Linear Scaling Solution of the Coulomb problem using wavelets

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The Coulomb problem for continuous charge distributions is a central problem in physics. Powerful methods, that scale linearly with system size and that allow us to use different resolutions in different regions of space are therefore highly desirable. Using wavelet based Multi Resolution Analysis we derive for the first time a method which has these properties. The power and accuracy of the method is illustrated by applying it to the calculation of of the electrostatic potential of a full three-dimensional all-electron Uranium dimer.

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The theory of wavelets is one of the most important recent developments in mathematics. It allows one to apply a multi-scale analysis to problems that exhibit widely varying length scales. Problems with this feature abound in all fields of physics. The problem we want to address here is the classical Coulomb problem for a continuous charge distribution \( \rho \), i.e. we want to solve Poisson’s equation

\[
\nabla^2 V = -4\pi \rho
\]

under the constraint that the potential \( V \) vanishes at infinity. This basic equation can be found in nearly any field of physics and it is therefore essential to have efficient solution methods for it. There are two important requirements for an algorithm that solves this problem. First it should scale linearly with the size of the charge distribution. Since in numerical applications the charge distribution is given on a grid a measure for the size of the charge distribution is the number of grid points necessary to represent it. This linear scaling property is of utmost importance because in many applications one needs grids consisting of a very large number of grid points. Second, it should allow for grids that are nonuniform, i.e. that have higher resolution in regions where this is required.

For discrete charge distributions several algorithms have been put forward to map the continuous problem onto a discrete and use the above mentioned algorithms for discrete systems. For continuous charge distributions proposals have been put forward to map the continuous problem onto a discrete and use the above mentioned algorithms for discrete systems. To the best of our knowledge, there exists however no linear scaling algorithm for nonuniform grids that can directly be applied to continuous charge distributions. If one constrains oneself to uniform grids and periodic boundary conditions, there are of course the well known Fourier techniques that show a nearly linear \( N \log_2(N) \) scaling with respect to the number of grid points \( N \). Non-periodic boundary conditions can be implemented in the context of Fourier techniques only by cutting off the long range Coulomb potential. Finite element methods allow nonuniform grids, but grid generation and preconditioning pose severe problems. Using a basis of wavelet functions, we will present in this paper a method that scales strictly linear and allows for nonuniform grids.

There are many families of wavelets and one has to choose the most appropriate one for a specific application. A widely used family are the compactly supported orthogonal wavelets of Daubechies. The orthogonality property is convenient if one has to expand an arbitrary function in a basis of wavelets. Their disadvantage is that they are not very smooth, i.e. only a small number of derivatives is continuous. One can however construct families of nonorthogonal wavelets that are much smoother. In general the mapping from the numerical values on a grid to the expansion coefficients of the wavelet basis is rather complicated and slow for biorthogonal wavelets. An exception are the second generation interpolating wavelets, whose special properties allow us to do this mapping easily. In addition they give rise to a particularly fast wavelet transform.

Wavelets have already been successfully applied in several areas of physics. In the context of electronic structure calculations a fairly small wavelet basis can describe the widely varying scales of both core and valence electrons. Self-consistent electronic structure calculations have also been done. In these self-consistent calculations the solution of Poisson’s equation was however done by traditional Fourier techniques.

Let us now briefly review the theory behind biorthogonal wavelets. As in ordinary orthogonal wavelet theory there are two fundamental functions, the scaling function \( \phi \) and the wavelet \( \psi \). In the biorthogonal case there are however still the complementary scaling function \( \tilde{\phi} \) and the complementary wavelet \( \tilde{\psi} \). Each scaling function and wavelet belongs to a hierarchical level of resolution. By analysing a function with respect to these different levels of resolution one can do a so-called Multi Resolution Analysis (MRA). The space belonging to a certain level of resolution \( k \) is spanned by all the integer translations
of the scaling function \( \phi_{i,k}(x) \propto \phi((\frac{1}{2})^k x - i) \). Any function can be expanded within this level of resolution.

\[
f(x) \approx \sum_i s_{i,k} \phi_{i,k}(x) ,
\]

Since the scaling functions and their complementary counterparts are orthogonal \(< \phi_{k,i}(x) | \phi_{k,j}(x) > = \delta_{ij} \), the expansion coefficients \( s_{i,k} \) are given by

\[
s_{i,k} = \langle f(x) | \phi_{i,k}(x) \rangle .
\]

The expansion becomes more accurate if one goes to a higher level of resolution, i.e. if one decreases \( k \) and it becomes exact in the limit \( k \to -\infty \). This is an important feature because it allows us to improve systematically the numerical accuracy in very much the same way as it is done with a basis of plane waves. In numerical application it is of course not possible to take this limit, and we will therefore denote the finest level of resolution that is used in the calculation by \( k = 0 \).

The scaling function satisfies a refinement relation

\[
\phi_{j,k}(x) = \sum_l \hat{h}_{l-2j} \phi_{l,k-1}(x)
\]
i.e. each scaling function of a lower resolution level can be expressed as a linear combination of higher resolution scaling functions. It is obviously not possible to express a scaling function of higher resolution by a linear combination of lower resolution scaling functions only. One can, however, write down such an expression if one still includes the wavelets \( \psi_{i,k}(x) \propto \psi((\frac{1}{2})^k x - i) \)

\[
\phi_{j,k-1}(x) = \sum_l h_{l-2j} \phi_{l,k}(x) + \sum_l g_{l-2j} \psi_{l,k}(x)
\]

The wavelets at level \( k \) thus reintroduces the resolution that is lost as one goes from level \( k \) to level \( k + 1 \) scaling functions. These transformation properties among the scaling functions and wavelets give rise to the wavelet transform. The wavelet expansion coefficients \( d_{i,k} \) are then either defined by this transform or equivalently by an expression analogous to Eq. 3.

\[
s_{i,k+1} = \sum_j \hat{h}_{l-2j} s_{j,k} ,
\]

\[
d_{i,k+1} = \sum_j g_{l-2j} s_{j,k}
\]

\[
s_{i,k} = \sum_j \left( \hat{h}_{l-2j} s_{j,k+1} + \hat{g}_{l-2j} d_{i,k+1} \right)
\]

Eq. 6 is called a forward fast wavelet transform, Eq. 7 being its inverse counterpart. If one has periodic boundary conditions for the data \( s_{i,k} \) the wavelet transform is one-to-one transformation between \( s_{i,k} \) and its spectral decompositions \( s_{i,k+1} \) and \( d_{i,k+1} \). To obtain a full wavelet spectral analysis the forward transform is applied recursively. In consecutive steps the output data \( s_{i,k+1} \) of the previous forward transform are the input data for the next transform. The size of the data set to be transformed is thus cut into half in each step. The total operation count is then given by a geometric series and scales therefore strictly linear. A full wavelet synthesis consists in the same way of a sequence of inverse transforms and gives back the original data set \( s_{i,k} \). The coefficients \( h_i \) and \( g_i \) and their complementary counterparts \( \hat{h}_i, \hat{g}_i \) are filters of finite length \( 2m \) and can be derived from the MRA requirements. The 8-th order lifted Lazy scaling function and wavelet that were used in this work are shown in Fig.1. Because it can represent polynomials up to degree 8 exactly, the expansion coefficients with respect to the wavelets \( d_{i,k} \) decay very rapidly for any smooth function with decreasing \( k \).

![FIG. 1. The scaling function (full line) and wavelet (dashed line) used in this work](image)

To do a multidimensional MRA, we use a scheme described by Daubechies. Even though all this work was done in the three-dimensional case, we will illustrate the principle just for the two-dimensional case. The space of all scaling functions of resolution level \( k \) is given by

\[
\phi_{i,j,k}(x,y) = \phi_{i,k}(x) \phi_{j,k}(y)
\]
The wavelet space is again defined as the space that rec-ompensates for the resolution that is lost by going up one level in the scaling functions space. Using Eq. 8 one obtains three kind of terms for the wavelet space

\[
\psi_{i,j,k}^{01}(x,y) = \phi_{i,k}(x) \psi_{j,k}(y) \]
\[
\psi_{i,j,k}^{10}(x,y) = \psi_{i,k}(x) \phi_{j,k}(y) \]
\[
\psi_{i,j,k}^{11}(x,y) = \psi_{i,k}(x) \psi_{j,k}(y)
\]

Let us now explain how to solve Poisson’s equation in wavelets. Expanding both the charge density and the potential in Eq. 1 into scaling functions

\[
\rho(x,y,z) = \sum_{i,j} \rho_{i,j} \phi_{i,k}(x) \phi_{j,k}(y) \]
\[
V(x,y,z) = \sum_{i,j} V_{i,j} \phi_{i,k}(x) \phi_{j,k}(y)
\]
one obtains the following system of equations.

\[
\sum_{j1,j2} L_{i,j1,j2} V_{j1,j2} = \rho_{i,j}
\]
where

\[ L^k_{i,j;\mu,\nu} = \langle \tilde{\phi}_{i,k}(x) \tilde{\phi}_{j,k}(y) | \nabla^2 | \phi_{\mu,k}(x) \phi_{\nu,k}(y) \rangle \quad (15) \]

Since the scaling functions have a finite support, the matrix \( L^k \) is a sparse matrix and its nonzero elements \( L^k_{1,i;2,j;1,2} \) can be calculated analytically. The natural boundary conditions for this scheme are periodic boundary conditions. As we stressed in the introduction, we however want to solve Poisson's equation with non-periodic boundary conditions. As is well known, boundary affects vanish whenever the boundary is sufficiently far away. Thus, one could in principle obtain natural boundary conditions (i.e. \( V(r) \to 0 \) if \( r \to \infty \)) within arbitrary precision if one uses a sufficiently large periodic box. Since the electrostatic potential decays fairly slowly a very large box is required and the numerical effort would be tremendous if one uses equally spaced grids within this huge periodic computational box. Far away from the charge distribution the variation of the potential is however small and less resolution is needed. The key idea is therefore to use a set of hierarchical grids as shown in Fig.2, where the resolution decreases as one goes out of the center. Expressed in the terms of wavelet theory this means that on the highest (periodic) level we have a basis of scaling functions. Resolution is then increased by adding wavelet basis functions near the center. By doing this repeatedly we obtain increasing resolution towards the center as shown in Fig.2.

Up to now the motivation for introducing grids of different resolution was to handle the natural boundary conditions. Additional levels of resolution can however be introduced to handle charge distributions that have different length scales and require therefore higher resolution in some parts of space. The theory of wavelets gives us also enough flexibility to increase the resolution not only around one center but around any number of centers in the computational box.

![Level 0](image1)

FIG. 2. A hierarchical multi resolution grid of the type used in this work. For simplicity only three levels of resolution are shown.

A full wavelet synthesis step can be done straightforwardly in this hierarchical grid setting. Any wavelet can be decomposed into scaling functions and therefore one can calculate the scaling function coefficients at any level of resolution and for any point in the computational volume. If one calculates these scaling functions for high resolution levels in a region of low resolution, one obtains however a highly redundant data set. To do a full wavelet analysis that brings back the original spectral decomposition data, it turns out that one needs actually a slightly redundant data set. In order to calculate the wavelet coefficients for a wavelet at a boundary to a lower resolution region, one needs the scaling function values corresponding to this higher resolution also in a strip of width \( m \) in the lower resolution region. A schematic diagram of a full hierarchical wavelet analysis and synthesis is shown in Fig.3.

![Level 1](image2)

FIG. 3. A schematic representation of a multi hierarchy wavelet synthesis. Data regions denoted by \( s,d^0, d^1 \) and \( d^2 \) contain expansion coefficients for basis functions of the type given by Eq. 8, 9, 10 and 11 respectively. CP stand for a copy step where one puts an additional layer of zeroes around the data set. SYN denotes a one level wavelet synthesis step. One starts the process at the coarsest (periodic) level and proceeds down to the finest resolution level. To do a multi hierarchy wavelet analysis one proceeds back up reversing all the copy operations and replacing the single level synthesis steps by analysis steps.

In this mixed representation, where one has scaling functions at the highest periodic level and wavelets all the refinement levels, the structure of the Laplace operator is much more complicated since one has coupling between all the hierarchical levels. An elegant way to cope with this additional complexity is the so-called non-standard operator form proposed by Beylkin, Coifman and Rokhlin which allows us to incorporate this coupling by a sequence of wavelet transforms (Fig.3), that are interleaved with the application of a simple one-level Laplace operator. For this one-level Laplace operator only the matrix elements of the Laplace operator among scaling functions and wavelets on the same resolution level, but not between different levels of resolution are needed.
Mathematically the nonstandard operator form is a telescopic expansion of the Laplace operator in the scaling function basis at the finest level $L^0$. If we define projection operators $P_k$ and $Q_k$, that project the whole space into the space of scaling functions and wavelets at the $k$-th level as well as their complementary counterparts $\tilde{P}_k$ and $\tilde{Q}_k$, they satisfy

$$P_k = P_{k+1} + Q_{k+1} \quad ; \quad \tilde{P}_k = \tilde{P}_{k+1} + \tilde{Q}_{k+1}$$

and we may write

$$L^k = \tilde{P}_k L^0 P_k = (\tilde{Q}_{k+1} + \tilde{P}_{k+1}) L^0 (Q_{k+1} + P_{k+1}) = L_{DD}^{k+1} + L_{SD}^{k+1} + L_{DS}^{k+1} + L_{SS}^{k+1}$$

where $L_{DD}^{k+1}$, $L_{SD}^{k+1}$, $L_{DS}^{k+1}$ are Laplace operators at the $(k + 1)$th level representing the coupling of wavelets with wavelets, wavelets with scaling functions and scaling functions with wavelets. Applying Eq. 17 recursively for $k = 0, 1, \ldots$, one obtains the nonstandard operator form.

In the basis of the wavelet functions at different resolution levels a simple diagonal preconditioning scheme is very efficient and we were able to reduce the residue by one order of magnitude with only 3 iterations.

To demonstrate the power of this method we applied it to a problem that can hardly be solved by any other methods, namely the potential arising from the nucleonic and electronic charge distribution of a fully three-dimensional all-electron Uranium dimer. The charge distribution of the nucleus was represented by a Gaussian charge distribution with an extension of $1 \text{ a.u.}$. Since the valence electrons have an extension which is of the order of one atomic unit, we have length scales that differ by more than 3 orders of magnitude. As can be seen from Fig.4, the potential also varies by many orders of magnitude. Using 22 hierarchical levels in our algorithm, we can represent resolutions that differ by 7 orders of magnitude and we are able to calculate the potential with at least 6 significant digits in the whole region from the nucleus to the valence region. In order to be able to determine the error we actually first fitted the electronic charge distribution by a small number of Gaussians, whose exact potential can be calculated analytically. This rather crude charge density was then used in all the calculations.

We also applied the method to clusters containing several $CO$ molecules that were described by pseudo-potentials. In this case there is only one length scale associated with the charge distribution and it is possible to reduce the number of grid points on higher levels such that the total amount of work increases only slightly with additional hierarchies. We were able to calculate the potential corresponding to the non-periodic boundary conditions with 8 significant digits.

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