From the dipole of a crystallite to the polarization of a crystal

Raffaele Resta\textsuperscript{1,2, a)\textsuperscript{1}}

\textsuperscript{1}Istituto Officina dei Materiali IOM-CNR, Strada Costiera 11, 34151 Trieste, Italy\textsuperscript{b)\textsuperscript{1}}

\textsuperscript{2}Donostia International Physics Center, 20018 San Sebastián, Spain

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The quantum-mechanical expression for the polarization of a crystalline solid does not bear any resemblance to the (trivial) expression for the dipole of a bounded crystallite; and in fact it has been proved via a conceptually different path. Here I show how to alternatively define the dipole of a bounded sample in a somewhat unconventional way; from such formula, the crystalline polarization formula follows almost seamlessly. Additionally, I discuss the issue of depolarization fields inside a spontaneously polarized crystallite (either pyroelectric or ferroelectric), and their relevance in order to ideally retrieve crystalline polarization from the large-crystallite limit.

I. INTRODUCTION

The dipole of a bounded and charge-neutral sample is a very trivial quantity; the macroscopic polarization of a crystalline solid, instead, has been a challenging problem for many years. In quantum mechanics the dipole of a bounded sample is the expectation value of the position operator \( \mathbf{r} \). The drawback is that solid state physics requires Born-von-Kármán periodic boundary conditions (PBCs),\textsuperscript{1} which define the Hilbert space where Schrödinger equation is solved. Unfortunately the multiplicative operator \( \mathbf{r} \) is not a legitimate operator in the PBC Hilbert space: it maps a state vector within the space into an entity which does not belong to the same space.

The ultimate solution of this long-standing problem was arrived at along the 1990s;\textsuperscript{2–6} by now, polarization theory is a mature topic.\textsuperscript{7,8} The historical development of the theory passed through abandoning the concept of polarization “itself”, addressing instead a polarization difference, which could be expressed as a time-integrated adiabatic current.\textsuperscript{2,3} Only afterwards it was realized\textsuperscript{4} that even polarization itself can be defined, although by means of a change of paradigm: bulk polarization is not a vector (as theretofore assumed), is a lattice. A counterintuitive corollary is that the polarization \( \mathbf{P} \) of an inversion-symmetric crystal is not necessarily zero. Since inversion symmetry requires \( \mathbf{P} = -\mathbf{P} \), the lattice must be symmetric: this may happen even if \( \mathbf{P} = 0 \) does not belong to the lattice. For a macroscopic bounded crystallite, the lattice ambiguity is fixed only after the sample termination is chosen.\textsuperscript{8,9}

What is disturbing is that the two definitions of essentially the same observable—dipole of a crystallite vs. polarization of a crystal—do not bear any formal resemblance. A basic tenet of statistical mechanics and condensed matter physics requires instead that crystalline polarization can also be expressed as the large-sample limit of the dipole of a bounded crystallite over its volume. In this work I am going to bridge this conceptual gap; it will be shown that when the dipole of a bounded sample is alternatively expressed in an unconventional way, with no reference to the \( \mathbf{r} \) operator, the crystalline expression follows somewhat naturally.

II. BOUNDED CRYSTALLITE

We assume that \( N \) electrons are confined in a macroscopic sample of volume \( V \), together with a neutralizing background of point–like classical nuclei. Let \( |\Psi_0\rangle \) be the singlet insulating ground eigenstate; the many-body wavefunction is square-integrable over \( \mathbb{R}^{3N} \) and vanishes far away from the sample. If the system is macroscopically homogeneous, the electronic term in polarization has the pretty trivial expression

\[
\mathbf{P}^{(el)} = -\frac{e}{\mathcal{V}} \langle \Psi_0 | \mathbf{r} | \Psi_0 \rangle, \quad \mathbf{r} = \sum_{i=1}^{N} \mathbf{r}_i;
\]

the nuclear classical contribution has to be added in order to obtain a meaningful observable.

It is expedient to address the family of many-body Hamil-
tonians parametrized by the parameter $\kappa$:

$$\hat{H}_{\kappa} = \frac{1}{2m}\sum_{i=1}^{N}(p_i + \hbar\kappa)^2 + \hat{V},$$  \quad (2)$$

where $\hat{V}$ includes one-body and two-body potentials, and whose ground eigenstate is $|\Psi_{0\kappa}\rangle$. In order to simplify notations we will set $\hat{H}_0 \equiv \hat{H}$ and $|\Psi_{n\kappa}\rangle = |\Psi_n\rangle$. The vector $\kappa$, having the dimensions of an inverse length, generalizes the Hamiltonian by including a constant vector potential: it is therefore a pure gauge. The gauge-transformed eigenstates are

$$|\tilde{\Psi}_{n\kappa}\rangle = e^{-i\kappa \hat{r}}|\tilde{\Psi}_n\rangle.$$  \quad (3)$$

We pause at this point to stress an important semantical issue. The choice of the (arbitrary) $\kappa$ value in Eq. (2) fixed the gauge in the Hamiltonian. Once this fixed, there is an additional freedom in choosing the arbitrary phase factor in front of each eigenstate: even this second choice goes under the rug" an issue of overwhelming importance. W e start from Eq. (1) into

$$P^{(el)} = \frac{i e}{\hbar} \langle \tilde{\Psi}_0 | \partial_\kappa \tilde{\Psi}_0 \rangle,$$  \quad (4)$$

but in this seemingly innocent transformation, I have "swept under the rug" an issue of overwhelming importance. We start noticing that Eq. (4) has not the standard form of an observable: it is in fact the real quantity $\Im \langle \tilde{\Psi}_0 | \partial_\kappa \tilde{\Psi}_0 \rangle$ is, in the language of quantum geometry, a Berry connection evaluated at $\kappa = 0$. Since Eq. (2) at two different $\kappa$’s yields two different Hamiltonians, an equally acceptable gauge-transformed eigenstate would be

$$|\tilde{\Psi}_{n\kappa}\rangle = e^{i\phi(\kappa)} e^{-i\kappa \hat{r}}|\tilde{\Psi}_n\rangle,$$  \quad (5)$$

with an arbitrary $\phi(\kappa)$. The physical observable obtains from Eq. (4) when the gauge of Eq. (3) is enforced; it is not allowed to adopt therein the most general gauge of Eq. (5).

The gauge dependence of Berry connections is a textbook fixture of quantum geometry.\textsuperscript{8} In the present case Eq. (4) acquires its physical meaning only after the above specific gauge fixing; the same gauge fixing—Eq. (8) below—will be essential for defining polarization in the crystalline case.

III. UNBOUNDED CRYSTAL

We adopt the same Hamiltonian as in Eq. (2), but now within the PBC Hilbert space: the many-body wavefunction is periodic in the cubic "supercell" of side $L$ in each electronic variable independently, and normalized to one therein.

Each Cartesian coordinate is then equivalent to the angle $\phi_i = 2\pi x_i/L$, and analogously for $y_i$ and $z_i$. The potential $\hat{V}$ enjoys the same periodicity: this means that the macroscopic field $E$ outside the sample vanishes: the issue is relevant in view of the discussion below, Sec. VI. We will indicate the eigenstates as $|\tilde{\Psi}_{n\kappa}\rangle$ without a tilde, in order to distinguish them from those of the bounded crystallite; as stressed above, the multiplicative $\hat{r}$ operator is "forbidden" in the PBC Hilbert space.\textsuperscript{6}

In order to address polarization, we need to ensure beforehand that the ground state is insulating. The many-body velocity operator is hence by Hellmann-Feynman theorem the macroscopic current density is

$$j_{\kappa} = -\frac{e}{\hbar L^3} \langle \Psi_{0\kappa} | \partial_{\kappa} \hat{H}_{\kappa} | \Psi_{0\kappa} \rangle = -\frac{e}{\hbar L^3} \partial_\kappa E_{0\kappa},$$  \quad (7)$$

where $E_{0\kappa}$ is the ground-state energy. Given that an insulator does not sustain a dc current, the ground-state energy is $\kappa$-independent (the opposite is true in metals).

The Hamiltonian of Eq. (2) was first introduced in 1964 in a milestone paper by W. Kohn, who noticed that PBCs violate gauge-invariance in the conventional sense.\textsuperscript{10} If we try the same gauge transformation as in Eq. (3), i.e.

$$|\Psi_{0\kappa}\rangle = e^{-i\kappa \hat{r}}|\Psi_0\rangle,$$  \quad (8)$$

the r.h.s. is a solution of Schrödinger equation with energy $E_0$, but it does not obey PBCs and therefore does not belong to the Hilbert space. At an arbitrary $\kappa$, the genuine PBC eigenstates $|\tilde{\Psi}_{n\kappa}\rangle$ have a nontrivial $\kappa$-dependence. There is, however, a discrete set of special $\kappa$ vectors for which the r.h.s. of Eq. (8) obeys PBCs and yields therefore the ground eigenstate of $\hat{H}_{\kappa}$: $\kappa_{\text{ran}} = \frac{2\pi}{L}(\ell, m, n)$, with integer $(\ell, m, n)$.

In order to define polarization, we proceed by adopting the analogue of Eq. (4), and in the analogous gauge. We start from the identity

$$\partial_\kappa \ln \langle \Psi_0 | \Psi_{0\kappa} \rangle = \frac{\langle \Psi_0 | \partial_\kappa \Psi_{0\kappa} \rangle}{\langle \Psi_0 | \Psi_{0\kappa} \rangle} = \frac{\langle \Psi_0 | \partial_\kappa \Psi_{0\kappa} \rangle}{\langle \Psi_0 | \Psi_{0\kappa} \rangle};$$  \quad (9)$$

since $\langle \Psi_0 | \partial_\kappa \Psi_0 \rangle$ is purely imaginary, a leading-order expansion in $\kappa$ yields

$$i \langle \Psi_0 | \partial_\kappa \Psi_0 \rangle \cdot \kappa \simeq -\Im \ln \langle \Psi_0 | \Psi_{0\kappa} \rangle.$$  \quad (10)$$

We pick a vector $\kappa_1$ in the special set: $\kappa_1 = \frac{2\pi}{L}(1, 0, 0)$, and we replace the derivative in Eq. (4) with a finite difference, in the large-sample limit:

$$P^{(el)} = \frac{e}{2\pi L^2} \Im \ln \langle \Psi_0 | \Psi_{0\kappa_1} \rangle.$$  \quad (11)$$

As it stands, Eq. (11) is gauge-dependent and cannot express an observable: it is in fact a discretized Berry connection. Eq. (11) only acquires physical meaning when we fix the
by adopting the one of Eqs. (3) and (8) (with no extra phase factor):

\begin{align*}
P_x^{\text{el}} &= \frac{e}{2\pi L^2} \text{Im} \ln \langle \Psi_0 | e^{-iK_1 \cdot \hat{r}} | \Psi_0 \rangle \\
&= \frac{e}{2\pi L^2} \text{Im} \ln \langle \Psi_0 | e^{-i\kappa_0 \cdot \Sigma_k \cdot X_k} | \Psi_0 \rangle. \tag{12}
\end{align*}

We have thus arrived at the main message of the present work: the bounded-crystallite formula, Eq. (4), and the crystalline formulation, Eq. (12), are essentially the same formula, within the same gauge, in two different frameworks.

The replacement of \(|\Psi_0(\kappa_1)\rangle\) in Eq. (11) with \(e^{-iK_1 \cdot \hat{r}} |\Psi_0\rangle\) in Eq. (12) is allowed in insulators only. We remind that \(|\Psi_0\kappa\rangle\) obtained by following the ground state \(|\Psi_0\rangle\) when the \(\kappa\) vector in \(\hat{H}_\kappa\) is adiabatically turned on; in the metallic case—as shown by Kohn\(^{10}\)—the energy \(E_{\kappa_0}\) of such state \(\kappa\) does depend on \(\kappa\), and therefore \(|\Psi_0(\kappa_1)\rangle\) is orthogonal to \(e^{-iK_1 \cdot \hat{r}} |\Psi_0\rangle\). We have shown above that in the insulating case the state \(|\Psi_0(\kappa_1)\rangle\) has instead the same energy as \(e^{-iK_1 \cdot \hat{r}} |\Psi_0\rangle\), and therefore the two states may be identified.

A double check that Eq. (12) is indeed the electronic term in polarization comes from addressing the macroscopic current density \(j_i(t)\) flowing through the insulating sample while the Hamiltonian is adiabatically varied in time: it was proved elsewhere\(^{6,11}\) that \(j_i(t)\) coincides indeed with \(dP_i(t)/dt\).

Finally, the nuclear term in polarization can be added to Eq. (12) in a very compact form. If the nuclei of charge \(Z_e\) sit at sites \(R_s\) in the supercell, the expression is

\[ P_s = \frac{e}{2\pi L^2} \text{Im} \ln \langle \Psi_0 | e^{i\Sigma_k Z_e X_k} | \Psi_0 \rangle, \tag{13} \]

where \(X_k = R_s - R\). Owing to charge neutrality, polarization is invariant by translation of the coordinate origin (as it must be). It is argued that Eq. (13) also holds when the quantum nature of the nuclei is considered.

### IV. MULTIVALUED POLARIZATION IN CRYSTALS

I have anticipated in the Introduction that bulk polarization is a lattice, not a vector, and in fact the main entry of Eqs. (12) and (13) is the multiple-valued function “\(\text{Im} \ln\)”. But it is also clear that for a three-dimensional system these equations cannot be accepted as they stand in the large-sample limit: the prefactor goes in fact to zero. The key point is that we have not exploited crystalline symmetry yet.

By definition, whenever a material is crystalline, a uniquely defined lattice can be associated with the real sample. The lattice is a “mathematical construction”,\(^1\) uniquely defined—by means of an appropriate average—even in cases with correlation, finite temperature, quantum nuclei, chemical disorder (i.e. crystalline alloys, a.k.a. solid solutions), where the actual wavefunction may require a supercell (multiple of the primitive lattice cell).

We consider—without loss of generality—a simple cubic lattice of constant \(a\), where the supercell side \(L\) is an integer multiple of \(a\): \(L = Ma\). Suppose the potential \(V\) in the Hamiltonian is adiabatically varied in time; we define the phase angle

\[ \gamma(t) = \text{Im} \ln \langle \Psi_0(t) | e^{i\Sigma_k Z_i X_i - \Sigma_s Z_s} | \Psi_0(t) \rangle, \tag{14} \]

where \(|\Psi_0(t)\rangle\) is the adiabatic ground eigenstate. The current flowing across a section of area \(L^2\) normal to \(x\) is

\[ I_x(t) = L^2 P_x(t) = \frac{e}{2\pi} \gamma(t). \tag{15} \]

Owing to crystalline periodicity, the current \(I_x(t)\) is the sum of \(M^2\) identical currents, each flowing through a microscopic section of area \(a^2\): one can therefore define a reduced crystalline phase angle \(\gamma_{\text{crystal}}^x\) such that \(\gamma(t) = M^2 \gamma_{\text{crystal}}^x(t)\). The crystalline polarization is thus expressed in terms of \(\gamma_{\text{crystal}}^x\) as

\[ P_x = \frac{e}{2\pi a^2} \gamma_{\text{crystal}}^x, \tag{16} \]

the case of independent electrons is presented in detail in the next Section.

A generic lattice is dealt with by means of a coordinate transformation:\(^{12}\) the bulk value of \(P\) is then ambiguous module \(eR/\gamma_{\text{cell}}\), where \(R\) is a lattice vector and \(\gamma_{\text{cell}}\) is the volume of a primitive cell. The quantity \(eR/\gamma_{\text{cell}}\) goes under the name of polarization “quantum”. By definition a primitive cell is a minimum-volume one: this choice is mandatory in order to make \(P\) an unambiguously defined multivalued observable. Finally we observe that the modulo ambiguity is only removed when the termination of the bounded sample is specified; it is also required that even the surfaces, as well as the bulk, are insulating.\(^5\) Insofar as the crystalline system is unbounded the modulo ambiguity cannot be removed.

### V. SINGLE-DETERMINANT WAVEFUNCTION

Within mean field (either Hartree-Fock or Kohn-Sham) the ground eigenstate \(|\Psi_0\rangle\) in the Schrödinger representation is a Slater determinant of \(N/2\) doubly occupied orbitals; in the crystalline case translational symmetry allows choosing the orbitals in the Bloch form. For the sake of simplicity we get rid of trivial factors of two, by considering a Slater determinant of singly occupied orbitals (so-called “spinless electrons”); furthermore we consider the contribution to \(P_x^{\text{el}}\) of a single occupied band.

In the simple cubic case, as dealt with above, the Bloch vectors are:

\[ k_m = \frac{2\pi}{Ma}(m_1, m_2, m_3), \quad m_s = 0, 1, \ldots, M - 1, \tag{17} \]

where \(m = (m_1, m_2, m_3)\). The Bloch orbitals \(|\psi_k\rangle = e^{ik\cdot x} |\psi_k\rangle\) are normalized over the crystal cell of volume \(a^3\). It is expedient to define the auxiliary Bloch orbitals \(|\phi_k\rangle = e^{ik\cdot x} |\phi_k\rangle\), and \(|\Phi_0\rangle\) as their Slater determinant; we also define \(q = (\frac{2\pi}{Ma}, 0, 0)\). Then

\[ \langle \Psi_0 | e^{i\Sigma_k q \cdot r} | \Psi_0 \rangle = \langle \Psi_0 | \Phi_0 \rangle = \frac{1}{M^3} \text{det} \mathcal{S}, \tag{18} \]
VI. DEPOLARIZATION FIELDS AND SHAPE ISSUES

This is indeed the single-band version of the depolarization; it is tacitly assumed that the macroscopic field $\mathbf{E}$ nucleates of charge $e$ inside field material (either pyroelectric or ferroelectric) the macroscopic sample is made of a spontaneously polarized in the vacuum region far away from the sample; then if the dependent.

$P = e \lim_{L \to \infty} \frac{1}{L} \left( \lim_{L_y \to \infty} \frac{1}{L_y} \langle \Psi_0 | \sum \mathbf{r} \mathbf{r} \mathbf{r} \mathbf{r} \mathbf{r} \mathbf{r} \mathbf{r} \mathbf{r} | \Psi_0 \rangle \right)$,

where $x_i = R_{i,x}$. The two limits in Eq. (24) can be spelled out as follows: the quantity in parenthesis is the dipole (parallel to the slab) per unit slab area, in the infinite-slab limit, at finite thickness; the outer limit is the dipole per unit thickness, in the infinite-thickness limit. The obvious reason why the limits in Eq. (24) do not commute is the long-range nature of Coulomb interaction; because of this very same reason Madelung sums are conditionally convergent.

One final observation is that Eq. (24) does not seem to yield a multivalued observable; yet in fact it does. The inner limit therein depends on the prescription for terminating the cuboid at the faces normal to $x$. The multivalued nature of macroscopic polarization, when addressed via bounded samples in the large-sample limit, has been indeed demonstrated via ab-initio simulations in the (much simpler) case of a stereoregular quasi-one-dimensional polymer with different terminations.

VII. CONCLUSIONS

The theory of polarization in condensed matter was developed along the 1990s and is now a staple of electronic structure theory. The relevant formulas adopt concepts from quantum geometry, and have no relationship to the (trivial) formula for the dipole of a bounded sample; this owes to the fact that the multiplicative position operator $\mathbf{r}$ is no longer a legitimate quantum-mechanical operator when the boundary conditions of condensed matter physics are adopted.

Here I show that it is possible to alternatively express the dipole of a bounded sample, making no use of the $\mathbf{r}$ operator: such expression has the virtue of being adoptable almost as such even in the case of an unbounded crystalline sample. The resulting formula defines bulk crystalline polarization as a multivalued observable. I also show that, in the special case of a single-determinant many-body wavefunction (either Hartree-Fock or Kohn-Sham), one retrieves the algorithm currently implemented in several electronic-structure codes.

Finally I show that—when the large-crystallite limit is ideally taken—one obtains the value of crystalline polarization only if the limit is taken in the appropriate way. The reason is that the crystalline formula assumes by construction zero macroscopic field, while instead a nonzero macroscopic field (depolarization field) is in general present inside a polarized macroscopic sample in vacuo.
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15We also assume that one of the principal axes of the dielectric tensor is along the P direction.