Selfconsistent order-$N$ density-functional calculations for very large systems

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(July 29, 2021)

We present a method to perform fully selfconsistent density-functional calculations, which scales linearly with the system size and which is well suited for very large systems. It uses strictly localized pseudoatomic orbitals as basis functions. The sparse Hamiltonian and overlap matrices are calculated with an $O(N)$ effort. The long range selfconsistent potential and its matrix elements are computed in a real-space grid. The other matrix elements are directly calculated and tabulated as a function of the interatomic distances. The computation of the total energy and atomic forces is also done in $O(N)$ operations using truncated, Wannier-like localized functions to describe the occupied states, and a band-energy functional which is iteratively minimized with no orthogonality constraints. We illustrate the method with several examples, including carbon and silicon supercells with up to 1000 Si atoms and supercells of $\beta$-$C_3N_4$. We apply the method to solve the existing controversy about the faceting of largeicosahedral fullerenes by performing dynamical simulations on $C_{60}$, $C_{240}$, and $C_{540}$.

A large effort has been devoted in the last few years to develop approximate methods to solve the electronic structure of large systems with a computational cost proportional to its size. Several approaches are now sufficiently accurate and robust to obtain reliable results for systems with thousands of atoms. So far, however, most of these schemes have been useful only with simple Hamiltonians, like empirical tight-binding models, which provide an ideal setting for order-$N$ calculations. First-principles order-$N$ calculations have been performed mainly in the non-selfconsistent Harris functional version of the local density approximation (LDA) for electronic exchange and correlation (XC) using minimal bases. Linear scaling algorithms in fully selfconsistent LDA have also been tried, but the results are far from the linear scaling regime, due to the relatively small number of manageable atoms in those simulations. Hernandez et al. have successfully produced LDA results in large silicon systems using a real-space grid method. The computational requirements that this kind of approach demands are, however, extremely large, and calculations must be performed in massive computational platforms.

We have developed a selfconsistent density-functional formulation with linear scaling, capable of producing results for very large systems, whose computational demands are not overwhelmingly large, so that systems with many hundreds of atoms can be treated in modest computational platforms like workstations, and much larger systems can be treated in massive platforms. The method is based on the linear combination of atomic orbitals (LCAO) approximation as basis of expansion of the electronic states. Non-orthogonal LCAO bases are very efficient, reducing the number of variables dramatically, compared to plane-wave (PW) or real-space-grid approaches, so that larger systems can be studied. Also, LCAO can provide up to extremely accurate bases, staying always in the range of a few valence orbitals per atom. As a first step, in this work we use minimal basis sets of one $s$ and three $p$ orbitals per atom, the extension to larger bases being perfectly possible within the present formulation. The choice of a basis obviously implies a possible error associated to its incompleteness. In the same way as for the error due to the linear scaling algorithm, the error in the basis can be reduced at the expense of an increase in computational effort. Its magnitude should be carefully checked, but also compared with other sources of error to ensure that an increase of the basis is really worthwhile.

Our method uses standard LDA techniques for the valence electrons, the core electrons being replaced by pseudopotentials. The basis orbitals used in this work are the $s$ and $p$ pseudoatomic orbitals defined by Sankey and Niklewski in the context of non-selfconsistent Harris functional methods. These are slightly excited pseudoatomic orbitals $\phi_{\mu}(r)$, obtained by solving the valence electron problem in the isolated atom, with the same pseudopotential and LDA approximations, and with the boundary condition that the atomic orbitals are strictly localized, vanishing outside a given radius $r_c$. This radius cutoff is in principle orbital dependent, but we do not make explicit this dependence in the equations only for simplicity in the notation. The great advantage of these orbitals is that they give rise to sparse overlap and Hamiltonian matrices (since matrix elements between distant orbitals exactly vanish exactly) and they display the same structure as in conventional tight-binding. The extent of the interactions and the sparseness of the matrices depend on the cutoff radius $r_c$ of each atom. These are not critical as long as the maxima of the atomic wave functions are well within $r_c$. For an analysis of the quality of pseudoatomic orbitals as a basis for solid state calculations we refer the reader to Ref. [10].
where $i$ is the spherical atomic charge density of the atom in the Kohn-Sham Hamiltonian by adding and subtracting techniques, adopting the most convenient one for each term of the Hamiltonian. In a prior step, to avoid dealing with the long range of the pseudopotentials, we rewrite the Kohn-Sham Hamiltonian by adding and subtracting the Hartree potential created by the neutral-atom charge $n_0(r)$, defined as

$$ n_0(r) = \sum_i n_{i}^{\text{NA}}(r - R_i) $$

where $i$ runs over the atoms in the system, and $n_i^{\text{NA}}$ is the spherical atomic charge density of the atom $i$ in its neutral, isolated state with $\rho_i^0$ electrons on each orbital $\phi_i$. If we define $\delta n(r) = n(r) - n_0(r)$ where $n(r)$ is the actual charge density, the Hartree potential can be decomposed into two contributions $V_{H}^0$ and $V_{\text{NL}}$, created by $\delta n(r)$ and $n_0(r)$ respectively. Using Eq. (1), $V_{H}^0$ can be expressed as a sum of atomic contributions. Also, the pseudopotential is decomposed into a short-range non-local term $V_{\text{NL}}$ and a long-range local term $V_L$. Following Sankey and Niklewski, we define the neutral atom potential of a given atom at $R_i$ as

$$ V_{\text{NA}}(r - R_i) = V_L(r - R_i) + e^2 \int \frac{n_i^{\text{NA}}(r - R_i)}{|r - r'|} dr'. $$

$V_{\text{NA}}$ is short ranged, since the core attraction and the electron Coulomb repulsion of the neutral atom charge cancel each other beyond $r_c$. The Kohn-Sham Hamiltonian is finally obtained as

$$ H^{\text{KS}} = \frac{p^2}{2m} + \sum_i [V_{\text{NL}}(r - R_i) + V_{\text{NA}}(r - R_i)] + V_{H}^0 + V_{XC}(r) $$

The overlap, kinetic energy term, neutral atom potential and non-local part of the pseudopotential, are all independent of the charge density $n(r)$, and their matrix elements between atomic orbitals can be expressed as sums of two center ($S_{\mu\nu} = \langle \phi_\mu | \phi_\nu \rangle$ and $\langle \phi_\mu | p^2 / 2m | \phi_\nu \rangle$) or three center ($\langle \phi_\mu | V_{\text{NL}}(r - R_i) | \phi_\nu \rangle$ and $\langle \phi_\mu | V_{\text{NA}}(r - R_i) | \phi_\nu \rangle$) integrals, which only depend on the relative positions of pairs or triplets of atoms. We follow the method proposed by Sankey and Niklewski to compute all these integrals: they are calculated beforehand and tabulated as a function of the relative position of the centers. These tables are used during the simulation, to calculate all the non-zero integrals by interpolation. The details of the procedure can be found in Ref. [12]. Since all these integrals are zero for distant enough atoms, their number scales linearly with the size of the system, as well as the computation time. The contributions of these terms to the Hamiltonian are computed only once for a given atomic configuration, since they do not depend on the selfconsistent charge.

The matrix elements of the Hartree potential $V_{H}^0(r)$ created by the charge density $\delta n(r)$ and the exchange-correlation potential $V_{XC}[n(r)]$ both depend on the selfconsistent charge. To calculate these integrals we compute $n_0(r)$, $n(r)$ and $\delta n(r)$ for a given LCAO density matrix at the points of a regular grid in real space. This is straightforward since the basis orbitals are defined in real space. Poisson’s equation for the Hartree potential can be then solved by the standard fast Fourier transform (FFT) method, assuming a supercell geometry, or by the multigrid method. In spite of its $N \log N$ scaling, we presently use FFT’s for simplicity, since this part represents a minor contribution to the total computational load. Note that only two FFT’s are necessary per cycle of selfconsistency (SCF cycle), in contrast with PW based calculations, where an FFT is required for each electronic state. The LDA XC potential is trivially computed on each point of the grid. Once the value of the Hartree and the XC potentials are known at every point, the integrals $\langle \phi_\mu | V_{H}^0 | \phi_\nu \rangle$ and $\langle \phi_\mu | V_{XC} | \phi_\nu \rangle$ are computed by direct summation on the grid. These sums are carefully done to minimize the amount of numerical workload involved. Only the non-zero integrals (between orbitals on atoms closer than 2$r_c$) are computed, and only the points of the grid for which both orbitals are non-zero contribute to each integral. We use sparse-matrix multiplication techniques optimized for this class of operations. As a result, the computation of the integrals scales linearly with the size of the system.

It is important to stress that the convergence with grid spacing of our method is different from that in standard PW calculations, which are known to require large PW cutoffs for systems containing atoms with hard pseudopotentials. In Figure 1 we show the convergence of the total energy per atom (referred to the converged value) for carbon, as a function of $E_{\text{cut}}$, the kinetic energy cutoff of the plane waves that the grid can represent. Full circles are for a diamond supercell of 64 atoms, whereas diamonds are for a cluster of 3 carbon atoms in a supercell of $15 \times 15 \times 15 \, \text{Å}^3$. In both cases, the results are converged.
to below 2 meV/atom for a cutoff of 30 Ry. This is in sharp contrast with results of PW calculations\(2^{3}\) (open circles) in which the cutoff necessary to achieve convergence (with the same pseudopotential) is much higher. Note, moreover, that the energy cutoff in our case refers to the grid representation of the charge density, whereas in the PW case it refers to the wave functions, which implies an even higher (four times) cutoff in the charge density. The reason for the fast convergence of our approach is that most of the Hamiltonian terms (most importantly the kinetic energy and the neutral atom potential) are not computed in the grid.

Once the Kohn-Sham Hamiltonian has been obtained, we use a recently proposed order-N method\(4^{13}\) to compute the band structure energy \(E_{BS}\) (sum of occupied eigenvalues). In this approach, a modified band energy functional\(4^{14}\) is minimized\(4^{14}\), with respect to the electronic orbitals by means of a conjugate gradients (CG) algorithm, to yield \(E_{BS}\). The orthonormality of the occupied states does not need to be imposed, but it is obtained as a result of the minimization of the energy functional. The elimination of the orthogonalization is the first step to achieve an order-N scaling. The second is the use of localized, Wannier-like wave functions (LWF) to describe the electronic states entering the minimization of the energy functional. Truncation of these localized functions beyond a given cutoff \(R_c\) from the center of the LWF provides a linear scaling algorithm. The errors involved in this truncation, which can be reduced arbitrarily by increasing the value of \(R_c\), are analyzed in detail in Ref.\(4^{14}\).

After the band energy has been minimized and the LWF’s obtained, the new charge density is computed, completing a so-called SCF cycle. From the density, a new Hamiltonian is produced, the procedure being repeated until selfconsistency in the charge density or the Hamiltonian is achieved. At this point, the total energy can be computed as

\[
E_{tot} = E_{BS} - \frac{e^2}{2} \int V_H(r)n(r)dr + \frac{e^2}{2} \int V_0(r)n_0(r)dr + \int [\epsilon_{XC}(n) - V_{XC}(n)] n(r)dr + U_{i\alpha - ee} \tag{4}
\]

where \(V_H(r)\) is the Hartree potential of the selfconsistent charge \(n(r)\), and, following Sankey and Niklewski,\(4^{14}\) we have defined

\[
U_{i\alpha - ee} = \frac{e^2}{2} \sum_{i\alpha} Z_i Z_{\alpha} |\mathbf{R}_i - \mathbf{R}_{\alpha}| - \frac{e^2}{2} \int V_0(r)n_0(r) \tag{5}
\]

As in the case of the Hamiltonian, we have added and subtracted the electrostatic energy of the neutral atom charge \(n_0(r)\) to obtain Eq.\(4^{14}\). The advantage, again, is that \(U_{i\alpha - ee}\) can be expressed as a sum of short range contributions, which is easy to evaluate in \(O(N)\) operations,\(4^{14}\) avoiding the problems related with the long range character of the ionic core interactions. The integrals appearing in Eq.\(4^{14}\) are calculated in the real space grid.

In molecular dynamics (MD) simulations and geometrical optimizations the atomic forces are needed. We compute them using a variation of the Hellman-Feynman theorem, which includes Pulay-like corrections to account for the fact that the basis set is not complete and moves with the atoms. The force on atom \(i\) is

\[
F_i = - \sum_{\mu\nu} \rho_{\mu\nu} \frac{\partial H^0}{\partial R_i} + \sum_{\mu\nu} E_{\mu\nu} \frac{\partial S_{\mu\nu}}{\partial R_i} - \frac{\partial U_{i\alpha - ee}}{\partial R_i} + 2 \sum_{\mu} \rho_{\mu} \frac{\partial \phi_{\mu}}{\partial R_i} [V_0^\mu |\phi_{\mu}|^2 - 2 \sum_{\mu\nu} \rho_{\mu\nu} \frac{\partial \phi_{\mu}}{\partial R_i} [V_0^\mu |\phi_{\mu}| - V_{XC} |\phi_{\mu}|] \tag{6}
\]

where \(H^0 = p^2/2m + V_{NL} + V_{NA}\), and \(\rho_{\mu\nu}\) and \(E_{\mu\nu}\) are the density and energy-density matrices, respectively.\(4^{14}\) The first three terms are calculated interpolating the tabulated data,\(4^{14}\) whereas the last two terms are computed by numerical integration in the grid, as was done for the matrix elements of the Hartree and XC potentials in the Kohn-Sham Hamiltonian.

In order to show the linear scaling of the method, we have performed calculations on supercells of silicon in the diamond structure, with different numbers of atoms from 64 to 1000. Only the \(\Gamma\) point was used to sample the Brillouin zone, the cutoff for the charge density grid was 12 Ry, and the LWF’s were truncated at 4.5 Å. Figure\(8\) shows the linear behavior of the CPU time and memory requirements with the number of atoms. The CPU time shown represents the average cost to perform a SCF step in a MD simulation, including the calculation of the charge density and Hamiltonian matrix elements, the minimization of the band structure energy, and the calculation of the atomic forces. The band structure energy minimization within each SCF cycle required an average of 20 CG iterations, while the number of SCF cycles depends largely on the simulation temperature, length of the time step and mixing algorithm for selfconsistency. So far, in comparable simulation conditions, no significant dependence of the number of CG iterations and SCF cycles with the size of the system has been observed. As we can see, in the present method both the CPU time and memory requirements are small enough to permit the calculation of a system of 1000 silicon atoms in a very modest workstation.
In conclusion, we have presented an efficient method for selfconsistent LDA calculations with linear scaling. We have analyzed the performance versus system size and grid cutoff, and shown that simulations of systems with hundreds of atoms are possible with small workstations. This should open the possibility of very large scale \textit{ab initio} simulations in the near future.

We acknowledge R. M. Martin and Paul von Allmen for many useful discussions, and D. A. Drabold and O. F. Sankey for allowing us the use of many of their codes. P. O. is indebted to R. M. Martin and J. B. Adams for continuous support and encouragement. This work was partially supported by DOE Grant No. DEFG 02-91ER45439 and DGICYT (Spain) Grant No. PB92-0169.

As an example of a system with partially ionic character, and with atoms with compact orbitals, we have performed calculations on the \(\beta\) phase of \(\text{C}_3\text{N}_4\), which was proposed as a potentially very hard material by Liu and Cohen.\cite{Liu} The calculations were done in supercells of 42 and 224 atoms, with nearly identical results. A cutoff of 200 Ry for the charge density grid was used. We obtain an accuracy better than 1\% in both the lattice constants and the several inequivalent bond lengths, and 10\% in the bulk modulus, compared to other LDA calculations.\cite{Poirier}

As a system with compact orbitals was studied, we have compared the results of the present work with those of Itoh \textit{et al.}\cite{Itoh} obtained with the Harris functional.

| This work | Itoh \textit{et al.}\cite{Itoh} |
|-----------|-------------------------------|
| \(\bar{r}\) (Å) | \(\bar{r}_\sigma\) | \(\bar{r}_{\sigma m}\) | \(\phi\) | \(\bar{r}_\sigma\) | \(\bar{r}_{\sigma m}\) | \(\phi\) |
| \(\text{C}_{60}\) | 3.59 | 0.000 | 0.000 | 12.0\% | 3.55 | 0.000 | 0.000 | 12.0\% |
| \(\text{C}_{240}\) | 7.18 | 0.023 | 0.027 | 8.5\% | 7.06 | 0.021 | 0.028 | 7.9\% |
| \(\text{C}_{540}\) | 10.69 | 0.038 | 0.054 | 9.6\% | 10.53 | 0.033 | 0.053 | 9.2\% |

\begin{table}[h]
\centering
\begin{tabular}{lcccccc}
\hline
 & \(r\) (Å) & \(\sigma_r\) & \(\sigma_{\sigma m}\) & \(\phi\) & \(r\) (Å) & \(\sigma_r\) & \(\sigma_{\sigma m}\) & \(\phi\) \\
\hline
\(\text{C}_{60}\) & 3.59 & 0.000 & 0.000 & 12.0\% & 3.55 & 0.000 & 0.000 & 12.0\% \\
\(\text{C}_{240}\) & 7.18 & 0.023 & 0.027 & 8.5\% & 7.06 & 0.021 & 0.028 & 7.9\% \\
\(\text{C}_{540}\) & 10.69 & 0.038 & 0.054 & 9.6\% & 10.53 & 0.033 & 0.053 & 9.2\% \\
\hline
\end{tabular}
\caption{Average radius (\(\bar{r}\)), standard (\(\sigma\)) and maximum deviation \(\sigma_m = (r_{\text{max}} - r_{\text{min}})/2\) of radii, and non-planarity angle \(\phi\) around pentagons (going from 0\% for a planar pentagonal site to 12\% for a truncated icosahedron) for the fullerene clusters. We compare the results of the present work with those of Itoh \textit{et al.}\cite{Itoh} obtained with the Harris functional.}
\end{table}

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