the authors with remarkably good agreement with experiments [61,62]. A similar calculation scheme has been applied by other researchers to studying the electron-phonon interactions in graphene [63,64]. The power of this \textit{ab initio} approach lies in the fact that individual carrier states can be analyzed separately in terms of the time and length scales associated
with their transport, given by the electron-phonon relaxation
time $\tau_{\text{eph}}$ and the carrier mean free path $l_{\text{mfp}} \equiv \tau_{\text{eph}} |v_{\text{th}}|$ that character-
izes the average length a carrier can travel before getting
scattered by a phonon, respectively. This information provides
crucial guidance in designing nanostructured thermoelectric
materials, as will be discussed in a later section.

III. RESULTS AND DISCUSSIONS

For brevity, the lattice parameters, the electronic structure,
and phonon dispersion are not shown here, since our results
are essentially the same as previous reports. The electronic
band gap is underestimated to be 0.8 eV, and is a well-known
problem of density functional theory. The more accurate band
gap of 2 eV from GW calculation [37] is imposed in the
following calculations by rigidly shifting the bands. Figure 1
shows the electronic density of states, where well-defined
steplike features specific to two-dimensional quadratic bands
are observed, as well as quasi-one-dimensional peaks near the
band edges as a result of the high anisotropy, since both the
lowest conduction band and the highest valence band are very
flat along the zigzag direction, resembling one-dimensional
bands along the armchair direction, which signals a high
Seebeck coefficient. This feature is reminiscent of the quasi-
two-dimensional bands in good bulk thermoelectrics, such as
PhTe and PbSe [65].

We proceed to calculate the scattering rates and mobility
for free carriers in phosphorene limited by electron-phonon
interactions. We exclude contributions from the flexural
phonon modes for two reasons: (1) out-of-plane vibrations do
not contribute to the first-order electron-phonon interactions
in two-dimensional materials due to the inversion symmetry
with respect to the material plane, as in the case of graphene
[64, 66] (although the flexural phonon modes in phosphorene
have small in-plane components as well, their contributions are
negligibly small); (2) so far in most experiments phosphorene
samples are studied on a substrate, by which the flexural
phonons will be largely suppressed. The scattering rates are
presented in Fig. 2, and compared with the deformation
potential calculation using parameters from Qiao et al. [43]. At
each electronic energy, electrons with different wave vectors
in general have different scattering rates due to the direction
dependence of the electron-phonon scattering, and thus the
plotted scattering rate is multivalued at each electron energy.
Although the deformation potential approximation is not
rigorously applicable in this case due to previously mentioned
reasons, in general it gives reasonable estimates of the average
strength of electron-acoustic-phonon interactions, except for
the case of holes in the zigzag direction, where the predicted
scattering rate using the deformation potential approximation
is three orders of magnitude lower than our result, and a hole
mobility of 26 000 cm$^2$/Vs was predicted accordingly [43].
Our results indicate that the contributions from optical phonons
are not negligible, especially for carriers slightly away from the
band edges. More importantly, we observe peak structures of
the scattering rates near the band edges, similar to the density
of states. This can be explained by the high anisotropy of the band
structure. According to Eq. (1), the scattering rate of a specific
carrier state depends on the available phase space for the final
states. In other words, the large number of carrier states along
the zigzag direction provides a large number of available final
states for carriers traveling along the armchair direction to be
scattered to. In this way the carrier transport along the two
directions is coupled through the electron-phonon interaction
and the electron-phonon scattering rates follow the trend of
the total density of states.

The peak structures of the scattering rates near the band
edges are expected to have a major impact on the carrier
mobility since carriers near the band edges contribute the most
to the transport. We show here in Fig. 3 the carrier mobilities
of electrons and holes along armchair and zigzag directions at
300 K, with respect to the carrier concentration. We simulate
the effect of carrier concentration by rigidly shifting the Fermi level, assuming the electronic band structure is not
greatly affected by free carriers. We predict the phonon-limited
carrier mobility of phosphorene is $\sim 170$ cm$^2$/Vs for both
electrons and holes along the armchair direction at 300 K and
a moderate anisotropy between the two transport directions.
Experimentally Xia et al. [28] measured the hole mobility of
a 15-nm-thick (~30 atomic layers) black phosphorus sample
to be $\sim 600$ cm$^2$/Vs and that of an 8-nm-thick (~15 atomic
layers) sample to be $\sim 400$ cm$^2$/Vs, and more recently Xiang
et al. [42] measured the hole mobility of a 4.8-nm-thick
(~8 atomic layers) sample to be $\sim 200$ cm$^2$/Vs, all along the
armchair direction. The decreasing trend of the hole mobility
with decreasing number of atomic layers was previously
attributed to the sample quality degradation [28]. As shown
by Qiao et al. [43], the hole effective mass along the zigzag
direction increases from 0.89 to 6.35 as the number of atomic
layers decreases from five to one, while the hole effective
mass along the armchair direction stays unchanged. As we
discussed in the previous section, the high anisotropy and
large effective mass along the zigzag direction provides large
phase space for the electron-phonon scattering processes and

![Figure 1](235419-3.png)

**FIG. 1.** (Color online) The electronic density of states of phosphorene. Steplike features characteristic to two-dimensional quadratic bands are present, as well as quasi-one-dimensional peaks, which reflect the fact that both the conduction band and valence band are very flat along the zigzag direction, resembling one-dimensional electronic bands. The arrows highlight the one-dimensional peak structures.
gives rise to the peak structures of the scattering rates near the band edges. Therefore our finding suggests that the drastically increased anisotropy with decreasing number of atomic layers [43] could also contribute to the observed reduction of the carrier mobility, and thus the single-layer phosphorene may not be the best option for high-mobility applications.

In Fig. 4 we show the calculated Seebeck coefficient and thermoelectric power factor along the armchair direction for both $p$-type and $n$-type phosphorene, at various carrier concentrations and temperatures. In calculating the electrical conductivity, the thickness of the phosphorene sheet is chosen as the interlayer distance in bulk phosphorus, 0.55 nm [37]. Although this conventional choice seems somewhat arbitrary, it will not affect the thermoelectric figure of merit $zT$ because the same factor appears in the thermal conductivity as well. Owing to the special features of the electronic structure mentioned above, the Seebeck coefficient is high, and the thermoelectric power factor reaches $\sim 70 \mu W/cm K^2$ in $p$-type phosphorene at room temperature. This number is comparable to that in state-of-the-art bulk thermoelectric materials, such as BiTeSb alloy [67] (the arbitrariness of choosing the film thickness may render this comparison unfair, to some extent).

With the recently reported phonon thermal conductivity of phosphorene calculated from first principles [48,49], we have all the ingredients for calculating the thermoelectric figure of merit $zT$ (here we used data from Ref. [49], where the full iterative solution of the Boltzmann equation was used instead of the relaxation time approximation), and the results are shown in Fig. 5 for $p$-type and $n$-type phosphorene along the armchair direction at temperatures up to 500 K. The optimal $zT$ is $\sim 0.06$ at 300 K and $\sim 0.14$ at 500 K for $p$ type, with the optimal carrier concentration around $5 \times 10^{12} \text{cm}^{-2}$ at 300 K and $8 \times 10^{12} \text{cm}^{-2}$ at 500 K. These values are for impurity-free phosphorene and should be regarded as an upper limit for the thermoelectric performance of phosphorene. A common strategy to improve the thermoelectric performance is to introduce nanoscale grain boundaries or precipitates [67,68].

![FIG. 2.](image-url) The calculated electron-phonon scattering rates (blue dots for acoustic-phonon scattering and orange dots for optical phonon scattering) for (a) electrons and (b) holes, compared to the deformation potential approximation (DPA) results (dash and dash-dot lines). The black arrows mark the peak structures of the scattering rates near the band edges.

![FIG. 3.](image-url) The calculated carrier mobility for electrons (solid line) and holes (dashed line) (a) along the armchair direction and (b) along the zigzag direction at different temperatures; all are plotted versus the carrier concentration.
FIG. 4. (Color online) The calculated (a) Seebeck coefficient and (b) thermoelectric power factor for $p$-type phosphorene and (c) Seebeck coefficient and (d) thermoelectric power factor for $n$-type phosphorene versus the carrier concentration along the armchair direction at different temperatures.

FIG. 5. (Color online) Thermoelectric figure of merit $zT$ versus the carrier concentration for (a) $p$-type and (b) $n$-type phosphorene along the armchair direction at different temperatures, limited by the electron-phonon scattering.
to strongly scatter phonons and suppress the thermal conductivity. The success of this nanostructuring approach relies on the separation of the transport length scales of electrons and phonons: In typical thermoelectric materials, the phonon mean free paths [69] are much longer than the electron mean free paths [61], so that nanostructures with characteristic sizes in between can effectively block the phonon flow while leaving the electrons intact. Therefore, whether this nanostructuring approach is effective for a certain material and what should be the proper length scale of the nanostructures depends on the detailed information of the mean free path distributions of electrons and phonons in that material. To evaluate the potential effectiveness of the nanostructuring approach [67] for further improving $zT$ of phosphorene, we calculate the accumulated contribution to the transport properties along the armchair direction from individual carrier states with respect to their mean free paths, as shown in Fig. 6, where we choose the optimal carrier concentration $5 \times 10^{12}$ cm$^{-2}$ at 300 K. The calculated accumulated contribution represents fictitious values of the transport properties if carriers with mean free paths longer than a certain value are strongly suppressed by nanostructures (removed from the calculation), which is then normalized by the corresponding bulk values. In doing so, we first sort the carrier states with respect to their mean free paths, and then calculate the transport properties using Eqs. (2)–(4), but only including in the summations carrier states with mean free paths below a certain value. For example, from Fig. 6(a), we can tell that holes with mean free paths below 6 nm contribute to about 40% of the total electrical conductivity and the power factor, and about 25% to the electronic thermal conductivity. It estimates the relative effectiveness of nanostructures with a characteristic size in affecting the transport properties. Figure 6 indicates that the major contribution to the transport comes from carriers with mean free paths below 10 nm at 300 K, or in other words, nanostructures with characteristic sizes above 10 nm do not have a strong impact on electron transport in phosphorene. Since the phonon thermal conductivity has contributions from phonons with mean free paths up to 1 μm [48,49], nanostructures with a characteristic size of $\sim$10 nm can significantly reduce the phonon thermal conductivity (down to about 1 W/mK according to Ref. [49]) and preserve the electronic properties. In this ideal case the figure of merit $zT$ at 300 K can be improved to around 1 in $p$-type phosphorene along the armchair direction (in reality phonons with mean free paths longer than 10 nm still conduct some heat, just with shorter mean free paths). We note here that a more realistic evaluation of the effectiveness of the nanostructuring approach is to study the scattering of phonons and electrons with the grain boundaries using, for example, the atomic Green’s function method [70]. In practice, however, since the atomic details of the grain boundaries can vary in numerous ways, a first-principles treatment has not been practical so far. Nevertheless, the mean-free-path distributions provide practical guidance for experimentalists in designing nanostructures with proper length scales to achieve improved thermoelectric performance of phosphorene.

IV. CONCLUSION

In summary, we study the potential thermoelectric performance of phosphorene via first-principles calculation of the electron-phonon interactions. Our calculation finds that previous deformation potential calculations overestimate the carrier mobility due to high anisotropy and contributions from optical phonons. We further calculate the figure of merit $zT$ for phosphorene and the carrier mean-free-path distribution, which, in comparison to the phonon mean-free-path distribution, indicates that nanostructuring can effectively enhance the thermoelectric performance of phosphorene.

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