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We introduce a new version of the Fast Multipole Method for the evaluation of potential fields in three dimensions. It is based on a new diagonal form for translation operators and yields high accuracy at a reasonable cost.

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1 Introduction

In this paper, we introduce a new version of the Fast Multipole Method (FMM) for the evaluation of potential fields in three dimensions. The scheme evaluates all pairwise interactions in large ensembles of particles, i.e. expressions of the form

\[ \Phi(x_j) = \sum_{i=1, \ i \neq j}^{n} \frac{q_i}{\|x_j - x_i\|} \]  

for the gravitational or electrostatic potential and

\[ E(x_j) = \sum_{i=1, \ i \neq j}^{n} q_i \cdot \frac{x_j - x_i}{\|x_j - x_i\|^3} \]  

for the field, where \( x_1, x_2, \ldots, x_n \) are points in \( \mathbb{R}^3 \), and \( q_1, q_2, \ldots, q_n \) are a set of (real) coefficients.

The evaluation of expressions of the form (1) is closely related to a number of important problems in applied mathematics, physics, chemistry, and biology. Molecular dynamics and Hartree-Fock calculations in chemistry, the evolution of large-scale gravitational systems in astrophysics, capacitance extraction in electrical engineering, and vortex methods in fluid dynamics are all examples of areas where simulations require rapid and accurate evaluation of sums of the form (1) and (2). When certain closely related interactions are considered as well, involving expressions of the form

\[ \Phi(x_j) = \sum_{i=1, \ i \neq j}^{n} q_i \cdot e^{ik\|x_j - x_i\|} \]  

the list of applications becomes even more extensive.

This paper is a continuation (after an interval of several years) of a sequence of joint papers by the authors, starting with (Greengard and Rokhlin 1987) and (Carrier et al. 1988) which introduced the Fast Multipole Method in two dimensions. Subsequent work extended the method to three dimensions (Greengard 1988; Greengard and Rokhlin 1988a,b), and there followed a number of versions of the scheme, both by the present authors and by other researchers (see, for example, Anderson 1992; Nabors et al. 1994; Berman 1995; Epton and Dembart 1995; Elliott and Board 1996). After about ten years of research, however, a somewhat unsatisfactory picture has emerged. In short, there now exist extremely efficient algorithms for the evaluation of the two-dimensional analogues of (1), (2) with (practically) arbitrarily high precision, as well as very efficient and accurate algorithms for a host of related problems (Rokhlin 1988; Alpert and Rokhlin 1991; Beylkin et al. 1991; Coifman and Meyer 1991; Greengard and Strain 1991; Strain 1991; Alpert et al. 1993). However, for the sums (1) and (2), there are few practical schemes, and these provide only limited accuracy. Since most real-world problems are three-dimensional, it can be said that analysis-based "fast" methods are a promising group of techniques, but that they have not yet lived up to all their expectations.

In the present paper, we try to remedy this situation. We describe a version of the Fast Multipole Method in three dimensions that produces high accuracy at an acceptable computational
cost. As will be seen from the numerical examples in Section 9, the new scheme has a break-even point of $n \sim 2000$ when compared with direct calculation in single precision; with 10-digit accuracy, the break-even point is $n \sim 5000$; with 3-digit accuracy, it is $n \sim 500$. The approach uses a considerably more involved mathematical (and numerical) apparatus than is customary in the design of fast multipole-type algorithms. This apparatus is based on a new diagonal form for translation operators acting on harmonic functions, extending the two-dimensional version introduced in (Hrycak and Rokhlin 1995). The overall approach bears some resemblance to that used in fast multipole methods for high frequency scattering problems, which are based on diagonal forms of translation operators for the Helmholtz equation (Rokhlin 1990, 1995; Epton and Dembart 1995).

2 Philosophical Preliminaries

We begin with an overview of analysis based "fast" numerical algorithms, concentrating on the evaluation of expressions of the form (1). Where possible, we summarize the current "state of the art" in the field.

If we define the $n \times n$-matrix $A$ by the formula

$$A_{ij} = \frac{1}{\|x_j - x_i\|},$$

we can rewrite (1) in the form

$$\Phi = Aq,$$

with $\Phi, q \in \mathbb{R}^n$ (the expression (2) can be rewritten in a similar fashion). Obviously, straightforward evaluation of either of the expressions (1), (2) requires $O(n^2)$ operations (evaluating $n$ potentials at $n$ points), and for large-scale problems this estimate is prohibitively large. On the other hand, the evaluation of expressions of the forms (1), (2) is an integral part of the numerical solution of many important problems in applied mathematics, and during the last decade, several "fast" schemes have been proposed for this purpose, i.e. schemes whose computational cost is less than $O(n^2)$. Typically, such methods require $O(n)$ or $O(n \cdot \log n)$ operations (Rokhlin 1985; Greengard and Rokhlin 1987; Carrier et al. 1988; Rokhlin 1988, 1990; Alpert and Rokhlin 1991; Beylkin et al. 1991; Coifman and Meyer 1991; Greengard and Strain 1991; Strain 1991, 1992; Epton and Dembart 1995). All of them are based on the straightforward observation that the potentials are smooth functions in $\mathbb{R}^3$, except when $x_i$ is near $x_j$, and as a result, large submatrices of $A$ are of low rank (to any finite precision). Clearly, applying a matrix of dimension $n \times n$ but rank $J$ to an arbitrary vector requires only $n \cdot J$ operations (as opposed to $n^2$); this simple observation leads directly to a variety of asymptotically "fast" schemes for the evaluation of (1); below, we illustrate the construction of such schemes with a simple example.

Suppose that, in the expression (1), the points $x_1, x_2, \ldots, x_n$ are equispaced and lie on the interval $[-1, 1]$, so that

$$x_1 = -1, \ x_2 = -1 + h, \ \ldots, \ x_{n-1} = 1 - h, \ x_n = 1,$$
Figure 1: Subdivision of matrix into well-separated blocks. The submatrices marked by an X are not well-separated from the diagonal.

where $h = 2/(n - 1)$. Given three integers $l$, $m$, $k$ such that

$$1 \leq l \leq n,$$
$$1 \leq m \leq n,$$
$$1 \leq k \leq n - l,$$
$$1 \leq k \leq n - m,$$

we will denote by $A_{l,m,k}$ the submatrix of $A$ consisting of such elements $A_{ij}$ that

$$l \leq i \leq l + k - 1,$$
$$m \leq j \leq m + k - 1,$$

and say that $A_{l,m,k}$ is separated from the diagonal if

$$| l - (m + k - 1) | > k,$$

and

$$| m - (l + k - 1) | > k.$$

In other words, we will say that the submatrix $A_{l,m,k}$ of the matrix $A$ is separated from the diagonal if its distance from the diagonal of $A$ is greater than or equal to its own size (Fig. 1). We will construct a rudimentary "fast" algorithm for the application of the matrix $A$ to an arbitrary vector by means of the following lemma; its proof is based on several well-known facts, all of which can be found in (Dahlquist and Bjork 1974).
Lemma 2.1 For any integer \( p \leq 1 \), and any \( l, m, k \) satisfying the conditions (7), there exists a matrix \( B_{i,m,k} \) of dimension \( k \times k \) and rank \( J \), such that

\[
\|A_{i,m,k} - B_{i,m,k}\| \leq \frac{1}{4^J}.
\]

In other words, any submatrix of \( A \) separated from the diagonal is of rank \( J \), to the precision \( 1/4^J \).

Outline of proof.

We start by defining the function \( f : \mathbb{R}^2 \rightarrow \mathbb{R}^1 \) by the formula

\[
f(x, y) = \frac{1}{|x-y|},
\]

and observing that \( f \) is smooth everywhere in \( \mathbb{R}^2 \), except when \( x = y \). We will say that the square \([a, a+c] \times [b, b+c] \subset \mathbb{R}^2\) is separated from the diagonal if

\[
|a + c - b| > c,
\]

and

\[
|b + c - a| > c,
\]

and observe that on any such square, the function \( f \) can be expanded in a two-dimensional Chebychev series, i.e. represented in the form

\[
f(x, y) = \sum_{p,q=0}^{\infty} \alpha_{pq} \cdot T_p\left(\frac{2 \cdot x - c + 2 \cdot a}{c}\right) \cdot T_q\left(\frac{2 \cdot y - c + 2 \cdot b}{c}\right),
\]

with \( T_j \) denoting the \( j \)-th Chebychev polynomial. Finally, we observe that for any \( a, b, c \) satisfying the conditions (13), (14), the convergence of the expansion (15) is given by the formula

\[
|f(x, y) - \sum_{p,q=0}^{J} \alpha_{pq} \cdot T_p\left(\frac{2 \cdot x - c + 2 \cdot a}{c}\right) \cdot T_q\left(\frac{2 \cdot y - c + 2 \cdot b}{c}\right)| < \frac{1}{4^J}.
\]

In other words, for any square separated from the diagonal, the expansion (15) converges to accuracy \( \varepsilon \) after no more than \( \log_4(\varepsilon) \) terms. Combining (9), (10) and (1) with (15) and (16), we observe that for any \( i, j \) satisfying the inequalities (8),

\[
|A_{ij} - \sum_{p,q=0}^{J} \alpha_{pq} \cdot T_p\left(\frac{2 \cdot x_i - c + 2 \cdot a}{c}\right) \cdot T_q\left(\frac{2 \cdot y_j - c + 2 \cdot b}{c}\right)| < \frac{1}{4^J},
\]

with \( a = (2 \cdot l)/n - 1 \), \( b = (2 \cdot m)/n - 1 \), \( c = (2 \cdot k)/n \). The matrix \( B_{i,m,k} \) defined by

\[
(B_{i,m,k})_{ij} = \sum_{p,q=0}^{J} \alpha_{pq} \cdot T_p\left(\frac{2 \cdot x_i - c + 2 \cdot a}{c}\right) \cdot T_q\left(\frac{2 \cdot y_j - c + 2 \cdot b}{c}\right)
\]
clearly satisfies the desired condition (11).

In order to develop a fast algorithm, we first subdivide the matrix $A$ into a collection of submatrices, as depicted in Fig. 1. Each of the submatrices in this structure is separated from the diagonal, except the submatrices near the diagonal whose ranks are small simply because their dimensionality is small. By virtue of Lemma 2.1, each of the separated submatrices is of rank $\ll J$, to the accuracy $4^{-j}$. In order to apply $A$ to an arbitrary vector with fixed but finite accuracy (which is always the case in numerical computations), we can apply each of the submatrices to the appropriate part of the vector for a cost proportional to $k \cdot J$, where $k$ is the size of the submatrix. Adding up the costs for all such submatrices, we obtain the operation count of

$$J \cdot n \cdot \log(n) \sim \log\left(\frac{1}{\varepsilon}\right) \cdot n \cdot \log(n),$$

instead of $n^2$.

The scheme outlined above is extremely simple, but representative of the current approach to the design of “fast” summation algorithms. Several comments are in order.

1. It is easy to see that the matrix $A$ defined in (4) with the spacing defined by (6) is in fact a Toeplitz matrix that can be applied to an arbitrary vector for a cost proportional to $n \cdot \log(n)$ via the Fast Fourier Transform. This situation occurs sometimes, both in one and higher dimensions. However, the Toeplitz nature of the matrix $A$ is lost when the points are not distributed on a uniform grid, and direct application of the FFT becomes impossible. For “somewhat uniformly” distributed points $x_i$, various types of local corrections have been successfully utilized. When the points are not distributed uniformly (for example, on a curve or surface), FFT-based methods become ineffective.

2. As described, the scheme is only applicable to one-dimensional problems, and under very limited conditions. In most situations, the subdivision of the matrix has to be modified, taking into account the geometric distribution of points in order to locate submatrices whose “numerical rank” is low. Examples of such subdivisions can be found in (Carrier et al. 1988; Van Dommelen and Rundensteiner 1989; Beylkin et al. 1991; Nabors et al. 1994)

3. The scheme is extremely simple and general. It is entirely unrelated to the detailed nature of the matrix $A$, needing only some inequality like (16). In other words, so long as the entries of the matrix $A$ are smooth functions of their indices away from the diagonal, a scheme of the type outline above will work. In fact, even that is not necessary; the elements of the matrix have only to be sufficiently smooth functions of their indices on a sufficiently large part of the matrix.

4. The scheme admits a large number of modifications; the most obvious ones replace the Chebychev expansion in (15) with other approximations; one should be careful in doing so, since under many conditions the Chebychev approximation is optimal (among polynomial approximations), or nearly so. Some of the special-purpose approximation schemes which have been used successfully employ wavelets and related bases (Beylkin et al. 1991; Alpert et al. 1993).

Another obvious modification is a change in the choice of submatrices of low rank; the use of
rectangular submatrices (as opposed to the square ones in Fig. 1) permits coarser subdivisions and tends to result in more efficient algorithms.

5. Algorithms of the type described above usually do not work for problems where the matrix $A$ is a discretization of an integral operator with an oscillatory kernel, since such discretizations (normally) have a more or less constant number of nodes per wavelength of the dominant oscillation. As a result, the rank of each submatrix is proportional to its size, and the resulting algorithms have CPU time estimates of the order $O(n^2)$. Sometimes, the calculation can be accelerated by reducing the size of the constant (Wagner et al. 1994), but the asymptotic complexity in such cases is the same as for the direct approach. For certain classes of oscillatory problems (such as Helmholtz and Schrödinger equations at high frequency), there exist asymptotically “fast” schemes based on a different (and considerably more involved) analytical apparatus (see, for example, Rokhlin 1988, 1990, 1993; Canning 1989, 1992, 1993; Coifman and Meyer 1991; Bradie et al. 1993; Coifman et al. 1993, 1994; Wagner and Chew 1994; Epton and Dembart 1995). As noted in the introduction, these schemes are related to the scheme we will present below. They are, however, outside the scope of this paper.

## 3 Mathematical Preliminaries I

In this section, we briefly derive the multipole expansion of a charge distribution and refer the reader to Kellogg (1953), Jackson (1975), Wallace (1984), and Greengard (1988) for more detailed discussions.

If a point charge of strength $q$ is located at $P_0 = (x_0, y_0, z_0)$, then the potential and electrostatic field due to this charge at a distinct point $P = (x, y, z)$ are given by

$$\Phi = \frac{1}{R}$$

and

$$\vec{E} = -\nabla \Phi = \left(\frac{x-x_0}{R^3}, \frac{y-y_0}{R^3}, \frac{z-z_0}{R^3}\right),$$

respectively, where $R$ denotes the distance between points $P_0$ and $P$.

We would like to derive a series expansion for the potential at $P$ in terms of its distance from the origin $r$. For this, let the spherical coordinates of $P$ be $(r, \theta, \phi)$ and of $P_0$ be $(\rho, \alpha, \beta)$. Letting $\gamma$ be the angle between the vectors $P$ and $P_0$, we have from the law of cosines

$$R^2 = r^2 + \rho^2 - 2r\rho \cos \gamma,$$

with

$$\cos \gamma = \cos \theta \cos \alpha + \sin \theta \sin \alpha \cos (\phi - \beta).$$

Thus,

$$\frac{1}{R} = \frac{1}{r\sqrt{1 - 2\rho^2 \cos \gamma + \frac{\rho^2}{r^2}}} = \frac{1}{r\sqrt{1 - 2u\mu + \mu^2}},$$

6
having set

\[ \mu = \frac{\rho}{r} \quad \text{and} \quad u = \cos \gamma. \quad (25) \]

For \( \mu < 1 \), we may expand the inverse square root in powers of \( \mu \), resulting in a series of the form

\[ \frac{1}{\sqrt{1 - 2u\mu + \mu^2}} = \sum_{n=0}^{\infty} P_n(u)\mu^n \quad (26) \]

where

\[ P_0(u) = 1, \quad P_1(u) = u, \quad P_2(u) = \frac{3}{2}(u^2 - \frac{1}{3}), \ldots \quad (27) \]

and, in general, \( P_n(u) \) is the Legendre polynomial of degree \( n \). Our expression for the field now takes the form

\[ \frac{1}{R} = \sum_{n=0}^{\infty} \frac{r^{n+1}}{r^{n+1}} P_n(u). \quad (28) \]

The angular parameter \( u \), however, depends on both the source and the target locations. A more general representation will require the introduction of spherical harmonics, which are solutions of the Laplace equation obtained by separation of variables in spherical coordinates.

Any harmonic function \( \Phi \) can be expanded in the form

\[ \Phi = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \left( L_n^m r^n + \frac{M_n^m}{r^{n+1}} \right) Y_n^m(\theta, \phi). \quad (29) \]

The terms \( Y_n^m(\theta, \phi)r^n \) are referred to as spherical harmonics of degree \( n \) or solid harmonics, the terms \( Y_n^m(\theta, \phi)/r^{n+1} \) are called spherical harmonics of degree \(-n - 1\) or multipoles, and the coefficients \( L_n^m \) and \( M_n^m \) are known as the moments of the expansion.

The spherical harmonics can be expressed in terms of partial derivatives of \( 1/r \) (Wallace 1984) as

\[ \frac{Y_n^0(\theta, \phi)}{r^{n+1}} = A_n^0 \cdot \frac{\partial^n}{\partial z^n} \left( \frac{1}{r} \right). \quad (30) \]

For \( m > 0 \), we have

\[ \frac{Y_n^m(\theta, \phi)}{r^{n+1}} = A_n^m \cdot \left( \frac{\partial}{\partial x} - i \frac{\partial}{\partial y} \right)^m \left( \frac{\partial}{\partial z} \right)^{n-m} \left( \frac{1}{r} \right), \quad (31) \]

and

\[ \frac{Y_n^{-m}(\theta, \phi)}{r^{n+1}} = A_n^m \cdot \left( \frac{\partial}{\partial x} + i \frac{\partial}{\partial y} \right)^m \left( \frac{\partial}{\partial z} \right)^{n-m} \left( \frac{1}{r} \right), \quad (32) \]

where

\[ A_n^m = \frac{(-1)^n}{\sqrt{(n-m)! \cdot (n+m)!}}. \quad (33) \]

They also satisfy the relation

\[ Y_n^m(\theta, \phi) \equiv \sqrt{\frac{(n-|m|)!}{(n+|m|)!}} \cdot P_n^{|m|}(\cos \theta) e^{im\phi}, \quad (34) \]
where we have omitted the normalization factor of \( \sqrt{(2n+1)/4\pi} \), to match the definitions (30) - (32) given above. The special functions \( P_n^m \) are called associated Legendre functions and can be defined by the Rodrigues' formula

\[
P_n^m(x) = (-1)^m (1 - x^2)^{m/2} \frac{d^m}{dx^m} P_n(x).
\]

**Theorem 3.1 (Addition Theorem for Legendre Polynomials)** Let \( P \) and \( Q \) be points with spherical coordinates \( (r, \theta, \phi) \) and \((\rho, \alpha, \beta)\), respectively, and let \( \gamma \) be the angle subtended between them. Then

\[
P_n(\cos \gamma) = \sum_{m=-n}^{n} Y_n^{-m}(\alpha, \beta) \cdot Y_n^{m}(\theta, \phi).
\]

Combining Theorem 3.1 and eq. (28), we have

\[
\frac{1}{R} = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \rho^n Y_n^{-m}(\alpha, \beta) \cdot \frac{Y_n^{m}(\theta, \phi)}{r^{n+1}}.
\]

It is now straightforward to expand the field due to a collection of sources in terms of multipoles.

**Theorem 3.2 (Multipole Expansion).** Suppose that \( k \) charges of strengths \( \{q_i, \ i = 1, \ldots, k\} \) are located at the points \( \{Q_i = (\rho_i, \alpha_i, \beta_i), \ i = 1, \ldots, k\} \) with \(|\rho_i| < a\). Then for any \( P = (r, \theta, \phi) \in \mathbb{R}^3 \) with \( r > a \), the potential \( \Phi(P) \) is given by

\[
\Phi(P) = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \frac{M_n^m}{r^{n+1}} \cdot Y_n^{m}(\theta, \phi),
\]

where

\[
M_n^m = \sum_{i=1}^{k} q_i \cdot \rho_i^n \cdot Y_n^{-m}(\alpha_i, \beta_i).
\]

Furthermore, for any \( p \geq 1 \),

\[
\left| \Phi(P) - \sum_{n=0}^{p} \sum_{m=-n}^{n} \frac{M_n^m}{r^{n+1}} \cdot Y_n^{m}(\theta, \phi) \right| \leq \frac{A}{r - a} \left( \frac{a}{r} \right)^{p+1},
\]

where

\[
A = \sum_{i=1}^{k} |q_i|.
\]

**Proof:** The formula (38) follows from eq. (36) and superposition. The error bound is obtained from the triangle inequality and the fact that the ratios \( \rho_i/r \) are bounded from above by \( a/r \).

Suppose now that \( r = 2a \) in the context of the preceding theorem. Then the error bound (39) becomes

\[
\left| \Phi(P) - \sum_{n=0}^{p} \sum_{m=-n}^{n} \frac{M_n^m}{r^{n+1}} \cdot Y_n^{m}(\theta, \phi) \right| \leq \frac{A}{a} \left( \frac{1}{2} \right)^{p+1},
\]

and setting \( p = \log_2(1/\varepsilon) \) yields a precision \( \varepsilon \) relative to the ratio \( A/a \).
4 An $N \log N$ Algorithm

Theorem 3.2 is all that is required to construct a simple fast algorithm of arbitrary precision. To reduce the number of issues addressed, we assume that the particles are fairly homogeneously distributed in a square so that adaptive refinement is not required.

In order to make systematic use of multipole expansions, we introduce a hierarchy of boxes which refine the computational domain into smaller and smaller regions. At refinement level 0, we have the entire computational domain. Refinement level $l + 1$ is obtained recursively from level $l$ by subdivision of each box into eight equal parts. This yields a natural tree structure, where the eight boxes at level $l + 1$ obtained by subdivision of a box at level $l$ are considered its children.

**Definition 4.1** Two boxes are said to be near neighbors if they are at the same refinement level and share a boundary point (a box is a near neighbor of itself).

**Definition 4.2** Two boxes are said to be well separated if they are at the same refinement level and are not near neighbors.

**Definition 4.3** With each box $i$ is associated an interaction list, consisting of the children of the near neighbors of $i$’s parent which are well separated from box $i$ (Fig. 4).

**Definition 4.4** With each box $i$ at level $l$ is associated a multipole expansion $\Phi_{l,i}$ about the box center, which describes the far field induced by the particles contained inside the box.

The basic idea is to consider clusters of particles at successive levels of spatial refinement, and to compute interactions between distant clusters by means of multipole expansions when possible. It is clear that at levels 0 and 1, there are no pairs of boxes which are well separated. At level 2, on the other hand, sixty-four boxes have been created and there are a number of well separated pairs. Multipole expansions can then be used to compute interactions between these well separated pairs (Fig. 2) with rigorous bounds on the error. In fact, it is easy to see that the bound (39) applies with the ratio $a/r < 1/\sqrt{3}$. Thus, to achieve a given precision $\varepsilon$, we need to use $p = \log_3(1/\varepsilon)$ terms.

It remains to compute the interactions between particles contained in each box with those contained in the box’s near neighbors, and this is done recursively. We first refine each level 2 box to create level 3. For a given level 3 box, we then seek to determine which other level 3 boxes can be interacted with by means of multipole expansions. Since those boxes outside the region of the parent’s nearest neighbors are already accounted for (at level 2), they can be ignored. Since interactions with near neighbors cannot be accounted for accurately by means of an expansion, they can also be ignored for the moment. The remaining boxes correspond exactly to the interaction list defined above (Fig. 3).

The nature of the recursion is now clear. At every level, the multipole expansion is formed for each box due to the particles it contains. The resulting expansion is then evaluated for each particle in the region covered by its interaction list (Fig. 4).
Figure 2: The first step of the algorithm, depicted in two space dimensions for clarity. Interactions between particles in box X and its near neighbors (grey) are not computed. Interactions between well separated boxes are computed via multipole expansions.

Figure 3: The second step of the algorithm, depicted in two space dimensions. After refinement, note that the particles in the box marked X have already interacted with the most distant particles (light grey). They are now well separated from the particles in the white boxes, so that these interactions can be computed via multipole expansions. The near neighbor interactions (dark grey) are not computed.
Figure 4: Subsequent steps of the algorithm. The interaction list for box X is indicated in white. In three dimensions, it contains up to 189 boxes.

Figure 5: At the finest level, interactions with near neighbors are computed directly. In three dimensions, there are up to 27 near neighbors.

We halt the recursive process after roughly $\log_8 N$ levels of refinement. The amount of work done at each level is of the order $O(N)$. To see this, note first that approximately $N p^2$ operations are needed to create all expansions, since each particle contributes to $p^2$ expansion coefficients. Secondly, from the point of view of a single particle, there are at most 189 boxes (the maximum size of the interaction list) whose expansions are computed, so that $189 N p^2$ operations are needed for all evaluations.

At the finest level, we have created roughly $8 \log_8 N = N$ boxes and it remains only to compute interactions between nearest neighbors. By the assumption of homogeneity, there are $O(1)$ particles per box, so that this last step requires about $27 N$ operations (Fig. 5). The total cost is approximately

$$189 N p^2 \log_8 N + 27 N.$$ (42)

The algorithm just described is, in essence, a nonadaptive version of the one proposed by Barnes and Hut (1986), except that it achieves arbitrary precision through the use of high
order expansions. Two-dimensional schemes of this type are due to Van Dommelen and Run-  
densteiner (1989) and Odlyzko and Schönhage (1988). Unfortunately, while such schemes have good  
asymptotic work estimates, the three-dimensional versions provide only modest speedups  
at high precision for the values of $N$ encountered in present day applications. At $N = 100,000$,  
for example, seven digits of accuracy require $p \approx 20$, and the $N \log N$ scheme is only two to three  
times faster than the direct $O(N^2)$ method. In order to significantly accelerate the calculation,  
we need some further analytic machinery.

5 Mathematical Preliminaries II

The FMM relies on three translation operators, acting on either multipole (far field) or solid  
harmonic (local) expansions. They are described in the next three theorems (Greengard and  
Rokhlin 1988a; Greengard 1988).

**Theorem 5.1 (Translation of a Multipole Expansion)** Suppose that $l$ charges of strengths  
$q_1, q_2, \ldots, q_l$ are located inside the sphere $D$ of radius $a$ with center at $Q = (\rho, \alpha, \beta)$, and that  
for points $P = (r, \theta, \phi)$ outside $D$, the potential due to these charges is given by the multipole  
expansion

$$
\Phi(P) = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \frac{O_n^m}{r^{n+1}} \cdot Y_n^m(\theta', \phi'),
$$

where $P - Q = (r', \theta', \phi')$. Then for any point $P = (r, \theta, \phi)$ outside the sphere $D_1$ of radius  
$(a + \rho)$,

$$
\Phi(P) = \sum_{j=0}^{\infty} \sum_{k=-j}^{j} M_j^k \cdot Y_j^k(\theta, \phi),
$$

where

$$
M_j^k = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \frac{O_j^{k-m} \cdot i^{|k|-|m|-|k-m|} \cdot A_n^m \cdot A_j^{k-m} \cdot \rho^n \cdot Y_j^m(\alpha, \beta)}{A_j^m},
$$

with $A_n^m$ defined by eq. (33). Furthermore, for any $p \geq 1$,

$$
\left| \Phi(P) - \sum_{j=0}^{p} \sum_{k=-j}^{j} \frac{M_j^k}{r^{j+1}} \cdot Y_j^k(\theta, \phi) \right| \leq \left( \sum_{i=1}^{p} |q_i| \right) \left( \frac{a + \rho}{r} \right)^{p+1}.
$$

**Definition 5.1** The linear operator mapping old multipole coefficients $\{O_j^k\}$, $j = 0, \ldots, p$ and  
k = $-j, \ldots, j$, to new multipole coefficients $\{M_j^k\}$, $j = 0, \ldots, p$ and $k = -j, \ldots, j$, according to  
eq (45) will be denoted by $T_{MM}$.

**Theorem 5.2 (Conversion of a Multipole Expansion into a Local Expansion)** Suppose  
that $l$ charges of strengths $q_1, q_2, \ldots, q_l$ are located inside the sphere $D_Q$ of radius $a$ with center  
at $Q = (\rho, \alpha, \beta)$, and that $\rho > (c+1)a$ with $c > 1$. Then the corresponding multipole expansion
converges inside the sphere $D_0$ of radius $a$ centered at the origin. Inside $D_0$, the potential due to the charges $q_1, q_2,\ldots, q_l$ is described by a local expansion:

$$\Phi(P) = \sum_{j=0}^{\infty} \sum_{k=-j}^{j} L_j^k \cdot Y_j^k(\theta, \phi) \cdot r^j,$$

(47)

where

$$L_j^k = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \frac{O_n^m \cdot j^{k-m|-m|-|m|} \cdot A_n^m \cdot A_j^k \cdot Y_{j+n}^{m-k}(\alpha, \beta)}{(-1)^n A_{j+n}^{m-k} \cdot \rho^{j+n+1}},$$

(48)

with $A_n^m$ defined by eq. (33). Furthermore, for any $p \geq 1$,

$$\left| \Phi(P) - \sum_{j=0}^{p} \sum_{k=-j}^{j} L_j^k \cdot Y_j^k(\theta, \phi) \cdot r^j \right| \leq \left( \frac{\sum_{i=1}^{l} |q_i|}{c a - a} \right) \left( \frac{1}{c} \right)^{p+1}.$$

(49)

**Definition 5.2** The linear operator mapping truncated multipole expansion coefficients $\{O_j^k\}$, $j = 0,\ldots,p$ and $k = -j,\ldots,j$, to local coefficients $\{L_j^k\}$, $j = 0,\ldots,p$ and $k = -j,\ldots,j$, according to eq. (48) will be denoted by $T_{ML}$.

**Theorem 5.3 (Translation of a Local Expansion)**

Let $Q = (\rho, \alpha, \beta)$ be the origin of a local expansion

$$\Phi(P) = \sum_{n=0}^{p} \sum_{m=-n}^{n} O_n^m \cdot Y_n^m(\theta', \phi') \cdot r'^n,$$

(50)

where $P = (r, \theta, \phi)$ and $P - Q = (r', \theta', \phi')$. Then

$$\Phi(P) = \sum_{j=0}^{p} \sum_{k=-j}^{j} L_j^k \cdot Y_j^k(\theta, \phi) \cdot r^j,$$

(51)

where

$$L_j^k = \sum_{n=0}^{p} \sum_{m=-n}^{n} \frac{O_n^m \cdot j^{k-m|-m|-|m|} \cdot A_n^m \cdot A_j^k \cdot Y_{j+n}^{m-k}(\alpha, \beta) \cdot \rho^{n-j}}{(-1)^n-j^+ A_n^m},$$

(52)

with $A_n^m$ defined by eq. (33).

**Definition 5.3** The linear operator mapping old local expansion coefficients $\{O_n^m\}$, $n = 0,\ldots,p$ and $m = -n,\ldots,n$, to new local expansion coefficients $\{L_j^k\}$, $n = 0,\ldots,p$ and $m = -n,\ldots,n$, according to eq. (52) will be denoted by $T_{LL}$.
6 The original FMM

We can now construct a scheme with cost proportional to $N$, by using Theorem 5.2 to convert the far field expansion of a source box into a local expansion inside a target box, rather than by direct evaluation of the far field expansion at individual target positions.

**Definition 6.1** With each box $i$ at level $l$ is associated a local expansion $\Psi_{l,i}$ about the box center, which describes the potential field induced by all particles outside box $i$'s near neighbors.

**Definition 6.2** With each box $i$ at level $l$ is associated a local expansion $\Psi_{l,i}$ about the box center, which describes the potential field induced by all particles outside the near neighbors of $i$'s parent.

---

**ALGORITHM**

[Comment The parent of a box $j$ will be denoted by $p(j)$. The list of children of a box $j$ will be denoted by $c(j)$. The interaction list of a box $j$ will be denoted by $i\text{list}(j)$.]

**Upward Pass**

**Initialization**

[Comment Choose number of refinement levels $n \approx \log_8 N$, and the order of the multipole expansion desired $p$. The number of boxes at the finest level is then $8^n$, and the average number of particles per box is $s = N/(8^n)$.]

**Step 1**

[Comment Form multipole expansions $\Phi_{n,i}$ of potential field due to particles in each box about the box center at the finest mesh level, via Theorem 3.2.]

**Step 2**

For levels $l = n - 1, \ldots, 2$,

Form multipole expansion $\Phi_{l,j}$ about the center of each box at level $l$ by merging expansions from its eight children via Theorem 5.1.

$$\Phi_{l,j} = \sum_{k \in \text{child}(j)} T_{M,M} \Phi_{l+1,k}.$$  

**Downward Pass**

**Initialization**

Set $\Psi_{1,1} = \Psi_{1,2} = \cdots = \Psi_{1,8} = (0, 0, \ldots, 0)$.  

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Step 3

For levels $l = 2, \ldots, n$,

For each box $j$ at level $l$,

- Form the expansion $\tilde{\Psi}_{l,j}$ by using Theorem 5.3 to shift the local $\Psi$ expansion of $j$'s parent to $j$ itself.

$$\tilde{\Psi}_{l,j} = T_{LL} \Psi_{l-1,p(j)}.$$

- Form $\Psi_{l,j}$ by using Theorem 5.2 to convert the multipole expansion $\Phi_{l,k}$ of each box $k$ in the interaction list of $j$ to a local expansion about the center of box $j$, adding these local expansions together, and adding the result to $\tilde{\Psi}_{l,j}$.

$$\Psi_{l,j} = \tilde{\Psi}_{l,j} + \sum_{k \in \text{list}(j)} T_{ML} \Phi_{l,k}.$$

Step 4

For each particle in each box $j$ at the finest level $n$,

- evaluate $\Psi_{n,j}$ at the particle position.

Step 5

For each particle in each box $j$ at the finest level $n$,

- compute interactions with particles in near neighbor boxes directly.

Since $s$ is the average number of particles per box at the finest level, there are approximately $N/s$ boxes in the tree hierarchy. Therefore, Step 1 requires approximately $Np^2$ work, Step 2 requires $(N/s)p^4$ work, Step 3 requires $189(N/s)p^4$ work, Step 4 requires $Np^2$ work, and Step 5 requires $27N$'s work. Thus, a reasonable estimate for the total operation count is

$$191 \left( \frac{N}{s} \right) p^4 + 2Np^2 + 27Ns. \quad (53)$$

With $s = 2p^2$, the operation count becomes approximately

$$150Np^2. \quad (54)$$

This would appear to beat the estimate (42) for any $N$, but there is a subtle catch. The number of terms $p$ needed for a fixed precision in the $N \log N$ scheme is smaller than the number of terms needed in the FMM described above. To see why, consider two interacting cubes $A$ and $B$ of unit volume, with sources in $A$ and targets in $B$. The worst-case multipole error decays like $(\sqrt{3}/3)^p$, since $\sqrt{3}/2$ is the radius of the smallest sphere enclosing cube $A$ and $3/2$ is the shortest distance to a target in $B$. The conversion of a multipole expansion in $A$ to a
local expansion in \( B \), however, satisfies an error bound which depends on the smallest sphere enclosing \( B \) as well as the smallest sphere enclosing \( A \). From eq. (49), the worst case error is less than \((0.76)^p\), although with more detailed analysis, one can show that the error is bounded by \((0.75)^p\) (Petersen et al., 1995).

In the original FMM (Greengard and Rokhlin 1988; Greengard 1988), it was suggested that one redefine the nearest neighbor list to include “second nearest neighbors,” so that boxes which interact via multipole expansions are separated by at least two intervening boxes of the same size. The error can then be shown to decay approximately like \((0.4)^p\). However, the number of near neighbors increases from 27 to 125 and the size of the interaction list increases from 189 to 875.

It is clear that the major obstacle to achieving reasonable efficiency at high precision is the cost of the multipole to local translations \((189p^4\) operations per box). There are a number of schemes which have been suggested for reducing the cost of applying translation operators. The simplest is based on rotating the coordinate system so that the vector connecting the source box \( B \) and the target box \( C \) lies along the \( z \)-axis, shifting the expansion along the \( z \)-axis, and then rotating back to the original coordinate system.

### 6.1 The FMM using rotation matrices

We begin with the following obvious result

**Lemma 6.1** Consider a harmonic function given by

\[
\Phi(P) = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \left( L_n r^n + \frac{M_n}{r^{n+1}} \right) Y_n^m(\theta, \phi),
\]

where \((r, \theta, \phi)\) are the spherical coordinates of the point \( P \). If we rotate the coordinate system through an angle \( \beta \) in the positive sense about the \( z \)-axis, then

\[
\Phi(P) = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \left( \tilde{L}_n r^n + \tilde{M}_n \right) Y_n^m(\theta', \phi'),
\]

where \((r, \theta', \phi')\) are the new coordinates of \( P \),

\[
\tilde{L}_n = L_n e^{im\beta}, \quad \text{and} \quad \tilde{M}_n = M_n e^{im\beta}.
\]

**Definition 6.3** Given a rotation angle \( \beta \), the diagonal operator mapping old multipole coefficients to rotated multipole coefficients \((O_n^m \rightarrow O_n^m e^{im\beta})\) will be denoted by \( R_z(\beta) \).

We also need to be able to rotate the coordinate system about the \( y \)-axis.

**Lemma 6.2** Consider a harmonic function given by

\[
\Phi(P) = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \left( L_n r^n + \frac{M_n}{r^{n+1}} \right) Y_n^m(\theta, \phi),
\]
where \((r, \theta, \phi)\) are the spherical coordinates of the point \(P\). If we rotate the coordinate system through an angle \(\alpha\) in the positive sense about the \(y\)-axis, then there exist coefficients \(R(n, m, m', \alpha)\) such that

\[
\Phi(P) = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \left( \tilde{I}_n^{m'} r^n + \frac{\tilde{M}_n^{m'}}{r^{n+1}} \right) Y_n^{m'}(\theta', \phi'),
\]

where \((r, \theta, \phi')\) are the new coordinates of \(P\),

\[
\tilde{I}_n^{m'} = \sum_{m=-n}^{n} R(n, m, m', \alpha) L_n^m
\]

and

\[
\tilde{M}_n^{m'} = \sum_{m=-n}^{n} R(n, m, m', \alpha) M_n^m.
\]

**Proof:** See (Biedenharn and Louck 1981) for a complete discussion and for a variety of methods which can be used to compute the coefficients \(R(n, m, m', \alpha)\).

**Lemma 6.3** In order to shift a multipole expansion a distance \(\rho\) along the \(z\)-axis, one can replace eq. (45) with the simpler formula

\[
M_j^k = \sum_{n=0}^{\infty} \frac{O_{j-n}^k \cdot A_n^0 \cdot A_{j-n}^k \cdot \rho^n \cdot Y_n^0(1, 0)}{A_j^k}.
\]

In order to convert a multipole expansion centered at the origin into a local expansion centered at \((0, 0, \rho)\), one can replace eq. (48) with the simpler formula

\[
L_j^k = \sum_{n=0}^{\infty} \frac{O_n^m \cdot A_n^0 \cdot A_j^k \cdot Y_n^0(1, 0)}{(-1)^n A_{j+n}^0 \cdot \rho^{n+1}},
\]

In order to translate the center of a local expansion from the origin to the point \((0, 0, \rho)\), one can replace eq. (52) with the simpler formula

\[
L_j^k = \sum_{n=j}^{\rho} \frac{O_n^m \cdot A_{n-j}^0 \cdot A_j^k \cdot Y_{n-j}^0(1, 0) \cdot \rho^{n-j}}{(-1)^{n+j} \cdot A_n^k}.
\]

**Definition 6.4** Given a rotation angle \(\alpha\), the diagonal operator mapping old multipole coefficients to rotated multipole coefficients according to formula (55) or (56) will be denoted by \(R_y(\alpha)\). The special cases of the linear operators \(T_{MM}, T_{ML}, \) and \(T_{LL}\) which shift a distance \(\rho\) in the \(z\)-direction according to the formulae (57), (58), and (59) will be denoted by \(T_{MM}(\rho), T_{ML}(\rho), \) and \(T_{LL}(\rho)\).
We can now combine Lemmas 6.1, 6.2 and 6.3 to obtain the desired factorizations of $T_{MM}$, $T_{ML}$, $T_{LL}$.

**Lemma 6.4**

\[
T_{MM} = R_z(-\beta)R_y(-\alpha)T_{MM}^\rho R_y(\alpha)R_z(\beta),
\]

\[
T_{ML} = R_z(-\beta)R_y(-\alpha)T_{ML}^\rho R_y(\alpha)R_z(\beta),
\]

\[
T_{LL} = R_z(-\beta)R_y(-\alpha)T_{LL}^\rho R_y(\alpha)R_z(\beta),
\]

where $(\rho, \alpha, \beta)$ is desired shifting vector.

Clearly, the cost of applying $T_{MM}$, $T_{ML}$, or $T_{LL}$ by means of the preceding factorization is $O(p^2) + O(p^3) + O(p^3) + O(p^3) + O(p^2)$.

Thus, the total computational cost of the FMM can be reduced to approximately

\[
191 \left( \frac{N}{s} \right) 3p^3 + 2Np^2 + 27Ns.
\]

With $s = 3p^{3/2}$, the operation count becomes

\[
270Np^{3/2} + 2Np^2. \tag{60}
\]

7 Mathematical Preliminaries III

Over the last few years, a number of “fast” or diagonal translation schemes have been developed which require $O(p^2)$ work (Greengard and Rokhlin 1988b; Berman 1995; Elliott and Board 1996). Unfortunately, these schemes are all subject to certain numerical instabilities. The instabilities can be overcome, but at additional cost, the details of which we leave to the cited papers.

The latest generation of fast algorithms is based on combining multipole expansions with exponential or “plane wave” expansions. The reason for using exponentials is that translation corresponds to multiplication and, like the earlier fast schemes, requires only $O(p^2)$ work. Unlike in the earlier diagonal schemes, however, no numerical instabilities are encountered. The two-dimensional theory is described in (Hrycak and Rokhlin 1995), and we present the three-dimensional theory here.

**Remark 7.1** A complicating feature of the new approach is that six plane wave expansions will be associated with each box, one emanating from each face of the cube. To fix notation, we will refer to the $+z$ direction as up, to the $-z$ direction as down, to the $+y$ direction as north, to the $-y$ direction as south, to the $+x$ direction as east, and to the $-x$ direction as west. The interaction list for each box will be subdivided into six lists, one associated with each direction.
Definition 7.1 The Uplist for a box $B$ consists of those elements of the interaction list which lie above $B$ and are separated by at least one box in the $+z$-direction (Fig. 6). The Downlist for a box $B$ consists of those elements of the interaction list which lie below $B$ and are separated by at least one box in the $-z$-direction. The Northlist for a box $B$ consists of those elements of the interaction list which lie north of $B$, are separated by at least one box in the $+y$-direction, and are not contained in the Up or Down lists. The Southlist for a box $B$ consists of those elements of the interaction list which lie south of $B$, are separated by at least one box in the $-y$-direction, and are not contained in the Up or Down lists. The Eastlist for a box $B$ consists of those elements of the interaction list which lie east of $B$, are separated by at least one box in the $+x$-direction, and are not contained in the Up, Down, North, or South lists. The Westlist for a box $B$ consists of those elements of the interaction list which lie west of $B$, are separated by at least one box in the $-x$-direction, and are not contained in the Up, Down, North, or South lists.

It is easy to verify that the original interaction list is equal to the union of the Up, Down, North, South, East and West lists. It is also easy to verify that

$$C \in \text{Uplist}(B) \iff B \in \text{Downlist}(C)$$

$$C \in \text{Northlist}(B) \iff B \in \text{Southlist}(C)$$

$$C \in \text{Eastlist}(B) \iff B \in \text{Westlist}(C)$$

Given a source location $P = (x_0, y_0, z_0)$ and a target location $Q = (x, y, z)$, our starting point is the well-known integral representation (Morse and Feshbach 1953, p. 1256)

$$\frac{1}{\sqrt{(x - x_0)^2 + (y - y_0)^2 + (z - z_0)^2}} = \frac{1}{2\pi} \int_0^\infty e^{-\lambda(z - z_0)} e^{i\lambda((x - x_0) \cos \alpha + (y - y_0) \sin \alpha)} d\alpha d\lambda$$

(62)
\[ \int_{0}^{\infty} e^{-\lambda(z-z_0)} J_0(\lambda \sqrt{(x-x_0)^2 + (y-y_0)^2}) \, d\lambda, \]

valid for \( z > z_0 \).

To get a discrete representation, we must use an appropriate quadrature formula. The inner integral, with respect to \( \alpha \), is easily handled by the trapezoidal rule (which achieves spectral accuracy for periodic functions), but the outer integral requires more care. Laguerre quadrature is an appropriate choice here, but even better performance can be obtained using generalized Gaussian quadrature rules (Yarvin and Rokhlin 1996). These have been designed with the geometry of the interaction list in mind.

Because of the restriction that \( z > z_0 \), we will assume, for the moment, that the source \( P \) is contained in a box \( B \) and that the target \( Q \) lies in a box \( C \in \text{Uplist}(B) \). The following lemma describes several discrete approximations of the double integral in (62) as double sums.

**Lemma 7.1** Let \( P \in B \) and \( Q \in C \in \text{Uplist}(B) \), where \( B \) is a box of unit volume. Then

\[
\left| \frac{1}{r_{PQ}} - \sum_{k=1}^{9} \frac{w_k}{M(k)} \sum_{j=1}^{M(k)} e^{-\lambda_k[(z-z_0)-i(x-x_0)\cos \alpha_j-(y-y_0)\sin \alpha_j]} \right| < 10^{-3},
\]

(63)

where \( \alpha_j = 2\pi j/M(k) \), and the weights \( w_1, \ldots, w_9 \), nodes \( \lambda_1, \ldots, \lambda_9 \), and values \( M(1), \ldots, M(9) \) are given in section 12, Table 1. (The total number of exponentials required is 109.)

\[
\left| \frac{1}{r_{PQ}} - \sum_{k=1}^{18} \frac{w_k}{M(k)} \sum_{j=1}^{M(k)} e^{-\lambda_k[(z-z_0)-i(x-x_0)\cos \alpha_j-(y-y_0)\sin \alpha_j]} \right| < 10^{-6},
\]

(64)

where \( \alpha_j = 2\pi j/M(k) \), and the weights \( w_1, \ldots, w_{18} \), nodes \( \lambda_1, \ldots, \lambda_{18} \), and values \( M(1), \ldots, M(18) \) are given in section 12, Table 2. (The total number of exponentials required is 558.)

\[
\left| \frac{1}{r_{PQ}} - \sum_{k=1}^{30} \frac{w_k}{M(k)} \sum_{j=1}^{M(k)} e^{-\lambda_k[(z-z_0)-i(x-x_0)\cos \alpha_j-(y-y_0)\sin \alpha_j]} \right| < 0.5 \cdot 10^{-10},
\]

(65)

where \( \alpha_j = 2\pi j/M(k) \), and the weights \( w_1, \ldots, w_{30} \), nodes \( \lambda_1, \ldots, \lambda_{30} \), and values \( M(1), \ldots, M(30) \) are given in section 12, Table 3. (The total number of exponentials required is 1751.)

**Remark 7.2** The formulae (63)-(65) are somewhat complex, but have a simple interpretation. The outer sums use the generalized Gaussian weights and nodes \( \{w_k, \lambda_k\} \) obtained in (Yarvin and Rokhlin 1996) to approximate the outer integral (with respect to \( \lambda \)), while the inner sums use the trapezoidal rule to approximate the inner integral (with respect to \( \alpha \)). The number of nodes in each inner integral depends on the value \( \lambda_k \) for which the integration is being performed, and is denoted by \( M(k) \). These are derived from standard estimates concerning Bessel functions (Watson 1944, pp. 227, 255; Rokhlin 1995).
Remark 7.3 In the remainder of this paper, we will assume that the desired precision \( \varepsilon \) is clear from the context and will write

\[
\frac{1}{r_{pq}} \sum_{k=1}^{s(\varepsilon)} \sum_{j=1}^{M(k)} \frac{w_k}{M(k)} e^{-\lambda_k (z-z_0)} e^{i \lambda_k ((x-x_0) \cos \alpha_j + (y-y_0) \sin \alpha_j)} < \varepsilon, \tag{66}
\]

where \( \alpha_j = 2\pi j/M(k) \). This is a mild abuse of notation, since the weights, nodes and values \( M(k) \) depend on \( \varepsilon \) as well. The total number of exponential basis functions used will be denoted by \( S_{\exp} \), so that

\[
S_{\exp} = \sum_{k=1}^{s(\varepsilon)} M(k).
\]

Corollary 7.1 Let \( B \) be a box of unit volume centered at the origin containing \( N \) charges of strengths \( \{q_l, \ l = 1, \ldots, N\} \), located at the points \( \{Q_l = (x_l, y_l, z_l), \ l = 1, \ldots, N\} \). Then for any \( P \) contained in \( Uplist(B) \), the potential \( \Phi(P) \) satisfies

\[
|\Phi(P) - \sum_{k=1}^{s(\varepsilon)} \sum_{j=1}^{M(k)} W(k,j) e^{-\lambda_k z} e^{i \lambda_k ((x \cos \alpha_j + y \sin \alpha_j)} | < A \varepsilon, \tag{67}
\]

where \( A = \sum_{l=1}^{N} |q_l| \) and

\[
W(k,j) = \sum_{l=1}^{N} q_l e^{\lambda_k z_l} e^{-i \lambda_k ((x_l \cos \alpha_j + y_l \sin \alpha_l)} \tag{68}
\]

Corollary 7.2 (Diagonal translation) Let \( B \) be a box of unit volume centered at the origin containing \( N \) charges of strengths \( \{q_l, \ l = 1, \ldots, N\} \), located at the points \( \{Q_l = (x_l, y_l, z_l), \ l = 1, \ldots, N\} \) and let \( C \) be a box in \( Uplist(B) \) centered at \( (x_1, y_1, z_1) \). For \( P \in C \), let the potential \( \Phi(P) \) be approximated by the exponential expansion centered at the origin

\[
\Phi(P) = \sum_{k=1}^{s(\varepsilon)} \sum_{j=1}^{M(k)} W(k,j) e^{-\lambda_k z} e^{i \lambda_k ((x \cos \alpha_j + y \sin \alpha_j)} + O(\varepsilon). \tag{69}
\]

Then

\[
\Phi(P) = \sum_{k=1}^{s(\varepsilon)} \sum_{j=1}^{M(k)} V(k,j) e^{-\lambda_k z} e^{i \lambda_k ((x-z_1) \cos \alpha_j + (y-y_1) \sin \alpha_j)} + O(\varepsilon), \tag{70}
\]

where

\[
V(k,j) = W(k,j) e^{-\lambda_k z_1} e^{i \lambda_k ((x_1 \cos \alpha_j + y_1 \sin \alpha_j)} \tag{71}
\]

Definition 7.2 The diagonal operator mapping the original set of exponential expansion coefficients \( \{W(k,j)\} \) to the shifted exponential expansion coefficients \( \{V(k,j)\} \) according to eq. (71) will be denoted by \( D_{BC} \), where \( BC = (x_1, y_1, z_1) \) is the vector from the center of \( B \) to the center of \( C \).
In the FMM, we will be given the multipole expansion of a charge distribution for a box $B$ rather than the charge distribution itself, and will need to convert it to an exponential expansion. This is accomplished by the following theorem.

**Theorem 7.1** Let $B$ be a box of unit volume centered at the origin containing $N$ charges of strengths $\{q_i, \ i = 1, \ldots, N\}$, located at the points $\{Q_i = (x_i, y_i, z_i), \ i = 1, \ldots, N\}$. Let $P \in C \in Uplist(B)$ and suppose that the potential $\Phi(P)$ is given as the multipole expansion

$$\Phi(P) = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \frac{M_n^m}{r^{n+1}} \cdot Y_n^m(\theta, \phi). \quad (72)$$

Then

$$\Phi(P) - \sum_{k=1}^{\infty} \sum_{j=1}^{\infty} W(k, j)e^{-\lambda_k \xi} e^{i\lambda_k (\pi \cos \alpha_j + y \sin \alpha_j)} < A \epsilon, \quad (73)$$

where $A = \sum_{i=1}^{N} |q_i|$ and

$$W(k, j) = \frac{w_k}{M(k)} \sum_{m=-\infty}^{\infty} (-i)^{|m|} e^{i m \alpha_j} \sum_{n=|m|}^{\infty} \frac{M_n^m}{\sqrt{(n-m)!(n+m)!}} \lambda_k. \quad (74)$$

**Proof:** The formula (74) follows from the definitions (30) (31) and (32). The estimate (73) follows from Corollary 7.1. □

**Definition 7.3** The linear operator mapping a finite multipole expansion $\{M_n^m\}, \ n = 0, \ldots, p$ and $m = -n, \ldots, n$, to the corresponding set of coefficients in an exponential expansion $\{W(k, j)\}$ according to eq. (74) will be denoted by $C_{MX}$.

Once the multipole expansion for a source box has been converted into an exponential expansion (via Theorem 7.1) and translated to a target box center (via Corollary 7.2), we will need to convert the exponential expansion back into a solid harmonic series. The following theorem provides the necessary machinery.

**Theorem 7.2** Let $B$ be a box of unit volume containing $N$ charges of strengths $\{q_i, \ i = 1, \ldots, N\}$, located at the points $\{Q_i = (x_i, y_i, z_i), \ i = 1, \ldots, N\}$. Let $P$ be contained in a box $C \in Uplist(B)$, centered at the origin, and suppose that the potential $\Phi(P)$ is given as the exponential expansion

$$\Phi(P) = \sum_{k=1}^{\infty} \sum_{j=1}^{\infty} W(k, j)e^{-\lambda_k \xi} e^{i\lambda_k (\pi \cos \alpha_j + y \sin \alpha_j)} < A \epsilon, \quad (75)$$

where $A = \sum_{i=1}^{N} |q_i|$. Then

$$\Phi(P) - \sum_{n=0}^{\infty} \sum_{m=-n}^{n} L_n^m \cdot Y_n^m(\theta, \phi) \cdot r^n < A \epsilon, \quad (76)$$

22
Proof: Eq. (77) follows easily from the formula (Hobson 1955, p. 123)

\[
(z + ix \cos \alpha + iy \sin \alpha)^n = r^n \left\{ P_n(\cos \theta) + 2 \sum_{m=1}^{n} (i)^{-m} \frac{n!}{(n+m)!} (-1)^m P_n^m(\cos \theta) \cos m(\phi - \alpha) \right\},
\]

where \((r, \theta, \phi)\) are the spherical coordinates of the point with Cartesian coordinates \((x, y, z)\).

Remark 7.4 Theorems 7.1 and 7.2, like Theorem 5.2, are not quite the right tools needed to obtain rigorous error estimates for the FMM. In both cases, we have ignored the fact that the multipole and local expansions are truncated. It is straightforward but tedious to derive precise estimates, and we ignore this issue in the present paper. We should note that the nature of such estimates depends on how the multipole-to-exponential, multipole-to-solid harmonic or exponential-to-solid harmonic conversion is carried out. Formulae (74), (77) and (48) are the easiest to derive, being the Taylor expansions of the potential \(\Phi\). However, each of these conversions is simply a linear mapping from one set of basis functions to another. The formulae (77), (74), and (48) can be shown to correspond to minimizing the \(L_2\) error on the surface of a sphere enclosing the given source or target box. One could choose a variety of other possible projections, such as minimizing the \(L_2\) or \(L_\infty\) error on the surface of the corresponding box itself.

Remark 7.5 By inspection of formula (74), it is clear that the cost of applying the operator \(T_{MX}\) is \(p^2 s(e) + p S_{exp}\). The same is true for the operator \(T_{XL}\). It is also worth noting that fast Fourier transforms can be used to reduce the cost of the outer sum in the truncated version of formula (74) and the inner sum in the truncated version of formula (77).

Corollary 7.3 (Multipole to local factorization) Let \(B\) be a box of unit volume and \(C\) a box in \(Uplist(B)\). If \(T_{ML}\) is the translation operator converting the multipole expansion centered in \(B\) to the local expansion centered in \(C\), then

\[
T_{ML} = C_{XL} D_{B \cap C} C_{MX}.
\]

Remark 7.6 It is important to note that Lemma 7.1 provides a carefully designed quadrature formula which assumes that the source box \(B\) has unit volume and that the target is in \(B\)'s
In order to use these quadrature weights and nodes, we need to rescale the multipole and local expansions so that the box dimension always has unit volume. To accomplish this, if

\[ \Phi(P) = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \frac{M_n^m}{r^{n+1}} \cdot Y_n^m(\theta, \phi) \]  

is the multipole expansion for a box \( B \) of volume \( d^3 \), we simply write

\[ \Phi(P) = \sum_{n=0}^{\infty} \sum_{m=-n}^{n} \frac{(M_n^m/d^{n+1})}{(r/d)^{n+1}} \cdot Y_n^m(\theta, \phi). \]  

The local expansion for a target box in \( B \)'s interaction list is accumulated as

\[ \Phi(P) = \sum_{j=0}^{\infty} \sum_{k=-j}^{j} L_j^k \cdot Y_j^k(\theta, \phi) \cdot \left( \frac{r}{d} \right)^j. \]

**Corollary 7.4 (Scaled multipole to local factorization)** Let \( B \) be a box of volume \( d^3 \) and \( C \) a box in \( Uplist(B) \), with the vector from the center of \( B \) to the center of \( C \) given by \((x_1, y_1, z_1)\). If \( T_{ML} \) is the translation operator converting the multipole expansion centered in \( B \) to the local expansion centered in \( C \), then

\[ T_{ML} = D_{d,M} C_{XL} D_{B,C} C_{MX} D_{d,M}, \]  

where

\[ D_{d,M} M_n^m = M_n^m/d^{n+1}, \quad D_{d,L} L_n^m = L_n^m/d^n, \]

and \( B,C = B,C/d \).

The cost of a single multipole-to-local translation using the factorization of Corollary 7.4 is

\[ 2p^2 + 2p^2s(\epsilon) + 2pS_{exp} \approx 2p^3, \]

since \( s \approx p \) and \( S_{exp} \approx p^2 \). If each translation were carried out in this manner, we would not improve on the rotation based scheme discussed in section 6.1. However, once the multipole expansion for a box \( B \) has been converted to an exponential expansion (via the application of \( D_{d,M} \) and \( C_{MX} \)), it can be translated to each box in its \( Uplist \) at a cost of \( S_{exp} \approx p^2 \) operations. Conversely, once a box \( B \) has accumulated all the exponential expansions transmitted from its \( Downlist \) (see eq. (61)), a single application of the operators \( C_{XL} \) and \( D_{d,L} \) yields the local harmonic expansion describing the field due to the sources in the \( Downlist \) of box \( B \) (Fig. 7).

Up to this point, we have considered only the exponential representation needed to shift information in the upward (+z) direction. As noted in the beginning of this section, however, there are six outgoing directions which need to be accounted for. The most straightforward way of generating the appropriate expansions is to rotate the coordinate system so that the z-axis points in the desired direction. The following lemma provides the necessary formulae.
Figure 7: In the new FMM, a large number of multipole-to-local translations, costing $O(p^3)$ or $O(p^4)$ work, can be replaced by a large number of exponential translations, costing $O(p^2)$ work.

Lemma 7.2 Let $B$ be a box of volume $d^3$ and $C$ a "target" box. Let $T_{ML}$ be the translation operator converting the multipole expansion centered in $B$ to the local expansion centered in $C$. If $C \in \text{Downlist}(B)$, then

$$T_{ML}^{\text{Down}} = D_{d,L} R_y(-\pi) C_{XL} D_{BC} C_{MX} R_y(\pi) D_{d,M}.$$  

If $C \in \text{Eastlist}(B)$, then

$$T_{ML}^{\text{East}} = D_{d,L} R_y(-\pi/2) C_{XL} D_{BC} C_{MX} R_y(\pi/2) D_{d,M}.$$  

If $C \in \text{Westlist}(B)$, then

$$T_{ML}^{\text{West}} = D_{d,L} R_y(\pi/2) C_{XL} D_{BC} C_{MX} R_y(-\pi/2) D_{d,M}.$$  

If $C \in \text{Northlist}(B)$, then

$$T_{ML}^{\text{North}} = D_{d,L} R_y(-\pi/2) R_z(-\pi/2) C_{XL} D_{BC} C_{MX} R_y(\pi/2) R_z(\pi/2) D_{d,M}.$$  

If $C \in \text{Southlist}(B)$, then

$$T_{ML}^{\text{South}} = D_{d,L} R_y(\pi/2) R_z(-\pi/2) C_{XL} D_{BC} C_{MX} R_y(-\pi/2) R_z(\pi/2) D_{d,M},$$

where $BC$ is the appropriately scaled vector from the center of $B$ to the center of $C$ in the rotated coordinate system. The operators $R_x$ and $R_y$ are defined in section 6.1.

Definition 7.5 Let $T_{ML}^{\text{Up}}$ be given by the operator $T_{ML}$ defined in eq. (82). Then, for $\text{Dir} \in \{\text{Up, Down, East, West, North, South}\}$, we will write

$$T_{ML}^{\text{Dir}} = Q^{\text{Dir}} \ P_{BC}^{\text{Dir}},$$

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so that

\[ Q^\uparrow_p = D_{d,L} C_{XL}, \]
\[ P^\uparrow_p = C_{MX} D_{d,M}, \]
\[ Q^{\text{Down}} = D_{d,L} R_y(-\pi) C_{XL}, \]
\[ P^{\text{Down}} = C_{MX} R_y(\pi) D_{d,M}, \]

etc.

We are now in a position to describe the new FMM in detail.

8 The New FMM

ALGORITHM

[Comment The parent of a box \( j \) will be denoted by \( p(j) \). The list of children of a box \( j \) will be denoted by \( c(j) \). For each box \( j \), the "outgoing" exponential expansion with coefficients \( \{W(n, m)\}, n = 1, \ldots, s(e); m = 1, \ldots, M(n) \), will be denoted by \( W_j \). We will also associate an "incoming" exponential expansion with each box, denoted by \( V_j \).]

Upward Pass

Initialization

[Comment Choose number of refinement levels \( n \approx \log_8 N \), and the order of the multipole expansion desired \( p \). The number of boxes at the finest level is then \( 8^n \), and the average number of particles per box is \( s = N/(8^n) \).]

Step 1

Form multipole expansions \( \Phi_{n,i} \) of potential field due to particles in each box about the box center at the finest mesh level, via Theorem 3.2.

Step 2

Do for levels \( l = n - 1, \ldots, 2, \)

Form multipole expansion \( \Phi_{l,j} \) about the center of each box at level \( l \) by merging expansions from its eight children via Theorem 5.1.

\[ \Phi_{l,j} = \sum_{k \in c\text{hild}(j)} T_{MM} \Phi_{l+1,k}. \]

(In applying \( T_{MM} \), use the factorization of Lemma 6.4.)

End do

26
Downward Pass

Initialization

Set $\Psi_{1,1} = \Psi_{1,2} = \cdots = \Psi_{1,8} = (0, 0, \ldots, 0)$.

Step 3A

Do for levels $l = 2, \ldots, n$,

Form the expansion $\tilde{\Psi}_{l,j}$ for each box $j$ at level $l$ by using Theorem 5.3 to shift the local $\Psi$ expansion of $j$'s parent to $j$ itself.

$$\tilde{\Psi}_{l,j} = T_{LL}(l-1,p(j)).$$

(In applying $T_{LL}$, use the factorization of Lemma 6.4.)

Set $\Psi_{l,j} = \tilde{\Psi}_{l,j}$.

Step 3B

[Comment For each direction $\text{Dir} = \text{Up}, \text{Down}, \text{North}, \text{South}, \text{East}, \text{West}$, the opposite direction will be denoted by $-\text{Dir}$, so that $-\text{Up} = \text{Down}$, $-\text{Down} = \text{Up}$, etc. Thus, if a box $B$ sends an outgoing expansion in direction $\text{Dir}$ to Box $C$ on its $\text{Dirlist}$, then $C$ can be viewed as receiving the expansion from $B$ which is an element of its $-\text{Dirlist}$. (see eq. (61)).]

Do for $\text{Dir} = \text{Up}, \text{Down}, \text{North}, \text{South}, \text{East}, \text{West}$,

For each box $j$ at level $l$, convert the multipole expansion $\Phi_{l,j}$ into the "outgoing" exponential expansion for direction $\text{Dir}$.

$$W_j = P^\text{Dir} \Phi_{l,j}.$$  

For each box $j$ at level $l$, collect the "outgoing" exponential expansions from the $-\text{Dirlist}$ of box $j$ as an "incoming" exponential expansion

$$V_j = \sum_{k \in \text{-Dirlist}} D_{k,j} W_k,$$

where $k,j$ is the appropriately scaled vector from the center of box $k$ to the center of box $j$ in the rotated coordinate system.

For each box $j$ at level $l$, convert the accumulated "incoming" exponential expansion $V_j$ into a local harmonic expansion and add result to $\Psi_{l,j}$.

$$\Psi_{l,j} = \Psi_{l,j} + Q^\text{Dir} V_j.$$  

End do

End do

Step 4

For each particle in each box $j$ at the finest level $n$, evaluate $\Psi_{n,j}$ at the particle position.
Step 5

For each particle in each box \( j \) at the finest level \( n \), compute interactions with particles in near neighbor boxes directly.

Since we are using the rotation scheme for applying \( \mathcal{T}_{MM} \) and \( \mathcal{T}_{LL} \) in Steps 2 and 3A, these now require a total of \( 3p^3 (N/s) \) work, where \( s \) is the number of particles per box on the finest level. In Step 3B, the applications of the multipole to exponential operators \( \mathcal{P}^{Dir} \) and the exponential-to-local-operators \( \mathcal{Q}^{Dir} \) require a total of approximately \( 6p^3 (N/s) \) work, while the exponential translations require approximately \( 189p^2 (N/s) \) work. The total operation count is therefore of the order

\[
189 \frac{N}{s} p^2 + 2Np^2 + 27Ns + 6\frac{N}{s} p^3.
\]

With \( s = 2p \), the total operation count is about

\[
150Np + 5Np^2.
\]

8.1 Current improvements

There are a number of ways in which the algorithm described above has been accelerated. Symmetry considerations, for example, allow the pairs of operators \( \{ \mathcal{P}^{Up}, \mathcal{P}^{Down} \}, \{ \mathcal{P}^{North}, \mathcal{P}^{South} \}, \) and \( \{ \mathcal{P}^{East}, \mathcal{P}^{West} \} \) to be applied simultaneously. The same is true for the adjoint pairs \( \{ \mathcal{Q}^{Up}, \mathcal{Q}^{Down} \}, \) etc. Thus, the \( 6p^3 (N/s) \) work needed in Step 3B can be replaced by \( 3p^3 (N/s) \) work.

Even more significant is the fact that the number of translations per box can be reduced from 189 to less than 40. To see why, suppose that a box \( B \) at level \( l \) has eight children, denoted \( B_1, \ldots, B_8 \), and that boxes \( C_1, \ldots, C_J \) lie in the \( Uplist \) of each child. In the new FMM described above, we accumulated an “incoming” exponential expansion in each box \( C_j \) as

\[
V_j = \sum_{k=1}^{8} D_{B_k C_j} W_k,
\]

where \( W_k \) is the “outgoing” exponential expansion for \( B_k \). Repeating this for \( j = 1, \ldots, J \) requires a total of \( 8J \) translations. Since all translations are diagonal, however, it is easy to verify that

\[
V_j = \sum_{k=1}^{8} D_{BC_j} D_{B_k B} W_k
\]

\[
= D_{BC_j} \sum_{k=1}^{8} D_{B_k B} W_k.
\]

Thus, by first merging the “outgoing” expansions, and then translating their sum to each target box \( C_j \), only \( 8 + J \) translations are needed. It should be emphasized that this improvement relies
on the diagonal form of the operators. One could try to merge expansions in this manner in the context of the original FMM, but the local expansion coefficients computed with and without merging would not be the same. There would be a significant loss of precision, consistent with the error bound (49).

8.2 Further improvements

There are several ways in which the scheme can be accelerated which have not been incorporated into the existing code. The most significant of these is probably a change in the choice of the translation operators $T_{MM}$ and $T_{LL}$, as well as the multipole-to-exponential and exponential-to-local conversion operators $C_{MX}$ and $C_{XL}$. As mentioned previously, the obvious formulae (45), (52), (74), and (77) are obtained via Taylor expansion and are clearly not optimal. Preliminary numerical experiments indicate that replacing them with more carefully optimized tools will reduce the cost of these calculations within the FMM by a factor of three. Furthermore, the improvement described in Remark 7.5 has not yet been implemented; we are using the explicit matrix form of the discrete Fourier transform in applying $C_{MX}$ and $C_{XL}$, rather than the FFT.

The incorporation of all these modifications is likely to reduce the overall cost by a factor of two.

9 Numerical Results

The new FMM has been implemented in Fortran 77 and tested on uniform random distributions. The results of our experiments are summarized in Tables 1-4, with all times calculated in seconds using a Sun Ultra-1/140 workstation. In each table, the first column lists the number of particles, the second column lists the number of levels used in the multipole hierarchy, the third column lists the order of the multipole expansion used, and the fourth column lists the corresponding number of exponential basis functions. Columns five and six indicate the times required by the FMM and the direct calculation, respectively, and column seven lists the $l^2$ norm of the error in the FMM approximation

$$E = \left( \frac{\sum_{i=1}^{N} |\Phi(x_i) - \tilde{\Phi}(x_i)|^2}{\sum_{i=1}^{N} |\Phi(x_i)|^2} \right)^{1/2}.$$  \hspace{1cm} (83)

For the largest simulations, with $N > 10000$, we have carried out the direct calculation on a subset of only 100 particles. The stated times, indicated in parentheses, are then computed by extrapolation and the errors are obtained by restricting the formula (83) to this subset.

10 Extensions and Generalizations

The scheme presented in this paper is not adaptive and assumes that the distribution of points is reasonably uniform in space. In order to handle more general distributions, one needs to allow some regions to be subdivided to finer refinement levels than others. Adaptive structures of this type have been designed by several groups (Carrier et al. 1988; Van Dommelen and
Table 1: Timing results for the FMM using fifth order expansions and twenty-eight exponential basis functions.

<table>
<thead>
<tr>
<th>N</th>
<th>Levels</th>
<th>p</th>
<th>( S_{exp} )</th>
<th>( T_{FMM} )</th>
<th>( T_{dir} )</th>
<th>Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>500</td>
<td>3</td>
<td>5</td>
<td>28</td>
<td>0.18</td>
<td>0.20</td>
<td>4.5 \cdot 10^{-3}</td>
</tr>
<tr>
<td>5000</td>
<td>4</td>
<td>5</td>
<td>28</td>
<td>1.9</td>
<td>20.1</td>
<td>7.6 \cdot 10^{-3}</td>
</tr>
<tr>
<td>40000</td>
<td>5</td>
<td>5</td>
<td>28</td>
<td>20</td>
<td>(1461)</td>
<td>7.0 \cdot 10^{-3}</td>
</tr>
<tr>
<td>300000</td>
<td>6</td>
<td>5</td>
<td>28</td>
<td>175</td>
<td>(82475)</td>
<td>1.3 \cdot 10^{-2}</td>
</tr>
</tbody>
</table>

Table 2: Timing results for the FMM using ninth order expansions and 109 exponential basis functions.

<table>
<thead>
<tr>
<th>N</th>
<th>Levels</th>
<th>p</th>
<th>( S_{exp} )</th>
<th>( T_{FMM} )</th>
<th>( T_{dir} )</th>
<th>Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>2000</td>
<td>3</td>
<td>9</td>
<td>109</td>
<td>1.4</td>
<td>3.37</td>
<td>1.4 \cdot 10^{-4}</td>
</tr>
<tr>
<td>10000</td>
<td>4</td>
<td>9</td>
<td>109</td>
<td>7.9</td>
<td>83</td>
<td>3.6 \cdot 10^{-4}</td>
</tr>
<tr>
<td>80000</td>
<td>5</td>
<td>9</td>
<td>109</td>
<td>111</td>
<td>(5838)</td>
<td>4.1 \cdot 10^{-4}</td>
</tr>
</tbody>
</table>

Table 3: Timing results for the FMM using eighteenth order expansions and 558 exponential basis functions.

<table>
<thead>
<tr>
<th>N</th>
<th>Levels</th>
<th>p</th>
<th>( S_{exp} )</th>
<th>( T_{FMM} )</th>
<th>( T_{dir} )</th>
<th>Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>4000</td>
<td>3</td>
<td>18</td>
<td>558</td>
<td>8.3</td>
<td>13.4</td>
<td>1.1 \cdot 10^{-7}</td>
</tr>
<tr>
<td>25000</td>
<td>4</td>
<td>18</td>
<td>558</td>
<td>68</td>
<td>(567)</td>
<td>1.5 \cdot 10^{-7}</td>
</tr>
<tr>
<td>150000</td>
<td>5</td>
<td>18</td>
<td>558</td>
<td>495</td>
<td>(20100)</td>
<td>1.9 \cdot 10^{-7}</td>
</tr>
</tbody>
</table>

Table 4: Timing results for the FMM using thirtieth order expansions and 1751 exponential basis functions.

<table>
<thead>
<tr>
<th>N</th>
<th>Levels</th>
<th>p</th>
<th>( S_{exp} )</th>
<th>( T_{FMM} )</th>
<th>( T_{dir} )</th>
<th>Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>5000</td>
<td>3</td>
<td>30</td>
<td>1751</td>
<td>22</td>
<td>20.8</td>
<td>6.2 \cdot 10^{-12}</td>
</tr>
<tr>
<td>50000</td>
<td>4</td>
<td>30</td>
<td>1751</td>
<td>316</td>
<td>(2280)</td>
<td>6.2 \cdot 10^{-12}</td>
</tr>
</tbody>
</table>
and we are in the process of incorporating these structures into the new FMM.

While a number of techniques now exist for high frequency scattering problems (Rokhlin 1988, 1990, 1993; Canning 1989, 1992, 1993; Coifman and Meyer 1991; Bradie et al. 1993; Coifman et al. 1993, 1994; Wagner and Chew 1994; Epton and Dembart 1995), an important generalization of the algorithm of this paper is to the calculation of potentials governed by the Helmholtz equation at low frequency. By this we mean an environment in which the region of interest is no more than a few wavelengths in size, but contains a large number of discretization points (for example, due to the complexity of some structure being modeled). Algorithms for such problems are currently being designed.

11 Conclusions

A new version of the FMM has been developed. It is based on a new diagonal form for translation operators, and is significantly faster than previous implementations at any desired level of precision. Of particular interest is the fact that high precision calculations have been brought within practical reach.

12 Tables: Quadrature weights and nodes

Table 5: Columns 1 and 2 contain the nine weights and nodes needed for discretization of the outer integral in (62) at three digit accuracy. Column 3 contains the number of discretization points needed in the inner integral, which we denote by $M(k)$.

<table>
<thead>
<tr>
<th>Node</th>
<th>Weight</th>
<th>$M(k)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0992739673971</td>
<td>0.24776441819008</td>
<td>4</td>
</tr>
<tr>
<td>0.47725674637049</td>
<td>0.49188566500464</td>
<td>7</td>
</tr>
<tr>
<td>1.05536613821383</td>
<td>0.65378749137677</td>
<td>11</td>
</tr>
<tr>
<td>1.7675934354008</td>
<td>0.76433038408784</td>
<td>15</td>
</tr>
<tr>
<td>2.57342629351471</td>
<td>0.84376180565628</td>
<td>20</td>
</tr>
<tr>
<td>3.44824339201583</td>
<td>0.90445883985098</td>
<td>20</td>
</tr>
<tr>
<td>4.37680983554726</td>
<td>0.95378613136833</td>
<td>24</td>
</tr>
<tr>
<td>5.34895757205460</td>
<td>0.99670261613218</td>
<td>7</td>
</tr>
<tr>
<td>6.35765785313375</td>
<td>1.10429422730252</td>
<td>1</td>
</tr>
</tbody>
</table>
Table 6: Columns 1 and 2 contain the eighteen weights and nodes for discretization of the outer integral in (62) at six digit accuracy. Column 3 contains the number of discretization points needed in the inner integral, which we denote by $M(k)$.

<table>
<thead>
<tr>
<th>Node</th>
<th>Weight</th>
<th>$M(k)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05278852766117</td>
<td>0.13438265914335</td>
<td>5</td>
</tr>
<tr>
<td>0.26949859838931</td>
<td>0.29457752727395</td>
<td>8</td>
</tr>
<tr>
<td>0.63220353174689</td>
<td>0.42607819361148</td>
<td>12</td>
</tr>
<tr>
<td>1.11307564277608</td>
<td>0.53189220776549</td>
<td>16</td>
</tr>
<tr>
<td>1.689396140213</td>
<td>0.61787306245538</td>
<td>20</td>
</tr>
<tr>
<td>2.34376200469530</td>
<td>0.6863156078905</td>
<td>25</td>
</tr>
<tr>
<td>3.0626982907806</td>
<td>0.74749099381426</td>
<td>29</td>
</tr>
<tr>
<td>3.83562941265296</td>
<td>0.79699192718599</td>
<td>34</td>
</tr>
<tr>
<td>4.65424734321562</td>
<td>0.83917454386997</td>
<td>38</td>
</tr>
<tr>
<td>5.51209386593581</td>
<td>0.8757092283745</td>
<td>43</td>
</tr>
<tr>
<td>6.40421268377278</td>
<td>0.90792943590067</td>
<td>47</td>
</tr>
<tr>
<td>7.3268001906175</td>
<td>0.93698393724261</td>
<td>51</td>
</tr>
<tr>
<td>8.27740099258238</td>
<td>0.96382546688788</td>
<td>56</td>
</tr>
<tr>
<td>9.25397180602489</td>
<td>0.9893295679673</td>
<td>59</td>
</tr>
<tr>
<td>10.2556072734640</td>
<td>1.01438284597917</td>
<td>59</td>
</tr>
<tr>
<td>11.2820829787774</td>
<td>1.04003654374165</td>
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Table 7: Columns 1 and 2 contain the thirty weights and nodes for discretization of the outer integral in (62) at ten digit accuracy. Column 3 contains the number of discretization points needed in the inner integral, which we denote by $M(k)$.

<table>
<thead>
<tr>
<th>Node</th>
<th>Weight</th>
<th>$M(k)$</th>
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<tbody>
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<td>0.16861844033714</td>
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<tr>
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<tr>
<td>2.75739199003868</td>
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<td>4.80897515497095</td>
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<tr>
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</table>
References


[34] L. Greenberg and V. Rokhlin (1989), "On the evaluation of electrostatic interactions in molecular modeling", Chemica Scripta 29A, 139-144.


