Piezoelectric Electron-Phonon Interaction from Ab Initio Dynamical Quadrupoles: Impact on Charge Transport in Wurtzite GaN

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First-principles calculations of e-ph interactions are becoming a pillar of electronic structure theory. However, the current approach is incomplete. The piezoelectric (PE) e-ph interaction, a long-range scattering mechanism due to acoustic phonons in non-centrosymmetric polar materials, is not accurately described at present. Current calculations include short-range e-ph interactions (obtained by interpolation) and the dipole-like Fröhlich long-range coupling in polar materials, but lack important quadrupole effects for acoustic modes and PE materials. Here we derive and compute the long-range e-ph interaction due to dynamical quadrupoles, and apply this framework to investigate e-ph interactions and the carrier mobility in the PE material wurtzite GaN. We show that the quadrupole contribution is essential to obtain accurate e-ph matrix elements for acoustic modes and to compute PE scattering. Our work resolves the outstanding problem of correctly computing e-ph interactions for acoustic modes from first principles, and enables studies of e-ph coupling and charge transport in PE materials.

When atoms move due to lattice vibrations, the potential seen by an electron quasiparticle changes due to both short-range and long-range forces. Early theories of such electron-phonon (e-ph) interactions focused on simplified models tailored to specific materials [1]. In metals, the so-called “deformation potential” quantifies the short-range e-ph interactions with acoustic phonons [2]. In ionic and polar covalent materials, in which atoms can be thought of as carrying a net charge, Fröhlich identified a dipole-like long-range e-ph interaction due to longitudinal optical (LO) phonons [3]. In polar materials lacking inversion symmetry, the piezoelectric (PE) e-ph interaction, due to strain induced by acoustic phonons, is also important [4, 5]. Its conventional formulation by Mahan expresses the e-ph coupling in terms of the macroscopic PE constants of the material [5].

Vogl unified these e-ph interactions [6], showing that the dipole, PE, and deformation-potential contributions originate from a multipole expansion [7] of the e-ph potential, and analyzed its behavior in the long-wavelength limit (phonon wave vector \( q \to 0 \)). The Fröhlich dipole term diverges as \( 1/q \) in this limit, and is proportional to the sum of atomic dipoles: If each atom \( \kappa \) is associated with a Born charge tensor, \( Z_\kappa \), contracting this tensor with the phonon eigenvector \( e_{\nu \mathbf{q}} \) gives the atomic contributions to the dipole field, \( Z_\kappa e_{\nu \mathbf{q}} \). Next in the multipole expansion is the quadrupole field generated by the atomic motions, which approaches a constant value as \( \mathbf{q} \to 0 \). If each atom is associated with a dynamical quadrupole tensor, \( Q_\kappa \), the atomic quadrupole contributions can be written as \( Q_\kappa e_{\nu \mathbf{q}} \). Both the dipole and the quadrupole terms contribute to the PE e-ph interaction [6]. The multipole expansion also gives octupole and higher terms, which vanish at \( \mathbf{q} \to 0 \) and can be grouped together into a short-range deformation potential.

Density functional theory (DFT) [8] and density functional perturbation theory (DFPT) [9] have enabled calculations of e-ph interactions from first principles. In turn, the e-ph interactions can be used in the Boltzmann transport equation (BTE) framework to predict electron scattering processes and charge transport [10–21]. The recent development of the ab initio Fröhlich e-ph interaction [22, 23] has been a first step toward implementing Vogl’s modern e-ph theory in first-principles calculations and correctly capturing the long-range e-ph contributions. However, a key piece is still missing in the ab initio framework: the quadrupole e-ph interaction, which critically corrects the dipole term in polar materials, is sizable in nonpolar materials, and is particularly important in PE materials such as wurtzite crystals or titanates.

In this Letter, we derive the ab initio quadrupole e-ph interaction and compute it for a PE material, wurtzite gallium nitride (GaN), using dynamical quadrupoles computed from first principles. We show that including the quadrupole term provides accurate e-ph matrix elements and is essential to obtaining the correct acoustic phonon contribution to carrier scattering. We compute the electron and hole mobilities in GaN including the quadrupole interaction, obtaining results in agreement with experiment. Our analysis highlights the large errors resulting from including only the Fröhlich term in GaN [24], which greatly overestimates the acoustic mode e-ph interactions [24, 25]. Our companion paper applies this framework to silicon and the PE material PbTiO₃. The e-ph quadrupole contribution is sizable in both cases, and essential to correctly compute the e-ph matrix elements. Taken together, the quadrupole interaction completes the theory and enables accurate ab initio e-ph calculations in all materials and for all phonon modes.
The key ingredient in first-principles e-ph calculations are the e-ph matrix elements, $g_{mn\nu}(k,q)$, which encode the probability for scattering from an initial Bloch state $|m_k\rangle$ with band index $m$ and crystal momentum $k$ into a final state $|n(k+q)\rangle$ by emitting or absorbing a phonon with branch index $\nu$ and wave vector $q$ [17, 26]. Following Vogl [6], we separate the long-range dipole and quadrupole contributions from the short-range part:

$$g_{mn\nu}(k,q) = g^{\text{dipole}}_{mn\nu}(k,q) + g^{\text{quad}}_{mn\nu}(k,q) + g^{S}_{mn\nu}(k,q) \quad (1)$$

where $e$ is the electron charge, $\Omega$ is the unit cell volume, $M_k$ and $\tau_{\alpha}$ are the mass and position of the atom with index $\kappa$, $G$ are reciprocal lattice vectors, $e_{\nu}(\kappa)$ is the phonon eigenvector projected on atom $\kappa$, and $\epsilon$ is the dielectric tensor of the material. Summation over the Cartesian indices $\alpha, \beta, \gamma$ is implied.

The dynamical quadrupoles $Q_{\kappa}$ entering Eq. (2) are third-rank tensors defined as the second order term in the long-wavelength expansion of the cell-integrated charge-density response to a monochromatic displacement $\text{e}^{i(q+G_k)\cdot(r-R_k)}|n_k\rangle$.

$$Q^{\alpha\gamma}_{\kappa,\beta} = i\Omega \left( \frac{\partial F^{\alpha\gamma}_{\kappa,\beta}}{\partial q_\gamma} \bigg|_{q=0} + \frac{\partial F^{\kappa\kappa}_{\gamma,\beta}}{\partial q_\alpha} \bigg|_{q=0} \right) \quad (3)$$

With the full first-principles e-ph matrix elements in Eq. (1) in hand, we compute the phonon-limited mobility at temperature $T$ using the BTE in both the relaxation time approximation (RTA) and with a full iterative solution [17]. Briefly, we compute the e-ph scattering rates (and their inverse, the relaxation times $\tau_{mn\nu}$), from the lowest order e-ph self-energy [26]. The mobility is then obtained as the energy integral [17]:

$$\mu_{\alpha\beta}(T) = \frac{e}{n_e \Omega N_k} \int dE \left( \frac{\partial}{\partial E} \right) \sum_{nk} F^{\alpha}_{nk}(T) v^{\beta}_{nk} \delta(E-\varepsilon_{nk}) \quad (4)$$

where $n_e$ is the carrier concentration, $f$ is the Fermi-Dirac distribution, $N_k$ is the number of k-points, and $v_{nk}$ and $\varepsilon_{nk}$ are electron band velocities and eigenvalues, respectively. The $F^{\alpha}_{nk}$ term is obtained as $\tau_{nk}(T) v^{\alpha}_{nk}$ in the RTA or by solving the BTE iteratively.

We carry out \textit{ab initio} calculations on wurtzite GaN with relaxed lattice parameters, using the same settings as in our recent work [31]. The ground state properties and electronic wave functions are computed using DFT in the generalized gradient approximation [32] with the QUANTUM ESPRESSO code [33, 34]. We include spin-orbit coupling by employing fully relativistic norm-conserving pseudopotentials [35, 36] (generated with Pseudo Dojo [37]) and correct the DFT band structure using GW results. We use DFPT [9] to compute phonon frequencies and eigenvectors, and obtain the e-ph matrix elements $g_{mn\nu}(k,q)$ on coarse $8 \times 8 \times 8$ k-point and q-point grids [38] using our PERTURBO code [17]. We obtain the dynamical quadrupoles $Q_{\kappa}$ by computing $\bar{T}^{\alpha\beta}_{\alpha\kappa\beta}$ in Eq. (3) with the methodology in Ref. [30] as implemented in the ABINIT code [39], and validate the results against DFPT clamped-ion PE constants (See Supplemental Material [40]). We subtract the long-range terms $g^{\text{dipole}} + g^{\text{quad}}$ and add back the long-range terms. The resulting matrix elements are employed to compute relaxation rates and mobilities [40] with the PERTURBO code [17].

Since DFPT can accurately capture the long-range e-ph interactions, it can be used as a benchmark for our approach of including the dipole plus quadrupole terms after interpolation. Note that due to computational cost, DFPT calculations cannot be carried out on the fine grids needed to compute electrical transport, so the long-range terms need to be added after interpolation. Following Ref. [22], we define a gauge-invariant e-ph coupling strength, $D^\kappa_{tot}$, proportional to the absolute value of the e-ph matrix elements:

$$D^\kappa_{tot}(q) = \hbar^{-1} \sqrt{2\epsilon_{ij} M_{uc} \sum_{\nu m_n} |g_{mn\nu}(k=\Gamma,q)|^2 / N_b} \quad (5)$$

with $M_{uc}$ the unit cell mass and $N_b$ the number of bands. We compute $D^\kappa_{tot}(q)$ with various approximations to analyze the role of the quadrupole e-ph interactions.
In Fig. 1, we use $D_{\nu}^{\text{tot}}(q)$ obtained from direct DFPT calculations of the matrix elements as a benchmark, and compare calculations that include only the long-range Fröhlich dipole interaction and both the dipole and quadrupole interactions. The short-range $e$-ph interactions are included in both cases as a result of the Wannier interpolation. Including the quadrupole term dramatically improves the accuracy of the $e$-ph matrix elements for the longitudinal acoustic (LA) and transverse acoustic (TA) modes at small $q$ (near $\Gamma$ in Fig. 1). The discrepancy between the dipole-only calculation and DFPT is completely corrected when including the quadrupole term, which reproduces the DFPT benchmark exactly. While the dipole-only scheme leads to large errors, the dipole and quadrupole contributions, as can be seen, cancel each other out since they are nearly equal and opposite for acoustic modes in GaN.

Both the dipole and quadrupole terms contribute to the PE $e$-ph interaction from the LA and TA acoustic modes [6]. Expanding the phonon eigenvectors at $q \to 0$ as $e_{\alpha q} \approx e_{\alpha q}^{(0)} + i q e_{\alpha q}^{(1)}$ [6], one finds two PE contributions of order $q^0$ for $q \to 0$ [6]. One is from the Born charges, $Z_{\kappa} (i q e_{\kappa q}^{(1)})$, and is a dipole-like interaction generated by atoms with a net charge experiencing different displacements due to an acoustic mode. The other is from the dynamical quadrupoles, $Q_{\kappa} e_{\kappa q}^{(0)}$, and is associated with a clamped-ion electronic polarization [42]. As a result, the ab initio Fröhlich term includes only part of the PE $e$-ph interaction, so the dipole-only scheme fails in GaN because it neglects the all-important quadrupole electronic contribution. We also implemented and tested Mahan’s phenomenological PE coupling [5],

$$g_{\nu e}^{\text{PE}}(q) = 4\pi \frac{e^2}{4\pi \varepsilon_0} \left( \frac{\hbar}{2\omega_{\nu q} M_{\text{mc}}} \right)^{\frac{1}{2}} q_{\alpha} e_{\alpha,\beta\gamma} e_{\nu q\beta\gamma} q_{\gamma}$$  \hspace{1cm} (6)

where $e_{\alpha,\beta\gamma}$ are the PE constants of GaN [40]. In the $q \to 0$ limit, this model includes both ionic-motion and electronic effects [6], and is a less computationally demanding alternative that does not require computing the dynamical quadrupoles. We find that the $e$-ph coupling $D_{\nu}^{\text{tot}}(q)$ obtained from the Mahan model improves over the dipole-only scheme [40], but exhibits discrepancies with direct DFPT calculations at finite $q$, and overall is inadequate for quantitative calculations.

Figure 2 shows the effect of using the more accurate quadrupole scheme on the $e$-ph scattering rates (defined as the inverse of the $e$-ph relaxation times, $\Gamma_{nk} = 1/\tau_{nk}$) computed at 300 K. We focus on the energy range of interest for charge transport near 300 K, namely an energy window within $\sim 100$ meV of the band edges. Since these energies are below the LO phonon emission threshold (90 meV in GaN), LO scattering is suppressed and
dominated by thermally activated LO phonon absorption processes. On the other hand, there is a large phase space for acoustic phonon scattering, especially for interband processes in the valence band. For electrons, including the quadrupole term greatly suppresses the acoustic mode contribution to the scattering rates, reducing it from nearly half of the total scattering rate to a negligible contribution (Fig. 2). This result is due to the cancellation of the dipole and quadrupole terms for acoustic phonons discussed above. As expected, the LO contribution becomes dominant for electrons in the conduction band, where small-$q$ intravalley scattering is controlled by the Fröhlich interaction with $1/q$ behavior rather than by the PE dipole and quadrupole terms, both of which have a $q^0$ trend at small $q$. Using correct $\epsilon$-$\phi$ matrix elements that include the quadrupole term, our calculation restores this physical intuition.

A similar but less pronounced trend is found for holes in Fig. 2, where at the peak of the mobility integrand (~50 meV below the valence band edge) acoustic phonon scattering is suppressed from 75% of the total scattering rate when only the Fröhlich interaction is included to less than 50% when including the quadrupole term. The large acoustic phonon scattering for holes found in recent calculations [24] is partially due to the fact that the quadrupole term was missing. The possibility of increasing hole mobility by engineering lower acoustic scattering based on these investigations [25] clearly does not properly take into account PE scattering (and PE charges induced by straining GaN), and should be revisited.

Analysis of the temperature dependent mobility in GaN, computed using Eq. (4) with both the RTA and iterative BTE approaches, highlights the key role of the quadrupole $\epsilon$-$\phi$ interaction. Figure 3 shows the electron and hole mobilities in the basal [1000] plane of GaN, computed both using Wannier interpolation plus the Fröhlich interaction and with our improved scheme including the quadrupole term. Experimental mobility measurements [43–49] are also given for comparison. Compared to calculations that include only the dipole Fröhlich interaction, including the quadrupole term removes the artificial overestimation of acoustic phonon scattering, thus increasing the computed mobility and correcting its temperature dependence, especially at lower temperatures, where acoustic scattering is dominant.

We find good agreement between our computed electron and hole mobilities and experimental results, especially when comparing with the highest mobilities measured in samples with low doping concentrations ($\sim 10^{15}$ cm$^{-3}$ for n-type [43] and $\sim 10^{16}$–$10^{17}$ cm$^{-3}$ for p-type [47, 50] GaN). In these high purity samples, charge transport is governed by phonon scattering in the temperature range we investigate, so these measurements are ideal for comparing with our phonon-limited mobilities.

We focus on the iterative BTE results, whose accuracy is superior to the RTA, also given for completeness [51]. For holes, in which acoustic scattering is significant, the temperature dependence of the mobility is improved after including the quadrupole interaction, as shown in the upper panel of Fig. 3. The exponent $n$ in the temperature dependence of the mobility, $\mu \sim T^{-n}$, is $n = 2.3$ after including the quadrupole term, the same value as the exponent obtained by fitting the experimental data (for comparison, $n = 2.0$ in the dipole-only calculation). The temperature dependence of the electron mobility is only in reasonable agreement with experiment. Including two-phonon processes may be needed to correctly predict the temperature trend, as we found recently for electrons in GaAs [52], which similar to GaN has dominant LO phonon scattering in the conduction band.

Our improved scheme increases the computed mobilities, placing them slightly above the experimental values. This trend is physically correct – our computed mobilities are an upper bound as experimental samples may exhibit additional scattering from defects and interfaces. In addition, a slight mobility overestimation is expected in polar materials because including two-phonon scattering processes would lower the mobility and bring it closer to experiment, as we have recently shown in GaAs [52].
In summary, we have presented a framework for computing the quadrupole e-ph interaction and including it in ab initio e-ph calculations. Our results show its crucial contribution to acoustic phonon and PE scattering. Our work enables accurate calculations of long-range acoustic phonon interactions and paves the way to studies of charge transport in PE materials.

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Note added.— While writing this manuscript, we became aware of a related preprint by another group [53]. Their article analyzes how the quadrupole term improves e-ph matrix element interpolation, while ours focuses more broadly on the physics of e-ph interactions, acoustic phonons, and piezoelectric materials.

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