Nonadiabatic Born Effective Charges in Metals and the Drude Weight

Cyrus E. Dreyer, Sinisa Coh, and Massimiliano Stengel

Department of Physics and Astronomy, Stony Brook University, Stony Brook, New York, 11794-3800, USA
Center for Computational Quantum Physics, Flatiron Institute, 162 5th Avenue, New York, New York 10010, USA
Institut de Ciencia de Materials de Barcelona (ICMAB-CSIC), Campus UAB, 08193 Bellaterra, Spain
ICREA-Instituciò Catalana de Recerca i Estudis Avançats, 08010 Barcelona, Spain

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In insulators, Born effective charges describe the electrical polarization induced by the displacement of individual atomic sublattices. Such a physical property is at first sight irrelevant for metals and doped semiconductors, where the macroscopic polarization is ill defined. Here we show that, in clean conductors, going beyond the adiabatic approximation results in nonadiabatic Born effective charges that are well defined in the low-frequency limit. In addition, we find that the sublattice sum of the nonadiabatic Born effective charges does not vanish as it does in the insulating case, but instead is proportional to the Drude weight. We demonstrate these formal results with density functional perturbation theory calculations of Al and electron-doped SnS₂ and SrTiO₃.

As we shall see shortly, one can circumvent the issues with defining and calculating BECs in metallic systems by relaxing the adiabatic (or Born-Oppenheimer) approximation, a key assumption in the PCM derivations. The adiabatic approximation is usually justified in insulators, where the electronic gap is much larger than typical optical phonon frequencies (< 200 meV for inorganic materials [18]), but generally breaks down in a metallic or doped semiconducting system. Manifestation of nonadiabaticity in lattice dynamics is well known, e.g., plasmon-phonon coupling [19,20] and renormalization of the phonon frequencies [21]. In particular, there exists a regime near the zone center where the phase velocity of an optical phonon is large compared to the Fermi velocity [22–24]. In such a regime, the nonadiabatic dynamical matrix may significantly differ from its adiabatic counterpart. Also, free carriers are unable to screen the long-range electric fields [25], and signatures of the BECs can be detected experimentally as resonances in the reflectivity spectra [26,27] or shifts in the plasmon frequency [19].

In this work, we show that nonadiabatic BECs (NABECs) are well-defined real quantities [as long as time-reversal (TR) symmetry is preserved] in the low-frequency regime relevant for lattice dynamics. Remarkably, we find that their sublattice sum in general does not vanish but tends to the Drude weight (DW), which gives the density of free electrons available for conduction. This generalized sum rule implies that the NABECs never vanish in metals or doped semiconductors (not even in elemental quasi-free-electron crystals). This result, in addition to clarifying the origin of the ASR breakdown that was pointed out in Ref. [27], provides a novel physical interpretation of the

Born effective charges (BECs) are defined as the polarization induced by the displacement of an atomic sublattice [1–3]. They are key quantities for the microscopic understanding of a wide variety of phenomena in insulators, including the interplay between electric fields and long-wavelength optical phonons [1,4–7], lattice contribution to dielectric screening [1,4] and electromechanical coupling [8–10], and polar properties of ferroelectrics [11–13]. BECs (also known as dynamical charges) were placed on a firm conceptual footing in the seminal work of Pick, Cohen, and Martin (PCM) [3], which formally established the acoustic sum rule (ASR), that a rigid translation of the crystal lattice does not produce either forces on the individual atoms or a net electrical polarization. This is a consequence of translational invariance and charge neutrality, and requires that the sublattice sum of the BECs vanish.

The macroscopic polarization and its parametric derivatives are only well defined in a gapped system [14], which at first sight rules out the calculation of BECs in metals. In addition, a hypothetical definition of the BECs as the dipolar moment of the first-order charge induced by an atomic displacement would also be problematic, since the free carriers screen any long-range electrostatic perturbation. Yet, the concept of BEC is routinely used to study free carriers screen any long-range electrostatic perturbation. The adiabatic approximation is usually justified in insulators, where the electronic gap is much larger than typical optical phonon frequencies (< 200 meV for inorganic materials [18]), but generally breaks down in a metallic or doped semiconducting system. Manifestation of nonadiabaticity in lattice dynamics is well known, e.g., plasmon-phonon coupling [19,20] and renormalization of the phonon frequencies [21]. In particular, there exists a regime near the zone center where the phase velocity of an optical phonon is large compared to the Fermi velocity [22–24]. In such a regime, the nonadiabatic dynamical matrix may significantly differ from its adiabatic counterpart. Also, free carriers are unable to screen the long-range electric fields [25], and signatures of the BECs can be detected experimentally as resonances in the reflectivity spectra [26,27] or shifts in the plasmon frequency [19].

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DW in terms of lattice dynamics (as opposed to electrical transport). Indeed, the DW emerges as the mechanical inertia of the conduction electrons to an acceleration of the underlying lattice potential, which cannot drag the carriers along if subjected to a sufficiently rapid oscillation. We demonstrate our formal results with a density-functional perturbation theory (DFPT) [28,29] based methodology for calculating the NABECs and DW for a prototypical metal (Al) and two doped semiconductors (SnS₂ and SrTiO₃). In the latter, we find that NABECs can deviate significantly from their values in the undoped material.

We shall frame our discussion in terms of generalized susceptibility functions, $\chi_{\lambda_1\lambda_2}(\omega, \mathbf{q})$, describing the $\lambda_1$ response to the perturbation $\lambda_2$, modulated by a wave vector $\mathbf{q}$ and at a frequency $\omega$. We focus henceforth on the cases where $\lambda_{1,2}$ are either an atomic displacement of sublattice $\kappa$ in direction $\alpha$ ($\tau_{\kappa\alpha}$) or a component of the vector potential ($A_\alpha$). The purely electromagnetic case is well known: $\chi_{A_\alpha A_\beta}(\omega, \mathbf{q})$ is the current-current response [30], whose long-wavelength limit relates to the macroscopic optical conductivity via [31]

$$\sigma_{\alpha\beta}(\omega) = \frac{i}{\omega} \lim_{\mathbf{q} \to 0} \chi_{A_\alpha A_\beta}(\omega, \mathbf{q}).$$

(As customary, we shall assume that the frequency has a small imaginary part $i\eta$ to ensure causality [32].) In close analogy with Eq. (1), we shall define NABECs as [26,27]

$$Z_{\kappa\alpha}(\omega) = \frac{i}{\omega} \lim_{\mathbf{q} \to 0} \chi_{A_\alpha \tau_{\kappa\alpha}}(\omega, \mathbf{q}),$$

(2)

which has the physical meaning of a current response to the atomic velocity [33].

For most materials, the interesting physics is contained in the small-$\omega$ behavior of $Z_{\kappa\alpha}(\omega)$, i.e., at the frequencies that are relevant for lattice dynamics. Within such a regime, the conductivity [Eq. (1)] vanishes in topologically trivial insulators [32,34] and Eq. (2) reduces to the standard linear-response formula [4,28,35] for the adiabatic BEC. Note that, in the insulating case, the result is also independent of the order of the long-wavelength and small-frequency limits.

In metals, Eq. (1) diverges as $\omega \to 0$, resulting in the Drude peak in the longitudinal conductivity [31]. Since Eq. (2) is formally similar to Eq. (1), it is reasonable to wonder whether $Z_{\kappa\alpha}(\omega) \to 0$ diverges as well, or tends to a finite constant. The divergence in $\sigma_{\alpha\beta}(\omega \to 0)$ is rooted in the fact that $\chi_{A_\alpha A_\beta}$ tends to a finite nonzero limit in all metals,

$$\lim_{\omega \to 0} \chi_{A_\alpha A_\beta}(\omega) = \frac{D_{\alpha\beta}}{\pi},$$

(3)

where $D_{\alpha\beta}$ is the DW tensor [32]. The situation regarding the mixed response to a vector potential and an atomic displacement clearly differs, as it requires time-reversal symmetry to be broken in order to be nonzero. (A is odd with respect to both TR and space inversion, while $\tau_{\kappa\alpha}$ is TR even.) This simple argument guarantees that in TR-symmetric metals, where a steady current in response to a static atomic displacement is forbidden, $\chi_{A_\alpha \tau_{\kappa\alpha}}(\omega \to 0)$ vanishes and therefore $Z_{\kappa\alpha}(\omega \to 0)$ is well defined. Note that, unlike the insulating case, here the order of the $\mathbf{q} \to 0$ and $\omega \to 0$ limits does not commute; our prescription of taking the $\mathbf{q} \to 0$ first is relevant for optical phonons, which retain a finite frequency in the long-wavelength limit.

Our arguments, in principle, do not concern acoustic phonons, since their frequency vanishes linearly with momentum in a neighborhood of the zone center. Yet, in the following we shall consider hypothetical long-wavelength acoustic phonons whose limiting $\mathbf{q} \to 0$ frequency is artificially assumed to remain finite. This assumption, while physically unrealistic, will serve as an intermediate step toward the derivation of a sum rule for the nonadiabatic BECs, which shall be our next goal. The rationale behind this procedure is that an acoustic phonon at $\mathbf{q} = 0$ reduces to a rigid translation of the whole crystal lattice. In the linear regime, a translation can be regarded as the sum of the displacements of all individual atomic sublattices; thus, by looking at the current density induced by a “zone-center acoustic phonon” that is externally modulated in time at a frequency $\omega$, one can directly infer the sublattice sum of the NABECs [Eq. (2)].

The advantage of such an approach is that the acoustic phonon perturbation enjoys a remarkably compact expression when rewritten, via a coordinate transformation, in the curvilinear frame that is comoving with the atoms [36–38]. Thereby, an acoustic phonon can be recast as a “static” strain contribution that only depends on the metric of the deformation, plus a “dynamical” effective vector potential that results from the inertia of the electrons upon local displacements of the coordinate frame [37]. In a Kohn-Sham [39] density-functional context [see Supplemental Material (SM), Sec. S1 [40]], we can then write the sublattice sum of the first-order phonon ($\tau_{\kappa\alpha}$) Hamiltonians as [37]

$$\sum_{\kappa} \hat{H}_{\kappa}(\omega, \mathbf{q}) = \hat{H}^{(\beta)}(\omega, \mathbf{q}) + i\omega \hat{H}^{(A)}(\omega, \mathbf{q}),$$

(4)

where $\hat{H}^{(\beta)}(\omega, \mathbf{q})$ is the “metric wave” Hamiltonian of Refs. [37,38], and $\hat{H}^{(A)}(\omega, \mathbf{q})$ is the additional effective vector-potential perturbation.

As a result of translational invariance, $\hat{H}^{(\beta)}(\omega, \mathbf{q})$ identically vanishes at the zone center [38], and thus the rigid displacement of the lattice reduces to the vector-potential perturbation term at $\mathbf{q} = 0$. By combining this observation with Eqs. (1) and (2), we readily arrive at
relating the sum of the NABECs to the optical conductivity ($\Omega$ is the cell volume). From Eq. (3) we find that the small-frequency limit of Eq. (5) becomes
\begin{equation}
\frac{1}{\Omega} \sum_k Z_{k,p}^{(a)}(\omega) = i\omega\sigma_{0j}(\omega), \tag{5}
\end{equation}
where $Z_{k,p}^{(a)} = Z_{k,p}^{(a)}(\omega \to 0)$ is the zero-frequency limit of the NABEC. We call Eq. (6) the NABEC sum rule for the NABEC. We call Eq. (6) the NABEC sum rule for
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\end{equation}

We now demonstrate the sum rule in Eq. (6) with DFT calculations performed on three systems: the face-centered cubic (fcc) phase of Al, electron-doped bulk ($P\bar{3}m1$) SnS$_2$, and electron-doped cubic ($Pm\bar{3}m$) SrTiO$_3$ (STO). The details of the implementation, computational parameters, and numerical convergence for these calculations are given in the SM, Secs. S1 D and S2 [40]. The doping in SnS$_2$ and STO is applied under the “rigid band” approximation, i.e., all matrix elements and structural parameters are taken from an undoped calculation, and just the occupation factors are changed to reflect the added electrons; the excess positive charge to compensate the electrons is added to the ionic BEC of a given sublattice, corresponding to chemical doping on that sublattice. In SM, Sec. S2 C [40], we compare these results to explicit doping via the virtual crystal approximation, demonstrating that the NABEC sum rule is satisfied regardless of the approximation.

We calculate the NABECs in the $\omega \to 0$ limit as (see SM, Sec. S1 [40], and Refs. [26,27])
\begin{equation}
Z_{a,k,p} = Z_{k}^{\text{ion}}\delta_{a,p} - \text{Im} \lim_{\eta \to 0} \int [d^3k] \sum_{n \neq m} \frac{f_{nk} - f_{nm}}{(e_{nk} - e_{nm})(e_{nk} - e_{nm} + i\eta)} \langle u_{nk} | \hat{H}_k^{A_n} | u_{mk} \rangle \langle u_{mk} | \hat{F}_k^{\alpha} | u_{nk} \rangle, \tag{7}
\end{equation}

where $Z_k^{\text{ion}}$ is the ionic (i.e., pseudopotential) charge, $e_{nk}$ and $f_{nk} = f(e_{nk})$ are the energy and Fermi occupation factor of band $n$ and $k$ point $k$, $u_{nk}$ is the cell-periodic part of the corresponding Bloch function, $\hat{H}_k^{A_n}$ is the $k$ derivative of the Hamiltonian, and $\hat{F}_k^{\alpha}$ is the screened (i.e., including self-consistent fields) first-order phonon Hamiltonian. The DW, in turn, is calculated via [32]
\begin{equation}
\Omega D_{0j} = \sum_k \frac{\partial f(e_{nk})}{\partial e} v_{nk}^{(a)} v_{nk}^{(j)}, \tag{8}
\end{equation}
where $v_{nk}^{(a)} = \langle u_{nk} | \hat{H}_k^{A_n} | u_{nk} \rangle$ is the $a$ component of the band velocity. Note that, because of a technical subtlety related to the use of nonlocal pseudopotentials (NLPSPs) in the calculations, the sum rule in Eq. (6) is exactly satisfied only if the second band velocity in Eq. (8) is replaced with the canonical momentum, $v_{nk}^{(j)} = \langle u_{nk} | \hat{P}_{jk} | u_{nk} \rangle$. We shall refer to this revised version of Eq. (8) as “modified” DW, $D_{0j}$ in the following.) We ascribe this outcome to the well-known ambiguities that arise when combining NLPSPs with electromagnetic fields [50–52]; in practice, we find that its quantitative impact is small.

In Fig. 1 we plot the NABEC $Z^*$, standard DW $D$, and modified DW $\hat{D}$ for fcc Al versus the number of $k$ points used to sample the Brillouin zone of the conventional cubic cell [53]. Since fcc Al has no optical modes, the non-adiabatic regime is hardly relevant here from a physical point of view; yet, its computational simplicity allows for a numerical test of our arguments. Remarkably, the NABEC of Al does not vanish in spite of it being an elemental metal, and converges to a value of around 2e. We see from the red dashed curve that the sum rule in Eq. (6) is accurately
satisfied for a large enough \( k \) mesh. We also see that \( \tilde{D} \) differs slightly from \( D \), and this results in a violation of the NABEC sum rule that is modest (\( \sim 0.1 e \)) but clearly discerned for the numerical accuracy of our calculations. We find a similar qualitative behavior for all materials in this study (see, e.g., SM, Figs. S9–S11 [40]); thus, we shall exclusively focus on \( \tilde{D}_{\alpha\beta} \) henceforth.

We now consider bulk SnS\(_2\), a layered insulator that has shown considerable promise for use in a wide range of device applications [54–58]. We dope SnS\(_2\) with electrons into the isolated, lowest-energy Sn(5s)-S(3p) conduction band [see Fig. S2(b) in SM [40]]. In order to accurately describe the Fermi surface, we perform the Brillouin-zone integrals of Eqs. (7) and (8) via Wannier interpolation to a dense \( k \) mesh (see SM, Sec. S2 B [40]).

In Fig. 2(a), we show the NABECs versus doping for the Sn and S sublattices, in the in-plane \( x \) and out-of-plane \( z \) directions. The dotted curves in Fig. 2(a) correspond to the excess ionic charge placed on the S sublattice, and the solid curves to the charge placed on the Sn sublattice. In Fig. 2(b) we see that the violation of the (sublattice-summed) BEC charge neutrality condition increases as we dope to roughly half of the conduction-band width, and then decreases back to zero when the conduction band is filled. The DW exhibits the same behavior, and we see from the red curves around zero that the NABEC sum rule [Eq. (6)] is accurately satisfied.

Finally, we calculate the NABECs and DW for the electron-doped cubic phase of STO, which has attracted significant attention due to its superconducting properties [59–61]. To illustrate the behavior of the NABECs, we present doping across the entire Ti \( t_{2g} \) conduction band manifold [see Fig. S2(a) [40]], even though it is not expected to be experimentally achievable. (Also, we should note that significant changes to the electronic and atomic structure are expected if such calculations at high doping are performed with atomic relaxations [62].)

In Figs. 3(a)–3(c) we plot the NABECs for the sublattices in STO with doping. The oxygens are labeled as “equatorial” (on faces parallel to \( \hat{t} \) for BEC element \( Z_{ij}^{\alpha} \)) and “apical” (on faces perpendicular to \( \hat{t} \)). Different trends in \( Z^\alpha \) result from different choices of sublattice for chemical doping, suggesting significant qualitative changes in frequencies of polar-phonon modes depending on the

![FIG. 1. Demonstration of nonadiabatic Born effective charge (NABEC) sum rule for Al versus \( k \) points in the Brillouin zone of the conventional 4-atom cell. Blue dot-dashed curve is the NABEC, the black solid (pink dotted) curve is minus the cell volume multiplied by the modified (normal) Drude weight, and red dashed (green dotted) curve is the sum of the two. Inset is a magnification around 0 \( e \) for the largest \( k \) mesh.](image1)

![FIG. 2. (a) Nonadiabatic Born effective charges versus doping above the conduction-band minimum (CBM) in the in-plane \( x \) direction and out-of-plane \( z \) direction. Solid (dashed) lines are for the excess ionic charge associated with the Sn (S) sublattice. (b) Demonstration of the NABEC sum rule, where the sublattice sum of the BECs equals the Drude weight. Blue shaded region is the DOS of the conduction band calculated from the Wannier interpolation.](image2)

![FIG. 3. Nonadiabatic Born effective charges of the sublattices in SrTiO\(_3\) versus doping above the conduction-band minimum on the (a) Sr, (b) Ti, or (c) O sublattice. Panel (d) demonstrates the NABEC sum rule, where the sublattice sum of the BECs cancels with the Drude weight. Blue shaded region is the density of states of the Wannierized Ti \( t_{2g} \) manifold.](image3)
sublattice that is doped versus the sublattice(s) involved in the phonon displacements. Figure 3(a) corresponds to doping on the Sr site, e.g., by La substitution; in this case, the Sr Z’ increases significantly to compensate the decrease of the Ti Z*, resulting from the fact that the electrons are doped into the Ti 3d orbitals; this effect is significantly reduced if we consider doping on the Ti site in Fig. 3(b) (e.g., by Nb substitution). For doping on the O site in Fig. 3(c) (e.g., by the formation of O vacancies), the decrease in the Ti Z* is now partially compensated by the increase in Z* for the O sublattices.

In Fig. 3(d), we see very similar behavior to Fig. 2(b); i.e., the sublattice sum of the BECs increasingly deviates from zero with doping toward the middle of the conduction-band manifold, and then decreases. As with SnS, the NABEC sum rule is accurately satisfied.

In conclusion, we have demonstrated that nonadiabatic Born effective charges are well defined in metals with time-reversal symmetry, and generalized the acoustic sum rule to the full nonadiabatic regime, where the sublattice sum of the nonadiabatic Born effective charges equals the Drude weight. The rigorous understanding and first-principles description of nonadiabatic Born effective charges provided by this work opens up several future directions for study. For example, it will allow quantitative predictions of plasmon-phonon coupling, and the implications for transport in doped semiconductor devices such as transparent conductors. Direct comparison with experimental probes such as infrared or Raman scattering will serve to validate the theory and aid in materials characterization. It may also shed light on the physics of ferroelectric metals, possibly providing additional tools to quantify the amplitude of the polar lattice distortion.

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