An ab-initio converse NMR approach for pseudopotentials

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(Dated: May 21, 2010)

We extend the recently developed converse NMR approach [T. Thonhauser, D. Ceresoli, A. Mostofi, N. Marzari, R. Resta, and D. Vanderbilt, J. Chem. Phys. 131, 101101 (2009)] such that it can be used in conjunction with norm-conserving, non-local pseudopotentials. This extension permits the efficient ab-initio calculation of NMR chemical shifts for elements other than hydrogen within the convenience of a plane-wave pseudopotential approach. We have tested our approach on several finite and periodic systems, finding very good agreement with established methods and experimental results.

PACS numbers: 71.15.-m, 71.15.Mb, 75.20.-g, 76.60.Cq

I. INTRODUCTION

The experimental technique of nuclear magnetic resonance (NMR) is a powerful tool to determine the structure of molecules, liquids, and periodic systems. It is thus not surprising that, since its discovery in 1938, NMR has evolved into one of the most widely used methods in structural chemistry.1,2 Unfortunately, one caveat of this successful method is that there is no basic, generally valid “recipe” that allows a unique determination of the structure given a measured spectrum. As a result, for more complex systems the mapping between structure and measured spectrum can be ambiguous.

It had been realized early on that ab-initio calculations could resolve some of these ambiguities and thus greatly aid in determining structures from experimental NMR spectra. For finite systems such as simple molecules, appropriate methods were first developed in the quantum-chemistry community.3 While highly accurate, these methods by construction were unable to calculate NMR shifts of periodic systems, which is important for the increasingly popular solid-state NMR spectroscopy. The underlying physical limitation is due to the fact that the description of any constant external magnetic field requires a non-periodic vector potential. Possible approaches to combine such a non-periodic vector potential with the periodic potential of crystals were found only within the last decade.4–8 All of these approaches have in common that they treat the external magnetic field in terms of the linear-response it causes to the system under consideration. Although these approaches are accurate and successful, the required linear-response framework makes them fairly complex and difficult to implement.

Recently, a fundamentally different approach for the calculation of ab-initio NMR shifts has been developed by some of us.9 In our converse approach we circumvent the need for a linear-response framework in that we relate the shifts to the macroscopic magnetization induced by magnetic point dipoles placed at the nuclear sites of interest. The converse approach has the advantage of being conceptually much simpler than other standard approaches and it also allows us to calculate the NMR shifts of systems with several hundred atoms. Our converse approach has already successfully been applied to simple molecules, crystals, liquids, and extended polycyclic aromatic hydrocarbons.9,10

While the converse method can be directly implemented in an all-electron first-principles computer code, an implementation into a pseudopotential code is more complicated due to nonlocal projectors usually used in the Kleinman-Bylander separable form.11 In this paper we present a mathematical extension to the converse formalism such that it can be used in conjunction with norm-conserving, non-local pseudopotential. This extension permits the efficient ab-initio calculation of NMR chemical shifts for elements other than hydrogen within the convenience of a plane-wave pseudopotential approach.

The paper is organized in the following way: In Sec. II we first review the converse-NMR method and its relation to the orbital magnetization at an all-electron level. Then, we apply the gauge including projector augmented wave (GIPAW) transformation to derive an expression for the orbital magnetization in the context of norm-conserving pseudopotentials. We discuss aspects of the implementation of the converse method in Sec. III. In this section we also show results of several convergence tests that we have performed. To validate our approach, we apply our converse approach to molecules and solids and the results are collected in Sec. IV. In Sec. V we discuss the main advantages of the converse method. Finally, we summarize and conclude in Sec. VI. The GI-PAW transformation and several details of the mathematical formalism—which would be distracting in the main text—are presented in Appendices A and B.

II. THEORY

The converse method for calculating the NMR chemical shielding has been introduced in Ref. [9] and can be
summarized as follows:

\[ \sigma_{s,\alpha\beta} = \delta_{\alpha\beta} - \Omega \frac{\partial M_{\beta}}{\partial m_{s,\alpha}} \]  

(1)

Thus, in the converse method the chemical shielding tensor \( \sigma_{s,\alpha\beta} \) is obtained from the derivative of the orbital magnetization \( M \) with respect to a magnetic point dipole \( m_s \), placed at the site of atom \( s \). \( \delta_{\alpha\beta} \) is the Kronecker delta and \( \Omega \) is the volume of the simulation cell. In other words, instead of applying a constant magnetic field to an infinite periodic system and calculating the induced field at all equivalent sites \( s \), and calculate the change in magnetization. Since the perturbation is now periodic, the original problem of the non-periodic vector potential has been circumvented.

In practice, the derivative in Eq. (1) is calculated as a finite difference of the orbital magnetization in presence of a small magnetic point dipole \( m_s \). Since \( M \) vanishes for \( m_s = 0 \) and is an odd function of \( m_s \), because of time-reversal symmetry (for a non-magnetic system in absence of spin-orbit interaction), it is sufficient to perform three calculations of \( M(m_s,e) \), where \( e \) are cartesian unit vectors.

Within density-functional theory (DFT) the all-electron Hamiltonian is, in atomic units:

\[ \mathcal{H} = \frac{1}{2} \left( p + \frac{1}{c} A_s(r) \right)^2 + V_{KS}(r) \]  

(2)

where

\[ A_s(r) = \frac{m_s \times (r-r_s)}{|r-r_s|^3} \]  

(3)

is the vector potential corresponding to a magnetic dipole \( m_s \), centered at the atom \( s \) coordinate \( r_s \). We neglect any explicit dependence of the exchange-correlation functional on the current density. In practice, spin-current density-functional theory calculations have shown to produce negligible corrections to the orbital magnetization.

In finite systems (i.e. molecules), the orbital magnetization can be easily evaluated via the velocity operator \( \mathbf{v} = -i[\mathbf{r}, \mathcal{H}] \):

\[ \mathbf{M} = \frac{1}{2c} \sum_n <\psi_n| \mathbf{r} \times \mathbf{v} | \psi_n> \]  

(4)

where \( |\psi_n> \) are molecular orbitals, spanning the occupied manifold. For periodic systems, the situation is more complicated due to itinerant surface currents and to the incompatibility of the position operator \( \mathbf{r} \) with periodic boundary conditions. It has been recently shown that the orbital magnetization in a periodic system is given by:

\[ \mathbf{M} = -\frac{1}{2c} \text{Im} \sum_{nk} f(\epsilon_{nk}) |\partial_{\mathbf{k}} u_{nk}| \times (\mathcal{H} + \epsilon_{nk} - 2\epsilon_F) |\partial_{\mathbf{k}} u_{nk}> \]  

(5)

where \( u_{nk} \) are the Bloch wavefunctions, \( \mathcal{H}_k = e^{-ikr} \mathcal{H} e^{ikr} \), \( \epsilon_{nk} \) are its eigenvalues and \( \epsilon_F \) is the Fermi level. The \( k \)-derivative of the Bloch wavefunctions can be evaluated as a covariant derivative, or by \( \mathbf{k} \cdot \mathbf{p} \) perturbation theory.

Equations (2) and (5) are adequate to evaluate the NMR shielding tensor Eq. (1) in the context of an all-electron method (such as FLAPW, or local-basis methods), in which the interaction between core and valence electrons is treated explicitly. However, in a pseudopotential framework, where the effect of the core electrons has been replaced by a smooth effective potential, Eqs. (2) and (5) are not sufficient to evaluate the NMR shielding tensor. One reason for this is that the valence wave functions have been replaced by smoother pseudo wave functions which deviate significantly from the all-electron ones in the core region.

In the following sections, we derive the formulas needed to calculate the converse NMR shielding tensor in Eq. (1), in the context of the pseudopotential method. Our derivation is based on the GIPAW transformation, (see also Appendix A), that allows one to reconstruct all-electron wave functions from smooth pseudopotential wave functions. For the sake of simplicity, we assume all GIPAW projectors to be norm-conserving.

A. The converse-NMR GIPAW hamiltonian

In this section we derive the pseudopotential GIPAW hamiltonian corresponding to the all-electron (AE) hamiltonian in Eq. (2). For reasons that will be clear in the next section, we include an external uniform magnetic field \( \mathbf{B} \) in addition to the magnetic field generated by the point dipole \( m_s \). For the sake of simplicity, we carry out the derivation for an isolated system (i.e. a molecule) in the symmetric gauge \( \mathbf{A}(\mathbf{r}) = (1/2)\mathbf{B} \times \mathbf{r} \). The generalization to periodic systems is then performed at the end.

We start with the all-electron hamiltonian:

\[ \mathcal{H}_{AE} = \frac{1}{2} \left[ p + \frac{1}{c} (\mathbf{A}(\mathbf{r}) + \mathbf{A}_s(\mathbf{r})) \right]^2 + V(\mathbf{r}) \]  

(6)

We now decompose Eq. (6) in powers of \( \mathbf{A}_s \) as \( \mathcal{H}_{AE} = \mathcal{H}_{AE}^{(0)} + \mathcal{H}_{AE}^{(1)} + \mathcal{H}_{AE}^{(2)} \) where

\[ \mathcal{H}_{AE}^{(0)} = \frac{1}{2} \left[ (p + 1/c A_s(\mathbf{r})) \right]^2 + V(\mathbf{r}) \]  

(7)

\[ \mathcal{H}_{AE}^{(1)} = \frac{1}{2c} \left( \mathbf{p} \cdot \mathbf{A}_s(\mathbf{r}) + \mathbf{A}_s(\mathbf{r}) \cdot \mathbf{p} + 2/c A_s(\mathbf{r}) \cdot A_s(\mathbf{r}) \right) \]  

(8)

\[ \mathcal{H}_{AE}^{(2)} = \frac{1}{2c^2} A_s(\mathbf{r})^2 \]  

We can neglect \( \mathcal{H}_{AE}^{(2)} \) in all calculations, since \( m_s \) is a small perturbation to the electronic structure.

We then apply the GIPAW transformation Eq. (4) to the two remaining terms, Eqs. (7) and (8), and we expand the results up to first order in the magnetic field.
At zeroth order in the external magnetic field $\mathbf{B}$, the GIPAW transformation of $\mathcal{H}^{(s0)}_{AE}$ and $\mathcal{H}^{(s1)}_{AE}$ yields the GIPAW Hamiltonian:

$$\mathcal{H}_{\text{GIPAW}} = \mathcal{H}^{(s0,0)}_{\text{GIPAW}} + \mathcal{H}^{(s1,0)}_{\text{GIPAW}}$$

$$\mathcal{H}^{(s0,0)}_{\text{GIPAW}} = \frac{1}{2} \mathbf{p}^2 + V_{\text{loc}}(\mathbf{r}) + \sum_{\mathbf{R}} V^{\text{NL}}_{\mathbf{R}}$$

$$\mathcal{H}^{(s1,0)}_{\text{GIPAW}} = \frac{1}{2} |\mathbf{p} \cdot \mathbf{A}_s(\mathbf{r}) + \mathbf{A}_s(\mathbf{r}) \cdot \mathbf{p}| + \sum_{\mathbf{R}} K^{\text{NL}}_{\mathbf{R}}$$

(9) (10) (11)

where $V_{\text{loc}}(\mathbf{r})$ is the local Kohn-Sham potential and $V^{\text{NL}}_{\mathbf{R}}$ is the non-local pseudopotential in the separable Kleinmann-Bylander (KB) form:

$$V^{\text{NL}}_{\mathbf{R}} = \sum_{nm} |\beta_{\mathbf{R},n} \rangle v_{\mathbf{R},nm} \langle \beta_{\mathbf{R},m}|$$

(12)

Similar to $V^{\text{NL}}_{\mathbf{R}}$, the term $K^{\text{NL}}_{\mathbf{R}}$ has the form of a non-local operator

$$K^{\text{NL}}_{\mathbf{R}} = \frac{1}{2c} \sum_{nm} |\tilde{p}_{\mathbf{R},n} \rangle k_{\mathbf{R},nm} \langle \tilde{p}_{\mathbf{R},m}|$$

(13)

$$k_{\mathbf{R},nm} = \langle \phi_{\mathbf{R},n} | \mathbf{p} \cdot \mathbf{A}_s(\mathbf{r}) + \mathbf{A}_s(\mathbf{r}) \cdot \mathbf{p} | \phi_{\mathbf{R},m} \rangle - \langle \tilde{\phi}_{\mathbf{R},n} | \mathbf{p} \cdot \mathbf{A}_s(\mathbf{r}) + \mathbf{A}_s(\mathbf{r}) \cdot \mathbf{p} | \tilde{\phi}_{\mathbf{R},m} \rangle$$

(14)

(15) (16)

The index $\mathbf{R}$ runs over all atoms in the system, and the indexes $n$ and $m$, individually run over all projectors associated with atom $\mathbf{R}$. For a definition of $|\phi_{\mathbf{R},n} \rangle$ and $|\tilde{\phi}_{\mathbf{R},n} \rangle$ see Appendix A. Note that the set of GIPAW projectors $|\tilde{\phi}_{\mathbf{R},n} \rangle$ need not be the same as the KB projectors $|\beta_{\mathbf{R},n} \rangle$. For instance, in the case of norm-conserving pseudopotentials, one KB projector per non-local channel is usually constructed. Conversely, two GIPAW projectors for each angular momentum channel are usually needed.

Equation (9) is the Hamiltonian to be implemented in order to apply a point magnetic dipole to the system. The first term of $\mathcal{H}^{(s1,0)}_{\text{GIPAW}}$ can be applied to a wave function in real space or in reciprocal space. The second term acts on the wave functions like an extra non-local term and requires very little change to the existing framework that applies the non-local potential.

At the first order in the magnetic field, the GIPAW transformation yields two terms:

$$\mathcal{H}^{(s0,1)}_{\text{GIPAW}} = \frac{1}{2c} \mathbf{B} \cdot \left( \mathbf{L} + \sum_{\mathbf{R}} \mathbf{R} \times \frac{1}{i} [\mathbf{r}, V^{\text{NL}}_{\mathbf{R}}] \right)$$

$$\mathcal{H}^{(s1,1)}_{\text{GIPAW}} = \frac{1}{2c} \mathbf{B} \cdot \left( \mathbf{r} \times \mathbf{A}_s(\mathbf{r}) + \sum_{\mathbf{R}} E^{\text{NL}}_{\mathbf{R}} + \sum_{\mathbf{R}} \mathbf{R} \times \frac{1}{i} [\mathbf{r}, K^{\text{NL}}_{\mathbf{R}}] \right)$$

(15) (16)

where $\mathbf{L} = \mathbf{r} \times \mathbf{p}$ and $E^{\text{NL}}_{\mathbf{R}}$ is the non-local operator

$$E^{\text{NL}}_{\mathbf{R}} = \sum_{nm} |\tilde{p}_{\mathbf{R},n} \rangle e_{\mathbf{R},nm} \langle \tilde{p}_{\mathbf{R},m}|$$

$$e_{\mathbf{R},nm} = \langle \phi_{\mathbf{R},n} | (\mathbf{r} - \mathbf{R}) \times \mathbf{A}_s(\mathbf{r}) | \phi_{\mathbf{R},m} \rangle - \langle \tilde{\phi}_{\mathbf{R},n} | (\mathbf{r} - \mathbf{R}) \times \mathbf{A}_s(\mathbf{r}) | \tilde{\phi}_{\mathbf{R},m} \rangle$$

(17) (18)

The two equations above will be used in the next section, in conjunction with the Hellmann-Feynman theorem, to derive the GIPAW form of the orbital magnetization.

**B. The orbital magnetization in the GIPAW formalism**

The orbital magnetization for a non spin-polarized system is formally given by the Hellmann-Feynman theorem as

$$M = - \left\langle \frac{\partial \mathcal{H}_{AE}}{\partial \mathbf{B}} \right\rangle_{B=0}$$

In the GIPAW formalism this expectation value can be expressed in terms of the GIPAW Hamiltonian and pseudo wave functions

$$M = - \left\langle \frac{\partial \mathcal{H}_{\text{GIPAW}}}{\partial \mathbf{B}} \right\rangle_{B=0}$$

(19) (20)

By using the results of the previous section we find:

$$M = \sum_{n \text{occ}} \langle \psi_n | [\mathbf{L} + \mathbf{r} \times \mathbf{A}_s(\mathbf{r}) + \sum_{\mathbf{R}} E^{\text{NL}}_{\mathbf{R}} + \sum_{\mathbf{R}} \mathbf{R} \times \frac{1}{i} [\mathbf{r}, K^{\text{NL}}_{\mathbf{R}}], \mathbf{r} \rangle | \psi_n \rangle$$

(21)

Note that in the expression above, $\psi_n$ are the eigenstates of the GIPAW Hamiltonian in absence of any external magnetic fields.

While the formula (21) for $M$ can directly be applied to atoms and molecules, it is ill-defined in the context of periodic systems, owing to the presence of the position operator—explicitly in $\mathbf{r}$, but also implicitly as in $\mathbf{L}$. This problem can be remedied by applying the modern theory of orbital magnetization.14-17 The goal is thus to reformulate Eq. (21) in terms of $\langle \mathbf{r} \times v_{\text{GIPAW}} \rangle$. We can calculate this operator as

$$\mathbf{r} \times v_{\text{GIPAW}} = \mathbf{r} \times \frac{1}{i} \left( \partial \mathcal{H}_{\text{GIPAW}} \right)_{B=0} = \mathbf{L} + \mathbf{r} \times \mathbf{A}_s(\mathbf{r}) + i \sum_{\mathbf{R}} \mathbf{r} \times [V^{\text{NL}}_{\mathbf{R}} + K^{\text{NL}}_{\mathbf{R}}, \mathbf{r}]$$

(22)

Replacing $\mathbf{L}$ in Eq. (21) by the corresponding expression calculated from Eq. (22), and regrouping the terms, we obtain the central result of this paper:

$$M = M_{\text{bare}} + M_{\text{NL}} + M_{\text{para}} + M_{\text{dia}}$$

$$M_{\text{bare}} = - \frac{1}{2c} \langle \mathbf{r} \times v_{\text{GIPAW}} \rangle$$

$$M_{\text{NL}} = - \frac{1}{2c} \left( \sum_{\mathbf{R}} \mathbf{r} \times [\mathbf{r} - \mathbf{R}, V^{\text{NL}}_{\mathbf{R}}] \right)$$

$$M_{\text{para}} = - \frac{i}{2c} \left( \sum_{\mathbf{R}} \mathbf{r} \times [\mathbf{r} - \mathbf{R}, K^{\text{NL}}_{\mathbf{R}}] \right)$$

$$M_{\text{dia}} = - \frac{1}{2c} \left( \sum_{\mathbf{R}} E^{\text{NL}}_{\mathbf{R}} \right)$$

(23) (24) (25) (26)
where \( \langle \ldots \rangle \) stands for \( \sum_{n}^{\text{occ}} \langle \psi_n | \ldots | \psi_n \rangle \). The naming of the various terms are in analogy to Ref. [6]. The set of equations (23)–(26) are now valid both in isolated and periodic systems, as shown in detail in Appendix B.

### III. IMPLEMENTATION AND COMPUTATIONAL DETAILS

We have implemented the converse NMR method and its GIPAW transformation into PWscf, which is part of the QUANTUM-ESPRESSO package.

In principle, the calculation of the NMR shielding is performed the following way: The vector potential corresponding to the microscopic dipole is included in the Hamiltonian and the Kohn-Sham equation is solved self-consistently under that Hamiltonian. In practice, however, in a first step one can equally as well find the ground state of the unperturbed system. Based on this ground state, in the second step one can then introduce the dipole perturbation and converge to the new ground state. Note that the convergence of the small dipole perturbation is usually very fast and only a small number of SCF steps is necessary in addition to the ground state calculation. In fact, tests have shown that a convergence is not even necessary—diagonalizing the perturbed Hamiltonian only once with the unperturbed wave functions gives results for the NMR shielding within 0.01 ppm of the fully converged solution. This yields a huge calculational benefit for large systems, where we calculate the unperturbed ground state once and then, based on the converged ground-state wave functions, calculate all shieldings of interest by non-SCF calculations.

In order to study the convergence of the NMR chemical shielding with several parameters, we performed simple tests on a water molecule in the gas phase. The molecule was relaxed in a box of 30 Bohr; for all our calculations we used Troullier-Martin norm-conserving pseudopotentials\(^\text{21}\) and a PBE exchange-correlation functional\(^\text{22}\).

First, we tested the convergence of the NMR shielding with respect to the kinetic-energy cutoff \( E_{\text{kin}} \) and the results are presented in Table I. The shielding \( \sigma \) is converged to within 0.02 ppm for a kinetic-energy cutoff of 80 Ryd. Similar tests on other structures show similar results.

Next, we tested the convergence of the NMR shielding with respect to the magnitude of the microscopic dipole \( |m_s| \) and the energy convergence criterion. At first sight it might appear difficult to accurately converge the electronic structure in the presence of a small microscopic magnetic point dipole. Thus, we tested using different magnitudes for the microscopic dipole spanning several orders of magnitude from \( 10^{-5} \mu_B \) (which is actually much less than the value of a core spin) to \( 10^3 \mu_B \) (which is obviously much more than an electron spin). On the other hand, the ability to converge the electronic structure accurately goes hand in hand with the energy convergence criterion \( E_{\text{conv}} \) that is used in such calculations. This criterion is defined such that the calculation is considered converged if the energy difference between two consecutive SCF steps is smaller than \( E_{\text{conv}} \). The results for the shielding as a function of \( |m_s| \) and \( E_{\text{conv}} \) are collected in Table II. It is interesting to see that it is just as simple to converge with a small dipole than it is to converge with a large dipole. In either case, using at least \( E_{\text{conv}} = 10^{-4} \) Ryd yields results converged to within 0.1 ppm. Such a convergence criterion is not even particularly “tight” and most standard codes use at least \( E_{\text{conv}} = 10^{-6} \) Ryd as default. Note that first signs of non-linear effects appear if large dipoles such as \( |m_s| = 100 \mu_B \) or \( |m_s| = 1000 \mu_B \) are used. In conclusion of the above tests, we use \( |m_s| = 1 \mu_B, E_{\text{conv}} = 10^{-8} \) Ryd, and \( E_{\text{kin}} = 100 \) Ryd for all calculations.

### A. Generation of GIPAW pseudopotentials

We have generated special-purpose norm-conserving pseudopotentials for our GIPAW calculations. In addition to standard norm-conserving pseudopotentials (PS), the GIPAW pseudopotentials include (i) the full set of AE core atomic functions and (ii) the AE \( \langle \phi_n \rangle \) and the PS \( \langle \bar{\phi}_n \rangle \) valence atomic orbitals. The core orbitals contribute to the isotropic NMR shielding is:

\[
\sigma_{\text{core}} = \frac{1}{2c} \sum_{n \in \text{core}} 2(2l_n + 1) \langle \phi_n | \frac{1}{r} | \phi_n \rangle
\]

The AE and PS valence orbitals are used to compute the coefficients \( k_{R,nn} \) and \( e_{R,nn} \) at the beginning of the calculation. The PS valence orbitals are also used to compute the GIPAW projectors \( \bar{\rho}_{R} \) from:

\[
\bar{\rho}_{R,n} = \sum_m (S^{-1})_{nm} |\bar{\phi}_{R,m}\rangle
\]

\[
S_{nm} = \langle \bar{\phi}_{R,n} | \bar{\rho}_{R,m} | \bar{\rho}_{R,m} \rangle_{H_c}
\]

\( S \) is the overlap between atomic PS wave function, integrated up to the cutoff radius of the corresponding pseudopotential channel.

We construct at least two projectors per angular momentum channel by combining each valence orbital with one excited state with the same angular momentum. For
TABLE II: Chemical shielding in ppm of a hydrogen atom in a water molecule. The shielding is given as a function of the magnitude of the microscopic dipole $|\mathbf{m}_i|$ (in units of Bohr magneton $\mu_B$) and the energy convergence criterion $E_{\text{conv}}$ (in units of Rydberg). $E_{\text{conv}}$ is defined such that the calculation is considered converged if the energy difference between two consecutive SCF steps is smaller than $E_{\text{conv}}$.

| $|\mathbf{m}_i| (\mu_B)$ | $E_{\text{conv}}$ (Ry) |
|-------------------------|----------------------|
|                         | $10^{-2}$ | $10^{-3}$ | $10^{-4}$ | $10^{-5}$ | $10^{-6}$ | $10^{-7}$ | $10^{-8}$ | $10^{-9}$ | $10^{-10}$ |
| 0.00001                 | 31.5170  | 31.2541  | 31.2541  | 31.1354  | 31.1403  | 31.1338  | 31.1356  | 31.1356  | 31.1356    |
| 0.0001                  | 31.5265  | 31.2664  | 31.2664  | 31.1392  | 31.1390  | 31.1322  | 31.1300  | 31.1300  | 31.1300    |
| 0.001                   | 31.5253  | 31.2667  | 31.2667  | 31.1395  | 31.1397  | 31.1328  | 31.1304  | 31.1304  | 31.1304    |
| 0.01                    | 31.5251  | 31.2665  | 31.2665  | 31.1395  | 31.1394  | 31.1325  | 31.1303  | 31.1303  | 31.1303    |
| 0.1                     | 31.5250  | 31.2664  | 31.2664  | 31.1393  | 31.1393  | 31.1323  | 31.1301  | 31.1301  | 31.1301    |
| 1.0                     | 31.5250  | 31.2664  | 31.2664  | 31.1393  | 31.1393  | 31.1323  | 31.1301  | 31.1301  | 31.1301    |
| 10.0                    | 31.5250  | 31.2664  | 31.2663  | 31.1392  | 31.1392  | 31.1322  | 31.1301  | 31.1301  | 31.1301    |
| 100.0                   | 31.5212  | 31.2586  | 31.2586  | 31.1350  | 31.1327  | 31.1243  | 31.1256  | 31.1256  | 31.1252    |
| 1000.0                  | 31.0167  | 30.5904  | 30.7197  | 30.6618  | 30.6334  | 30.6408  | 30.6403  | 30.6404  | 30.6405    |

TABLE III: Electronic configuration and cutoff radii for the norm-conserving pseudopotentials used in the present work.

| Atom | configuration | $r_c$(s) | $r_c$(p) | $r_c$(d) |
|------|---------------|----------|----------|----------|
| H    | 1s$^2$        | 0.50     |          |          |
| B    | [He]2s$^2$2p$^1$ | 1.40     | 1.40     |          |
| C    | [He]2s$^2$2p$^2$ | 1.50     | 1.50     |          |
| N    | [He]2s$^2$2p$^3$ | 1.45     | 1.45     |          |
| O    | [He]2s$^2$2p$^4$ | 1.40     | 1.40     |          |
| F    | [He]2s$^2$2p$^5$ | 1.30     | 1.30     |          |
| P    | [Ne]3s$^2$3p$^2$5$^3$d$^0$ | 1.90     | 2.10     | 2.10     |
| Si   | [Ne]3s$^2$3p$^1$3$^3$d$^0$ | 2.00     | 2.00     | 2.00     |
| Cl   | [Ne]3s$^{1.75}$3p$^{4.5}$5d$^0$.25 | 1.40     | 1.40     | 1.40     |
| Cu   | [Ar]4s$^2$4p$^6$3d$^{10}$ | 2.05     | 2.20     | 2.05     |

example, for hydrogen we include the 2s orbital in the set of atomic wave functions. For all second row elements, we add the 3s and 3p orbitals, and so on. If any excited state turns out to be unbound (as in the case of oxygen and fluorine), we generate an atomic wave function as a scattering state at an energy 0.5 Ry higher than the corresponding valence state. This procedure ensures that the GIPAW projectors are linearly independent and that the matrix $S$ is not singular.

We found that the accuracy of the calculated NMR chemical shifts depends critically on the cutoff radii of the pseudopotentials. Whereas the total energy and the molecular geometry converge more quickly with respect to reducing the pseudopotential radii, the NMR chemical shift converges more slowly. Therefore, GIPAW pseudopotentials have to be generated with smaller radii compared to the pseudopotentials usually employed for total energy calculations. Table III reports the atomic configuration and the cutoff radii used to generate the pseudopotentials.

IV. RESULTS

In this section we present results for molecules and solids. We first calculated the absolute shielding tensor of some small molecules by two different approaches, the direct (linear response) and the converse method, in order to check that the two yield the same results. Then, we compared the chemical shifts of fluorine compounds, calculated by the converse method and by all-electron large basis set quantum-chemistry calculations. Finally, we report the calculated $^{29}$Si chemical shifts of three SiO$_2$ polymorphs and the Cu shift of a metalorganic compound.

A. Small molecules

We calculated the chemical shift of hydrogen, carbon, fluorine, phosphorus, and silicon atoms of various small molecules. First, the structures were relaxed using PWSCF in a box of 30 Bohr and a force convergence threshold of $10^{-4}$ Ry/Bohr. Using the relaxed positions, the chemical shifts were calculated using both the direct and converse method, and the results are shown in Table IV. This benchmark calculation shows that the direct and the converse methods agree to within less than 1%.

B. Fluorine compounds

In structural biology $^{19}$F NMR spectroscopy plays an important role in determining the structure of protein membranes. The advantage over $^{15}$N and $^{17}$O labeling is twofold: the natural abundance of $^{19}$F is nearly 100%, and $^{19}$F has spin 1/2, i.e. a vanishing nuclear quadrupole moment. Quadrupole interactions in high-spin nuclei (e.g. $^{17}$O) are responsible for the broadening of the NMR spectrum. On the contrary, $^{19}$F NMR yields...
TABLE IV: Results for the absolute NMR shielding $\sigma$ in ppm of small molecules for the direct and converse methods. The core contribution to the shielding is also shown.

| molecule | direct | converse | core |
|----------|--------|----------|------|
| **H shielding** | | | |
| CH$_4$ | 30.743 | 30.670 | 0.0 |
| C$_6$H$_6$ | 22.439 | 22.403 | 0.0 |
| SiH$_4$ | 27.444 | 27.413 | 0.0 |
| TMS | 30.117 | 30.125 | 0.0 |
| **C shielding** | | | |
| CH$_4$ | 185.435 | 186.027 | 200.333 |
| C$_6$H$_6$ | 36.887 | 37.205 | 200.333 |
| CH$_3$F | 93.704 | 94.250 | 200.333 |
| TMS | 175.774 | 176.094 | 200.333 |
| **F shielding** | | | |
| CH$_3$F | 448.562 | 447.014 | 305.815 |
| PF$_3$ | 277.148 | 275.819 | 305.815 |
| SiF$_4$ | 378.857 | 376.227 | 305.815 |
| SiH$_3$F | 423.456 | 422.253 | 305.815 |
| **P shielding** | | | |
| P$_2$ | -323.566 | -320.201 | 908.854 |
| PF$_3$ | 150.603 | 150.856 | 908.854 |
| **Si shielding** | | | |
| SiF$_4$ | 431.438 | 432.495 | 837.913 |
| SiH$_3$F | 337.648 | 337.677 | 837.913 |
| Si$_2$H$_4$ | 230.830 | 230.489 | 837.913 |
| TMS | 320.958 | 320.636 | 837.913 |

very sharp and resolved lines. In addition, it has been found that the substitution of $\text{CH}_3$ groups with $\text{CF}_3$ in some amino-acids does not perturb the structure and the activity of protein membranes, allowing for in vivo NMR measurements.

In order to benchmark the accuracy of our method, we calculated the $^{19}\text{F}$ chemical shifts of ten fluorine compounds utilizing the converse method, and we compared our results to all-electron gaussian-basis set calculations, as well as to experimental data. The molecules were first relieved with Gaussian03, with the 6-311+g(2d,p) basis set at the B3LYP level. Then, we calculated the IGAIM chemical shift with Gaussian03, with the cc-pVTZ, cc-pVQZ, cc-pV5Z and cc-pV6Z basis sets.

To calculate the relative chemical shifts, we used C$_6$F$_6$ as a secondary reference compound, and we used the experimental C$_6$F$_6$ chemical shift, to get the primary reference absolute shift (CF$_3$Cl). The results are shown in Table V and in Fig. 1. While compiling Table V we suspected that the experimental values of p-C$_6$H$_4$F$_2$ and C$_6$H$_2$F have been mistakenly exchanged in Ref. [26]. A quick inspection of the original paper, confirmed our suspicion. The overall agreement of the converse method with experimental data is very good, and of the same quality as the cc-pV6Z basis set, which comprises 140 basis functions for 2nd row atoms. The calculation time required by our plane-wave converse method is comparable to that of cc-pV5Z calculations.

C. Solids

In this section we present the $^{29}\text{Si}$ and $^{17}\text{O}$ chemical shifts calculated by our converse method in four SiO$_2$ polymorphs: quartz, $\beta$-cristobalite, coesite and stishovite. Coesite and stishovite are metastable phases that form at high temperature and pressures developed during a meteor impact. Besides their natural occurrence in meteorite, they can also be artificially synthesized by shock experiments.

We adopted the experimental crystal structures and atom positions in all calculations. We used a cutoff of 100 Ryd and a k-point mesh of $8 \times 8 \times 8$ for quartz, $\beta$-cristobalite and stishovite. In the case of coesite, having the largest primitive cell (48 atoms), we used a k-point mesh of $4 \times 4 \times 4$. In Table VI we show a comparison between the calculated and the experimental chemical shifts for the four crystals. We determined the $^{29}\text{Si}$ and $^{17}\text{O}$ reference shielding as the intercept of the least-square linear interpolation of the ($\sigma_{\text{calc}}, \delta_{\text{expt}}$) pairs. Note that the nuclear magnetic dipole $\mathbf{m}_s$ breaks the symmetry of the hamiltonian. Thus, we retained only the symmetry operations that map site $s$ to $s'$ without changing the orientation of the magnetic dipole (i.e. $s \rightarrow s', \mathbf{m}_s = \mathbf{m}_{s'}$).

Another important point is that in periodic systems we are not just including one nuclear dipole, but rather
TABLE V: Experimental and calculated $^{19}$F chemical shifts in ppm, with respect to CF$_3$Cl. In all calculations (Gaussian03 and GIPAW plane waves), we used C$_6$F$_6$ as the reference compound. The experimental values of p-C$_6$H$_4$F$_2$ and C$_6$H$_5$F are exchanged with respect to Ref. [26]. For molecules with inequivalent F atoms, the average chemical shift is reported. MAE is the mean absolute error in ppm, with respect to experiment.

| Molecule | Expt.$^{26}$ | Gaussian cc-pVTZ | Gaussian cc-pVQZ | Gaussian cc-pV5Z | Gaussian cc-pV6Z | GIPAW converse |
|----------|--------------|------------------|------------------|------------------|------------------|----------------|
| CH$_2$FCN | -251 | -253.07 | -253.47 | -254.25 | -254.79 | -258.31 |
| C$_6$F$_6$ | -164.9 | -164.90 | -164.90 | -164.90 | -164.90 | -164.90 |
| BF$_3$ | -131.3 | -145.93 | -139.55 | -136.37 | -135.49 | -135.65 |
| p-C$_6$H$_4$F$_2$ | -113.15 | -115.77 | -113.98 | -114.07 | -113.78 | -111.84 |
| C$_6$H$_5$F | -106 | -106.84 | -104.94 | -104.83 | -104.35 | -104.68 |
| (CF$_3$)$_2$CO | -84.6 | -90.37 | -82.12 | -78.81 | -77.83 | -76.63 |
| CF$_3$COOH | -76.55 | -87.82 | -81.38 | -77.88 | -76.80 | -75.90 |
| C$_6$H$_5$CF$_3$ | -63.72 | -78.24 | -69.42 | -66.21 | -65.28 | -64.16 |
| CF$_4$ | -62.5 | -74.48 | -73.94 | -68.76 | -66.66 | -66.05 |
| F$_2$ | 422.92 | 367.92 | 375.36 | 383.03 | 385.72 | 390.02 |
| MAE | - | 13.19 | 9.40 | 7.35 | 6.69 | 5.64 |

TABLE VI: Calculated and experimental $^{29}$Si and $^{17}$O NMR chemical shifts of four SiO$_2$ crystals in ppm. Experimental data was taken from Refs. [28] and [29]. In the case of coesite all inequivalent chemical shifts are reported.

| Mineral | Calc. $\delta$ [ppm] | Expt. $\delta$ [ppm] |
|---------|----------------------|---------------------|
| quartz | -107.10 | -107.73 |
| $\beta$-cristobalite | -108.78 | -108.50 |
| stishovite | -184.13 | -191.33 |
| coesite | -107.30 | -113.35 |
| | -113.35 | -114.33 |
| quartz | -74.48 | 43.52 |
| $\beta$-cristobalite | 40.35 | 37.2 |
| stishovite | 116.35 | N/A |
| coesite | 26.35 | 29 |
| | 39.66 | 41 |
| | 52.70 | 53 |
| | 56.84 | 57 |
| | 59.03 | 58 |

D. Large Systems

Reactive sites in biological systems such as organometallic molecules, as well as inorganic materials, are of great importance. In particular, there is a surge of interest in studying copper(I) reactive sites using solid-state NMR. NMR experiments on these materials are challenging because of the large nuclear quadrupole moments of $^{63}$Cu and $^{65}$Cu. Here, we present the results for the copper-phosphine metallocene, tetramethylcyclopentadienyl copper(I) triphenylphosphine (CpCuPPh$_3$), which as a solid contains 228 atoms in a primitive orthorhombic unit cell. The molecular structure is shown in Fig. 2. The properties of the shielding tensor for the copper environment were observed experimentally for the solid material, and simulated using quantum-chemical methods on the molecular complex.$^{30}$

While the converse approach can calculate the chemical shift for this large system easily, it is more challenging for the linear-response method, which in our experience took much longer in general, did not finish at all, or was unable to handle such large systems. We calculated the copper chemical shift for CpCuPPh$_3$ using the converse method with an energy cutoff of 80 Ry in the self-consistent step and PBE pseudopotentials. While previous quantum-chemical calculations were able to reasonably reproduce the experimental span (1300 ppm) and the skew (0.95) of the chemical shielding tensor, they were not able to calculate the chemical shift itself (0 ppm relative to copper (I) chloride), with an inaccuracy of several hundred ppm.$^{30}$ In addition to yielding excellent agreement with experiment to within 2 ppm for the chemical shift, our calculations also gave good results for the span (1038 ppm) and the skew (0.82) of the chemical shielding tensor.
FIG. 2: (Color online) The molecular structure of the metalloocene, tetramethylcyclopentadienyl copper(I) triphenylphosphine (CpCuPPh3). The copper is shown in gold (light gray) forming the metallocene bond, while the phosphorus is green (gray). The crystal structure of this material consists of a 228-atom orthorhombic unit cell.

V. DISCUSSION

The results presented in the previous sections and in a previous work\textsuperscript{9} show that DFT is able to predict accurately the chemical shift of molecules and solids. In general, we expect this to be true for any weakly-correlated system, well described by the generalized-gradient approximation (GGA). In addition to that, relativistic corrections to the NMR chemical shifts are negligible for all light elements in the periodic table, and become important starting from 4th row elements.

However, there will always exist “difficult” cases in which relativistic corrections cannot be neglected and/or one has to go beyond DFT with standard local functionals. This is an active field of research in quantum chemistry\textsuperscript{31,32} and today it is customary to compute NMR chemical shifts with semi-local hybrid DFT functionals (such as B3LYP). Most quantum-chemistry codes allow the inclusion of relativistic effects (spin orbit) by perturbation theory; furthermore, fully-relativistic (four component) solutions of the Dirac-Breit equation have recently been implemented.\textsuperscript{31}

To the best of our knowledge, all existing ab-initio codes, calculate NMR shifts by perturbation theory. Among them, localized-basis sets are the most popular choice to expand wave functions. This leads to very complicated mathematical expressions and to gauge-dependent results. Only two plane-wave, linear-response implementations\textsuperscript{5,6} have been reported. Our converse-

NMR method is built on Mauri's GIPAW method, but has the advantage of circumventing the need for a linear response framework.

The main advantage of our converse-NMR method is that it requires only the ground state wave functions and Hamiltonian to calculate the orbital magnetization. Since no external magnetic field is included in the calculation, our method solves the gauge-origin problem. Moreover, “difficult cases” can be treated easily by our converse method, provided that relativistic corrections and many-body effects are included in the Hamiltonian. Thus, one can concentrate effectively all efforts in developing advanced post-DFT theories (i.e. DFT+U, DMFT, hybrid functionals, self-interaction-free methods) and benchmark them against NMR experiments.

VI. SUMMARY

In this paper we have generalized the recently developed converse NMR approach\textsuperscript{9} such that it can be used in conjunction with norm-conserving, non-local pseudopotentials. We have tested our approach both in finite and periodic systems, on small molecules, four silicate minerals and a molecular crystal. In all cases, we have found very good agreement with established methods and experimental results.

The main advantage of the converse-NMR method is that it requires only the ground state wave functions and Hamiltonian, circumventing the need of any linear response treatment. This is of paramount importance for the rapid development and validation of new methods that go beyond DFT.

Currently, we are applying the converse NMR method to study large biological systems such as nucleic acids\textsuperscript{33} and drug-DNA interactions\textsuperscript{34,35} in conjunction with a recently developed van der Waals exchange-correlation functional.\textsuperscript{36,37} We are also exploring the possibility to calculate non-perturbatively the Knight shift in metals. Finally, the converse method can be used to calculate the EPR g-tensor in molecules and solids.\textsuperscript{38}

VII. ACKNOWLEDGMENTS

This work was supported by the DOE/SciDAC Institute on Quantum Simulation of Materials and Nanостructures. All computations were performed on the Wake Forest University DEAC Cluster with support from the Wake Forest University Science Research Fund. D.C. acknowledges partial support from ENI.
Appendix A: The GIPAW transformation

The starting point is the projector augmented wave (PAW) transformation:  
\[
T = 1 + \sum_{\mathbf{R},n} \left( |\phi_{\mathbf{R},n}| - |\tilde{\phi}_{\mathbf{R},n}| \right) \langle \tilde{\phi}_{\mathbf{R},n} | (A1)
\]
which connects an all-electron wave function \( |\psi\rangle \) to the corresponding pseudopotential wave function \( |\tilde{\psi}\rangle \) via: 
\[|\psi\rangle = T |\tilde{\psi}\rangle.\] 
Here, \(|\phi_{\mathbf{R},n}\rangle\) are all-electron partial waves, \(|\tilde{\phi}_{\mathbf{R},n}\rangle\) are pseudopotential partial waves, and \(|\tilde{\phi}_{\mathbf{R},n}\rangle\) are PAW projectors. The sum runs over the atom positions \(\mathbf{R}\). \(n\) is a combined index that runs over the set of projectors attached to atom \(\mathbf{R}\). In the original PAW formalism, there are two sets of projectors per angular momentum channel \((0, \ldots, l_{\text{max}})\), each with \((2l+1)\) projectors for a total of \(2(l_{\text{max}} + 1)^2\) PAW projectors.

The expectation value of an all-electron operator \(O_{\text{AE}}\) between all-electron wave functions can then be expressed as the expectation value of a pseudo operator \(O_{PS}\) between pseudo wave functions as \(\langle \psi | O_{\text{AE}} | \psi \rangle = \langle \tilde{\psi} | O_{PS} | \tilde{\psi} \rangle\), where the \(O_{PS}\) is given by:
\[
O_{PS} \equiv T^+ O_{\text{AE}} T =
O_{\text{AE}} + \sum_{\mathbf{R},n,m} |\tilde{\phi}_{\mathbf{R},n}\rangle \left( \langle \phi_{\mathbf{R},n} | O_{\text{AE}} | \phi_{\mathbf{R},m} \rangle - \langle \phi_{\mathbf{R},n} | O_{\text{AE}} | \tilde{\phi}_{\mathbf{R},m} \rangle \right) |\tilde{\phi}_{\mathbf{R},m}\rangle (A2)
\]
In the presence of external magnetic fields the PAW transformation is no longer invariant with respect to translations (except in the very simple case of only one augmentation region). This deficiency was resolved by Mauri et al. who developed the GIPAW (“gauge including PAW”) method, which is similar to the PAW transformation from Eq. (A2) but with the inclusion of phase factor compensating the gauge term arising from the translation of a wave function in a magnetic field. The GIPAW transformation in the symmetric gauge reads:
\[
T_G = 1 + \sum_{\mathbf{R},n} e^{i(2\pi/c) \mathbf{R} \times \mathbf{B}} \left( |\phi_{\mathbf{R},n}| - |\tilde{\phi}_{\mathbf{R},n}| \right). 
\]
\[
|\tilde{\phi}_{\mathbf{R},n}| e^{-i(2\pi/c) \mathbf{R} \times \mathbf{B}} (A3)
\]
and the corresponding pseudopotential operator \(O_{PS}\) is:
\[
O_{PS} \equiv T_G^+ O_{\text{AE}} T_G = O_{\text{AE}} + \sum_{\mathbf{R},n,m} e^{i(2\pi/c) \mathbf{R} \times \mathbf{B}} |\tilde{\phi}_{\mathbf{R},n}\rangle \left( \langle \phi_{\mathbf{R},n} | O_{\text{AE}} | \phi_{\mathbf{R},m} \rangle - \langle \phi_{\mathbf{R},n} | O_{\text{AE}} | \tilde{\phi}_{\mathbf{R},m} \rangle \right) |\tilde{\phi}_{\mathbf{R},m}\rangle (A4)
\]
In the following we will refer to the often occurring part \(O_{\text{AE}} = e^{-(i/2c) \mathbf{r} \times \mathbf{B}} O_{\text{AE}} e^{i(2\pi/c) \mathbf{r} \times \mathbf{B}}\) as inner operator and denote it with a hat. The above expression allows us to calculate accurate expectation values of operators within a pseudopotential approach. In this work we carry out the derivation working in the symmetric gauge \(\mathbf{A}(\mathbf{r}) = (1/2)\mathbf{B} \times \mathbf{r}\). This is not an issue, since all physical quantities we are working with, are gauge invariant.

One useful property of the GIPAW transformation is:
\[
e^{-i(2\pi/c) \mathbf{r} \times \mathbf{B}} \left[ \mathbf{p} + \frac{1}{c} \mathbf{A}(\mathbf{r}) \right]^n e^{i(2\pi/c) \mathbf{r} \times \mathbf{B}} =
\left[ \mathbf{p} + \frac{1}{c} \mathbf{A}(\mathbf{r} - \mathbf{R}) \right]^n (A5)
\]

Appendix B: Periodic systems

Note that the set of equations (23) are well defined also in periodic boundary conditions. In fact, \(M_{\text{bare}}\) can be calculated by the Modern Theory of the Orbital Magnetization14–17 as:
\[
M_{\text{bare}} = -\frac{1}{2c} \text{Im} \sum_{nk} f(\epsilon_{nk}) \langle \partial_k u_{nk} | \langle \mathcal{H}_k + \epsilon_{nk} - 2\epsilon_F | \partial_k u_{nk} \rangle \rangle_{\text{GIPAW}} \] (B1)
\]
where \(\mathcal{H}_k\) is the GIPAW hamiltonian, \(\epsilon_{nk}\) and \(|u_{nk}\rangle\) are its eigenvalues and eigenvectors.

The position operator appearing in the other terms in Eq. (23) is well defined because the projectors \(|\tilde{\phi}_{\mathbf{R},n}\rangle\) and \(|\tilde{\phi}_{\mathbf{R},n}\rangle\) are non-vanishing only inside an augmentation sphere, centered around atom \(\mathbf{R}\). The expression of \(M_{\text{NL}}\), \(M_{\text{para}}\), and \(M_{\text{para}}\), can be further manipulated in order to work with Bloch wave functions. In the following part, we show it only for \(M_{\text{NL}}\), since \(M_{\text{para}}\) is similar. The term \(M_{\text{dia}}\) can be manipulated instead in a trivial way.
Let’s consider the expectation value of
\[(r - R) \times \langle r - R, V_{NL}^{RL} \rangle = -(r - R) \times \langle V_{NL}^{RL} \rangle (r - R)\]  
(B2)
on a Bloch state \(|\psi_{nk}\rangle = e^{ikr}|u_{nk}\rangle\):
\[- \sum \langle u_{nk}| e^{-ikr} (r - R) \times \langle V_{NL}^{RL} \rangle (r - R) e^{ikr}|u_{nk}\rangle = \]
\[- \sum_{L \tau} \sum_{ij} \langle u_{nk}| e^{-ikr} (r - L - \tau) |\beta_{L+\tau, i}\rangle \times v_{ij}^{L+\tau} \langle \beta_{L+\tau, j} | (r - L - \tau) e^{ikr}|u_{nk}\rangle, \]  
(B4)
where \(L\) are the real space lattice vectors (not to be confused with the angular momentum operator) and \(\tau\) are the position of the atoms in the unit cell. Inserting two canceling phase factors:
\[\cdots = - \sum_{L \tau} \sum_{ij} \langle u_{nk}| e^{-ik(r-L-\tau)} (r - L - \tau) |\beta_{L+\tau, i}\rangle \times v_{ij}^{L+\tau} \langle \beta_{L+\tau, j} | (r - L - \tau) e^{ik(r-L-\tau)} |u_{nk}\rangle \]  
(B5)
one can recognize immediately the \(k\) derivative of the KB projectors. In addition, since the KB projectors vanish outside their augmentation regions, it is possible to insert a second sum over \(L'\) running on the right hand side of the cross product:
\[\cdots = - \sum_{L L' \tau} \sum_{ij} \langle u_{nk}| (-1/i) \partial_k \left( e^{-ik(r-L-\tau)} |\beta_{L+\tau, i}\rangle \right) \times v_{ij}^{L+\tau} (1/i) \partial_k \left( \langle \beta_{L' + \tau, j} | e^{ik(r-L-\tau)} \right) |u_{nk}\rangle \]  
(B6)
\[\cdots = - \sum_{\tau} \sum_{ij} \langle u_{nk}| \partial_k \left( \sum_{L} e^{-ik(r-L-\tau)} |\beta_{L+\tau, i}\rangle \right) \times v_{ij}^{\tau+\tau} \partial_k \left( \sum_{L'} \langle \beta_{L' + \tau, j} | e^{ik(r-L'-\tau)} \right) |u_{nk}\rangle \]  
(B7)
In periodic systems the structure factors can be absorbed by the projectors:
\[V_{NL}^{k} = \sum_{\tau} \sum_{ij} |\beta_{\tau, i}^{k}\rangle \langle \beta_{\tau, j}^{k}| \]  
(B8)
\[|\beta_{\tau, i}^{k}\rangle = \sum_{L} e^{-ik(r-L-\tau)} |\beta_{L+\tau, i}\rangle \]  
(B9)
Finally:
\[M_{NL} = \frac{1}{2c} \sum_{nk} \sum_{\tau \sigma} \sum_{ij} \langle u_{nk}| \partial_k \beta_{\tau, i}^{k} \rangle \times v_{\tau, ij} \langle \beta_{\tau, j}^{k}| u_{nk}\rangle \]  
(B10a)
\[M_{para} = \frac{1}{2c} \sum_{nk} \sum_{\tau \sigma} \sum_{ij} \langle u_{nk}| \partial_k \beta_{\tau, i}^{k} \rangle \times k_{\tau, ij} \langle \beta_{\tau, j}^{k}| u_{nk}\rangle \]  
(B10b)
\[M_{dia} = -\frac{1}{2c} \sum_{nk} \langle u_{nk}| \hat{p}_{\tau, i}^{k} \rangle e_{\tau, ij} \langle \hat{p}_{\tau, j}^{k}| u_{nk}\rangle \]  
(B10c)
This completes the main result and it allows us to calculate the orbital magnetization in the presence of non-local pseudopotentials. With this result we can now easily and efficiently calculate the NMR chemical shift for elements heavier then hydrogen using Eq. (1).

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