Accelerating the Multipole to Local Field Translations

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1 Introduction

The fast multipole method (FMM) was originally motivated by N-body problems, such as those found in electrostatics and gravitation. This $O(N^2)$ problem is analogous to a matrix vector product (matvec). The FMM accelerates the computation of this matvec to just O(N) or $O(N \log N)$ depending on the underlying kernel function. The former is typically achieved when the problem is such that a subset of particles can be thought to interact only with a constant number of other particles, or groups, and is the case for the kernels we study, which can be considered to be asymptotically smooth. e.g.

$$\frac{1}{(r^2+c^2)^{\frac{n}{2}}}\tag{1}$$

where r = ||x-y||, c is a given parameter and $n \in \mathbb{N}$. The main idea of the FMM is to separate the near $(||x-y|| \to 0)$ and far-field $(||x-y|| \to \infty)$ components of a field actions of a set of particles (e.g. the electric/gravitational potential generated by static charges/masses). The near-field is computed directly, and the far-field is approximated using a divide and conquer strategy by recursion over a hierarchical tree that partitions the region of interest. The algorithm is explained in the literature, and we won't recount it here [3]. The algorithmic step of concern is the so-called 'M2L' operator, which acts as an approximation of a far-field at a given tree node. This is the most computationally intensive portion of the FMM algorithm, as it needs to be applied to all nodes in a hierarchical tree up to 189 times (in the 3D algorithm).

The computational optimisation of this operation is the focus of this note. Our choice of kernel functions is deliberate as they are compatible with so-called 'black box' FMM methods [2, 9]. These methods emerged two decades ago, and remain the basis for the highest performance FMM operators as of today [5, 8], performance in the FMM community is measured in terms of throughput (i.e. particles processed per second). The major benefit of these methods is that unlike the original 'analytical' formulations of the FMM, which rely on hand derived representations of multipole/local expansions for each kernel function, they rely *only* on kernel evaluations and are thus highly suitable for generic software implementations that can be specialised to work with a wide variety of kernels.

In this note I summarise different approaches to calculate the M2L operator for black box FMM methods, and various computational & mathematical optimisations that can be taken to accelerate its calculation. Our group is developing a new distributed FMM software in Rust as a part of a wider project to develop massively scalable integral equation software. We base our work on the KiFMM introduced by Ying et. al in [9], which relies on the method of fundamental solutions (MFS) to approximate the potentials generated by a set of particles, building on the work in [4].

In this note I summarise the main differences between black box FMMs, and common algorithmic approaches to calculate the M2L. We proceed to discuss a composite approach to approximating the M2L operation, building on previous work in [5, 6], which we believe will achieve a new benchmark performance for this operation.

...[TODO change eventually when I can take benchmarks!]

We also include an appendix on mathematical details, for example the FFT and SVD algorithms, which are used in approximations.

1.1 FMM Operator Basics

In 2D the single-layer Laplacian Green's function is.

$$G(\mathbf{x}, \mathbf{y}) = -\frac{1}{2\pi} \log \rho \tag{2}$$

where $\mathbf{r} = \mathbf{x} - \mathbf{y}$ and $\rho = |\mathbf{r}|$. It is useful to reformulate this using complex numbers, where $G(\mathbf{x}, \mathbf{y}) = \text{Re}\{\log(z_x - z_y)\}$ where z_x and z_y are complex numbers corresponding to source and target points on the plane. The key idea of the FMM is to encode the potentials fo a set of source densities using

a multipole expansion, and a local expansion at places far away from these sources.

Suppose that source densities are supported on a disk centered at z_c with radius r. Then for all z outside the disk with radius R, (R > r) we can represent the potential at z from the source densities using a set of coefficients $\{a_k, 0 \le k \le p\}$ as,

$$q(z) = a_0 \log(z - z_c) + \sum_{k=1}^{p} \frac{a_k}{(z - z_c)^k} + \mathcal{O}\left(\frac{r^p}{R_p}\right), \text{ Multipole Expansion}$$
(3)

On the other hand, if the source densities are outside the disk with radius R, the potential at a point z inside the disk with radius r can be represented by another set of coefficients $\{c_l, 0 \le k \le p\}$ as,

$$q(z) = \sum_{k=0}^{p} c_k (z - z_c)^k + \mathcal{O}\left(\frac{r^p}{R_p}\right), \text{ Local Expansion}$$
 (4)

The M2L translation transforms a multipole expansion of a box to a local expansion of another non-adjacent box. Instead of Laurent series, in 3D the far-field is represented using spherical-harmonics.

N.B KIFMM [9] relies on smoothness of kernel, as well as uniqueness of properly posed interior/exterior Dirichlet problem.

How do you prove that black box methods offer good approximations? Something that I need to ask about and write down at some point.

The formulation of the operator depends on the algorithm taken [9, 2], however in general we will get some kind of matvec, where a translation matrix is applied to a vector of multipole expansion coefficients.

1.2 Interaction Lists

This section is based on a similar discussion in [6], we adapt it here adding expository notes and calculations.

1.3 bbFMM

The basic idea is in this (and other interpolation based FMMs) is as follows. Letting $w_l(x)$ denote interpolating functions,

$$K(x,y) \approx \sum_{l} \sum_{m} K(x_{l}, y_{m}) w_{l}(x) w_{m}(y)$$
 (5)

ie. finding a low-rank approximation of the kernel. The advantage of such methods is that we only require the ability to evaluate the kernel at various points, no kernel dependent analytical expansion is required. The drawback is that the number of terms can be relatively large for a given error tolerance (verify?).

In Fong et al's approach, a Chebyshev interpolation scheme is used to approximate the far-field behaviour of the kernel. The M2L operator then consists of evaluating the field due to particles located at Chebyshev nodes, this can be effectively compressed using the SVD. If the kernel is translation invariant i.e. of the form K(x-y), the cose of the SVD precomputations reduces to $O(\log N)$ instead of O(N) as we only have to precompute for each level, reduces further if kernel can be scaled between levels.

1.4 **KIFMM**

The KIFMM uses the method of fundamental solutions (MFS) to approximate the field due to a set of discrete charges, an overview of which is given in the appendix. It slightly differs to the bbFMM in formulation, namely through the use of fictitious surfaces, however the fundamental computation remains the same - a matvec.

1.5 Summary

Both formulations essentially result in the same kind of computation being taken, some kind of matrix vector product which descrives a convolution operation. There are a couple of approaches that have been taken to accelerate this calculation, we provide an overview of each of these in the following sections, before concluding with a discussion on how we choose to implement the M2L for our software, in light of hardware and software optimisations taken in previous attempts.

Accelerating the M2L with the SVD 2

This is the method first presented in [2]. Consider the application of the M2L operator K to a multipole expansion w to get the check potential g.

$$g = Kw (6)$$

This can be approximated with a rank k SVD,

$$\tilde{g} = U_k \Sigma_k V_k^T \tag{7}$$

Stacking the M2L operators for all the source nodes in a given target node's interaction list can be done in two ways, column wise,

$$K_{\text{fat}} = [K^1, ..., K^3 16]$$
 (8)

$$= U\Sigma \left[V^{(1)T}, ..., V^{(316)T} \right]$$
 (9)

where we use the fact that there are at most 316 unique orientations for the M2L operator in 3D. Similarly they can be stacked row wise,

$$K_{\text{thin}} = \left[K^{1}; ...; K^{316}\right]$$

$$= \left[R^{(1)T}; ...; R^{(316)T}\right] \Lambda S^{T}$$
(10)

$$= [R^{(1)T}; ...; R^{(316)T}] \Lambda S^T$$
 (11)

we note that

$$K_{\text{thin}} = K_{\text{fat}}^T \tag{12}$$

for symmetric kernels.

We can do some algebra to reduce the application cost of K when we've done these two SVDs. Consider the application of a single M2L operator corresponding to a single source box in a target box's interaction list,

$$K^{(i)}w = R^{(i)}\Lambda S^T w \tag{13}$$

Using the fact that S is unitary, $S^TS = I$, we can insert into the above equation,

$$K^{(i)}w = R^{(i)}\Lambda S S^T S^T w \tag{14}$$

$$=K^{(i)}SS^Tw\tag{15}$$

$$= U\Sigma V^{(i)T} S S^T w \tag{16}$$

(17)

Now using the fact that U is also unitary, such that $U^TU=I$, we find

$$K^{(i)}w = UU^T U \Sigma V^{(i)T} S S^T w \tag{18}$$

$$= U[U^T U \Sigma V^{(i)T} S] S^T w \tag{19}$$

$$= U[U^T K^{(i)} S] S^T w (20)$$

The term in the brackets can be calculated using the low rank (k-rank) terms from the SVD,

$$[U^T K^{(i)} S] = \Sigma V^{(i)T} S \tag{21}$$

$$= U^T R^{(i)} \Lambda \tag{22}$$

We call this previous equation the compressed M2L operator,

$$C^{i,k} = U^T K^{(i)} S (23)$$

This object can be pre-computed for each unique interaction. The M2L operation can be then broken down into 4 steps

1. Find the 'compressed multipole expansion'

$$w_c = S^T w (24)$$

2. Compute the convolution to find the compressed check potential

$$g_c = \sum_{i \in I} C^{i,k} w_c \tag{25}$$

where the sum is over the interaction list I.

3. A post processing step to recover the check potential

$$g = Ug_c \tag{26}$$

4. The calculatation of the local expansion, as usual, in the KIFMM.

Doing this the convolution step is reduced to matrix vector products involving the compressed M2L matrix, which is only of size $k \times k$, rather than $6(p-1)^2 + 2$ where p is the expansion order.

2.1 Taking Advantage of Modern Compute Architectures

BLAS3 vs BLAS2

3 Accelerating M2L with FFT

The M2L operation can also be accelerated with a fast fourier transform (FFT). The M2L accelerated this way is quite natural, as it's simply a convolution operation, however computing it in practice can be be tricky. Here I document how I've managed to compute it, as well as a summary of the relevant FFT theory as a background.

3.1 The M2L Translation as a Fourier Convolution

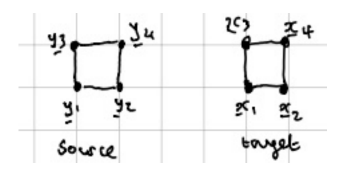
For the M2L operation we're computing the following convolution,

$$\phi(x) = \int G(x - y)q(y)dy \tag{27}$$

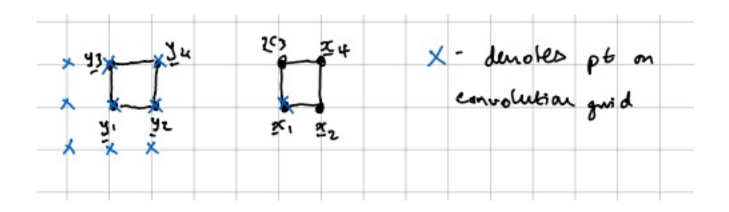
where we're attempting to compute the far-field potential as a convolution of the Green's function with a charge distribution (multipole expansion) at some local box. This is definitely somewhere we can apply the FT/FFT. How is this actually done in practice though, we're only concerned about the BBFMM [2] case where the charge distributions/multipole expansions are placed at regular intervals on the surface of a box enclosing a node in

the octree. In the literature there is a significant gap in describing how to actually setup the convolution operation such that we can apply the FFT to accelerate it, I illustrate it pictorially below.

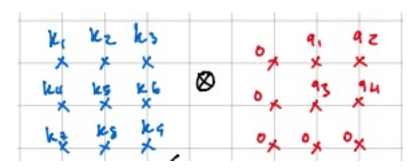
Consider two boxes (source and target) in 2D for simplicity, the expansion order is set to p=2



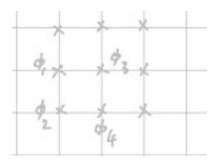
The boxes are referred to as lying in a 'surface grid', with the equivalent charges, $q_1, ..., q_4$, placed at $y_1, ... y_4$. We embed the unique kernel interactions between these two boxes on a so called 'convolution grid', we define them wrt to a fixed point - we can take this to be just x_1 . These can be pre-computed and stored.



The unique interactions define the convolution grid points. We label these $K_1, ..., K_9$. This is how the convolution is defined practically.



When we go ahead and compute this, taking care to 'flip' the kernel values, we find the potentials we're looking for embedded in at the following corresponding points on the convolution grid (flipped wrt to the positions of the equivalent densities). Only the four positions that correspond to the positions of the original equivalent densities are significant, the remainder can be ignored.



We can accelerate this convolution computation using the FFT as normal. Expressing this more explicitly, when we compute the check potential during the M2L operator during the KIFMM, we compute an approximation of the following integral,

$$\phi^{c}(\mathbf{x}) = \int_{y \in Y} G(\mathbf{x} - \mathbf{y}) q(\mathbf{y}) d\mathbf{y}$$
 (28)

where $\mathbf{x} \in X$ are points on the target surface, $\mathbf{y} \in Y$ are points on the source surface and $q(\mathbf{y})$ are discrete charges placed at source surface points and $\phi^c(\mathbf{x})$ is the check potential at a target surface point. Applying Fourier convolution theorem,

$$\phi^{c}(\mathbf{x}) = \mathcal{F}^{-1} \left[\mathcal{F}G(\xi) * \mathcal{F}q(\xi) \right] (\mathbf{x})$$
(29)

We notice that $G(\xi)$ must contain all the unique kernel evaluations between points on the source/target surface, and that when convolved with $q(\xi)$ we must recover the potentials in a predictable way (in order to create index maps to the write matrix elements).

This is done by defining a 'convolution grid', which is an extension of the surface discretisation of the KIFMM. This extension is defined by the unique evaluations of the Green's function with respect to a single point on the target surface grid. By doing this, the convolution can be drawn around the source surface grid in a formulaic way each time, as represented by the above figure. Representing the points on the target and surface grids by their indices in their discrete form, i.e. $\mathbf{y} = y_{ijk}$, we can write the matrix elements of the kernel evaluations represented on this convolution grid as,

$$\underline{G}_{ijk} = G(x_{000} - y_{ijk}) \tag{30}$$

where x_{000} corresponds to the lower left corner of the target surface, though in principle any point on the target surface could be used, they will result in different mappings between the final convolved form and the potentials we're seeking to identify. This results in a 3D sequence,

$$G[i,j,k] = \underline{G}_{ijk} \tag{31}$$

where the indices, e.g. $i \in I$, extend over the indices of the axes of the convolution grid.

Next we discuss padding. For optimum performance, the size of the FFT in each dimension should be a power of two. So we begin by padding the convolution grid with zeros,

$$G^{pad}[i', j', k'] = \begin{cases} & \text{if } 0 \le i' \le P - M \\ 0 & \text{and } 0 \le j' \le Q - N \\ & \text{and } 0 \le k' \le R - K \end{cases}$$
(32)

Where P, Q, R correspond to the dimensions of the sequence before padding, and M, N, K correspond to the next largest power of two of the size each of these dimensions, i', j', k' are the indices of the padded array. The remainder of the array is filled by the original sequence G[i, j, k].

Similarly, we must pad the sequence of discrete charges to match the dimensions of the padded kernel sequence in order to compute the FFT. We start by creating a creating a new convolution grid, and placing the discrete charges at their corresponding positions from the source surface grid. Once this is complete, we have a sequence q[i, j, k] with the same dimensions as the kernel sequence. We then choose the following padding,

$$q^{pad}[i',j',k'] = \begin{cases} & \text{if } P - M \le i' \le P \\ 0 & \text{and } Q - N \le j' \le Q \\ & \text{and } R - K \le k' \le R \end{cases}$$
 (33)

Where P,Q,R are the same as for the kernel sequence, note M,N,K are the same for the kernel and charge sequence now. This choice of padding for both sequences as well as taking the convolution grid with respect to x_{000} is fortuitous. Noting that in the computation of the FFT convolution we must flip the kernel, we find that our sequence of potentials lie at the indices [P-M-1:P,Q-N-1:Q,R-K-1:R] of the final FFT computed result. Looking up the potentials associated with each point on the target surface grid is exactly equivalent to looking up their associated index in the subsequence of the result indexed by [P-M-1:P,Q-N-1:Q,R-K-1:R].

3.1.1 Accelerating the Hadamard Product

The Hadamard product is the most computationally intensive part of the above scheme. Here I spell out how to accelerate it using explicit SIMD instructions and careful data organisation.

Instead of computing the convolution in the preceding section for a single source and target box, we now consider a set of siblings together,

$$S = \bigcup_{i=1}^{N=8} S_i \tag{34}$$

For a given M2L interaction, we'll have a sequence corresponding to the unique kernel interactions,

$$G_l[i,j,k] (35)$$

We generally pre-compute and store these for use, so we'll have single sequence for a given M2L interaction,

$$\hat{G}_l[i,j,k] \tag{36}$$

Now we can compute, the Hadamard product of each sibling's discrete charge sequence with this sequence. By computing this way, we retain $\hat{G}[i,j,k]$ in cache, and the element wise Hadamard product is an 8×8 operation which we can write as an explicit SIMD instruction. The result is a matrix, H, the element H_{pq} corresponds to the p^{th} element of the q^{th} sibling in the sibling set's Hadamard product.

We can extend this scheme by 'stacking' together sibling sets for this M2L translation, and processing them together. Iterating over all unique M2L translations in a given level of the FMM algorithm, which can be done in parallel.

Algorithm 1 M2L Convolution

```
Require: level l

1: for Translation Vector t do

2: in parallel do

3: Result for these sibling set, H

4: for Siblings FFT sequence, \hat{S}, Kernel FFT sequence, \hat{G} do

5: H[subI, subJ] = \text{hadamard} \cdot 8\text{x}8(\hat{S}_{8x8}, \hat{G}_{8x8})

6: end for

7: end for
```

3.2 Taking Advantage of Modern CPU Architectures

We want to batch together computations sibling interactions and take advantage of SIMD to maximally take advantage of the cache hierarchies in modern CPUs. One strategy is as follows, we conclude by contrasting it with the BLAS3/SVD approach, and include some numerical benchmarks to contrast the two ...

4 A New Software and Algorithm for M2L operators

4.1 Rusty Field

Software design of Rust field.

4.2 FFT based M2L with Symmetries

Algorithm description that leverages symmetries in the FFT M2L.

5 Appendix

5.1 Fourier Transform Theoretical Background

A lot of the theoretical background I want to keep at hand is taken from the excellent course notes [7]. I summarise the key aspects here as related to the FFT, especially when discussing padding/indexing, as these issues come up most pertinently in real implementations.

5.1.1 Going from Fourier Series to Fourier Transforms

Starting off with Fourier Series (FS), i.e. representing periodic functions using a periodic (trig) basis, and generalising to non-periodic (i.e. ∞ period) functions takes us to Fourier Transforms (FT).

Q: Is the sum of two periodic functions also periodic?

A: No if you're a mathematician, e.g. cos(t) and $cos(\sqrt{2}t)$ are each periodic with periods 2π and $2\pi/\sqrt{2}$ resp. But the sum is not periodic. ie. no common divisors in the periods.

When considering a sum of sinusoids, as Fourier pitched,

$$\sum_{n=1}^{N} A_n \sin(n\theta + \phi_n) \tag{37}$$

The sum is also periodic as the frequencies are multiples of the fundamental frequency $1/2\pi$.

It's more common to write a general trig sum as,

$$\frac{a_0}{2} + \sum_{n=1}^{N} (a_n \cos(2\pi nt) + b_n \sin(2\pi nt))$$
 (38)

Where the zeroth component is often referred to as a DC component (from electrical engineering contexts). The half is a simplifying factor that comes up. Expressing this instead using complex exponentials, the sum can be written as,

$$\sum_{n=-N}^{N} c_n e^{2\pi i n t} \tag{39}$$

One can refer to RHB to see how the coefficients are related between forms. In particular we find $c_0 = a_0/2$. The complex conjugate property of the coefficients,

$$c - n = \bar{c_n} \tag{40}$$

is important, it allows us to group terms such that

$$\sum_{n=-N}^{N} c_n e^{2\pi i n t} = 2 \operatorname{Re} \left\{ \sum_{n=0}^{N} c_n e^{2\pi i n t} \right\}$$
 (41)

Our goal is to express a general periodic function f(t) as an FS.

$$f(t) = \sum_{-N}^{N} c_n e^{2\pi i n t} \tag{42}$$

Take a given coefficient, can we solve for it?

$$f(t) = \sum_{-N}^{N} c_n e^{2\pi i n t} \tag{43}$$

$$e^{-2\pi ikt}f(t) = e^{-2\pi ikt} \sum_{-N}^{N} c_n e^{2\pi int}$$
 (44)

Therefore,

$$c_k = e^{-2\pi i k t} f(t) - \sum_{n=-N, n \neq k}^{N} c_n e^{2\pi i (n-k)t}$$
(45)

We've pulled the coefficient out, but the expression involves all the other coefficients! Instead, we can try and integrate both sides over 0 to 1 (any function can be made to have this period if it's periodic). The integrals in the sum all cancel out,

$$\int_{0}^{1} e^{2\pi(n-k)t} dt = \frac{1}{2\pi i(n-k)} e^{2\pi i(n-k)t} \Big|_{t=0}^{t=1} = 0$$
 (46)

With this trick, the expression for the coefficient reduces to,

$$c_k = \int_0^1 e^{-2\pi i k t} f(t) dt \tag{47}$$

We haven't stated whether any periodic function *can* be expressed in such a way that we can apply this analysis, but if we can express it in the periodic form we started off with, we have a way of evaluating the coefficients.

Note in particular that the zeroth coefficients corresponds to an average value of the function over its period.

$$\hat{f}(0) = \int_0^1 f(t)dt \tag{48}$$

The case when all the coefficients are real is when the signal is real and even. For then,

$$\bar{\hat{f}}(n) = \hat{f}(-n) = \int_0^1 e^{-2\pi i(-n)t} f(t)dt = \int_0^1 e^{2\pi i nt} f(t)dt$$
 (49)

$$= -\int_0^{-1} e^{-2\pi i n s} f(-s) ds, \text{ subs t} = -s, \text{ changing lims}$$
 (50)

$$= \int -1^0 e^{-2\pi i n s} f(-s) ds, \text{ even f(s)}$$

$$\tag{51}$$

$$=\hat{f}(n) \tag{52}$$

So the coefficients are real. The evenness of f seems to pass over into its fourier coefficients too.

We haven't yet answered when a periodic function can be approximated by a fourier series . . . We're basically allowed to if $f(t) \in L^2([0,1])$ as then the integral defining its Fourier coefficients exists. The fourier approximation is the best approximation in $L^2([0,1])$ by a trigonemtric polynomial of degree N. The complex exponentials form a basis for this space, and the partial sums converge to f(t) in its norm,

$$\lim_{N \to \infty} \left\| \sum_{n = -N}^{N} \hat{f}(n) e^{-2\pi i n t} - f(t) \right\| = 0$$
 (53)

For Fourier Transforms, lets start off by considering a box function.

$$\Pi(t) = \begin{cases} 1 & \text{if } |t| < 1/2, \\ 0 & \text{if } |t| \ge 1/2. \end{cases}$$
 (54)

This isn't periodic, and doesn't have an FS. However, if we make it repeat with intervals T, we can find a representation with coefficients given by,

$$c_n = \frac{1}{T} \int_0^T e^{-2\pi i n t/T} f(t) dt = \frac{1}{T} \int_{-T/2}^{T/2} e^{-2\pi i n t/T} f(t) dt = \frac{1}{\pi n} \sin(\frac{\pi n}{T})$$
 (55)

The coefficients tend to 0 for large T as 1/T, to compensate for this we can scale by T. Using a change of variables s = n/T we can write,

$$\Pi(s) = \frac{\sin(\pi s)}{\pi s}$$

We can now take a limit as $T \to \infty$,

$$\hat{\Pi}(s) = \int_{-\infty}^{\infty} e^{-2\pi i s t} \Pi(t) dt = \int_{-1/2}^{1/2} e^{-2\pi i s t} \cdot 1 dt = \frac{\sin(\pi s)}{\pi s}$$
 (56)

We are lead to the same idea - scale the Fourier coefficients by T - if we had started off periodising any function that is zero outside of some interval and letting the period tend to infinity. This gives us the following definition for Fourier Transforms,

$$\hat{f}(s) = \int_{-\infty}^{\infty} e^{-2\pi i s t} f(t) dt \tag{57}$$

where the coefficients are in general complex. FTs produce continuous spectra, in contrast to a discrete set of (potentially infinitely many) frequencies as in FS.

We can push this to get a definition for the dual, the inverse transform. Again supposing that we have a non-periodic function that we can say is zero outside of an interval, we find an expression for its FS, and fourier coefficients

$$f(t) = \sum_{n = -\infty}^{\infty} c_n e^{2\pi i n t/T}$$
(58)

$$c_n = \frac{1}{T} \int_{-T/2}^{T/2} e^{-2\pi i n t/T} f(t) dt = \frac{1}{T} \int_{-\infty}^{\infty} e^{-2\pi i n t/T} f(t) dt$$
 (59)

$$=\frac{1}{T}\hat{f}(\frac{n}{T}) = \frac{1}{T}\hat{f}(s) \tag{61}$$

Plugging back in, and thinking of Riemann sum to approximate an integral,

$$f(t) = \sum_{-\infty}^{\infty} \frac{1}{T} \hat{f}(s_n) e^{2\pi i s_n t} = \sum_{-\infty}^{\infty} \hat{f}(s_n) e^{2\pi i s_n t} \Delta s \approx \int_{-\infty}^{\infty} \hat{f}(s_n) e^{2\pi i s_n t} ds \quad (62)$$

5.1.2 The Convolution

In general we want to modify signals by each other. Is there a combination of signals f(t) and g(t) such that in the frequency domain the FT is:

$$\mathcal{F}g(s)\mathcal{F}f(s)$$

i.e. is there a combination of the signals such that frequency components are scaled by each other?

Very roughly, we find,

$$\mathcal{F}g(s)\mathcal{F}f(s) = \int_{-\infty}^{\infty} e^{-2\pi i s t} g(t) ds \int_{-\infty}^{\infty} e^{-2\pi i s x} f(x) dx \tag{63}$$

$$= \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} e^{-2\pi i s(t+x)} g(t) f(x) dt dx \tag{64}$$

using the change of variable u = t + x for the inner integral,

$$\int_{-\infty}^{\infty} \left(\int_{-\infty}^{\infty} e^{-2\pi i s u} g(u - x) du \right) f(x) dx \tag{65}$$

switching the order of integration,

$$\int_{-\infty}^{\infty} e^{-2\pi i s u} \left(\int_{-\infty}^{\infty} g(u - x) f(x) dx \right) du \tag{66}$$

The inner integral can be seen to be a function of u, we can write it as h(u), the outer integral reduces to:

$$\int_{-\infty}^{\infty} e^{-2\pi i s u} h(u) du = \mathcal{F}h(s) \tag{67}$$

This defines our convolution,

$$(g * f)(t) = h(t) = \int_{-\infty}^{\infty} g(t - x)f(x)dx \tag{68}$$

And the following theorem,

$$\mathcal{F}(g * f)(s) = \mathcal{F}g(s)\mathcal{F}f(s) \tag{69}$$

Most significantly for us, convolving in the time domain reduces to a multiliplication in the frequency domain.

The convolution is defined by flipping the kernel, and dragging it over the signal.

5.1.3 Discrete Fourier Transforms, and the Fast Fourier Transform

We want to find a discrete analogue to the FT for real signals which are sampled at a certain rate.

Let's suppose that f(t) is zero outside of an interval $0 \le t \le L$, similarly the FT $\mathcal{F}f(s)$ is assumed zero outside of $0 \le s \le 2B$ (indexing is easier if we ignore negative frequencies), L and B are both integers.

According to Shannon, we can reconstruct f(t) perfectly if we sample at a rate of 2B per second. So in total we want,

$$N = \frac{L}{1/2B} = 2BL$$

evenly spaced samples, notice that this is even. Sampled at points,

$$t_0 = 0, t_1 = \frac{1}{2B}, ..., t_{N-1} = \frac{N-1}{2B}$$

$$f_{discrete}(t) = \sum_{n=0}^{N-1} \delta(t - t_n) f(t_n)$$
 (70)

and therefore,

$$\mathcal{F}f_{discrete}(t) = \sum_{n=0}^{N-1} f(t_n) \mathcal{F}\delta(t - t_n) = \sum_{n=0}^{N-1} f(t_n) e^{-2\pi i s t_n}$$
 (71)

which is almost what we need, it's the continuous FT of the sampled form of f(t).

Shifting to the frequency domain, we find the number of sample points to be,

$$N = \frac{2B}{1/L} = 2BL$$

the same as in the time domain. We base the discrete version of the FT using the discrete version of the signal,

$$F(s_0) = \sum_{n=0}^{N-1} f(t_n) e^{-2\pi i s_0 t_n}$$
(72)

etc. We now have a way of converting from the discrete signal to the discrete FT,

$$F(s_m) = \sum_{n=0}^{N-1} f(t_n) e^{-2\pi i s_m t_n}$$
(73)

It's possible to link this to the continuous case by discretising the integral defining a continuous FT, we see that this sum (up to a scaling) comes out. using,

$$t_n = \frac{n}{2B}, \quad s_m = \frac{m}{L} \tag{74}$$

we can write in terms of indices,

$$F(s_m) = \sum_{n=0}^{N-1} f(t_n) e^{-2\pi i n m/2BL} = \sum_{n=0}^{N-1} f(t_n) e^{-2\pi i n m/N}$$
 (75)

Thinking about io as sequences of numbers, we can write in 'array' form, where the transform is just defined on a sequence.

$$\mathbf{F}[m] = \sum_{n=0}^{N-1} \mathbf{f}[n] e^{-2\pi i m n/N}, \quad m = 0, 1, ..., N-1$$
 (76)

The input sequence can be complex, it's not less valid, but the output sequence is always complex.

A common notation is to write the complex exponentials as,

$$\omega = e^{2\pi i/N} = \omega_N$$

s.t.

$$\omega_N^N = 1$$

for any integer n and k,

$$\omega_N^{Nn} = 1$$

$$\omega_N^{Nn+k} = \omega_N^k$$

and,

$$\omega_N^{N/2} = -1$$

so,

$$\omega_N^{kN/2} = (-1)^k$$

We write a vector of the Nth roots of unity as,

$$\omega = (1, \omega, \omega^2, ..., \omega^{N-1})$$

the components,

$$\omega^k[m] = \omega^{km}$$

The DFT can be thought of as a linear transfrom between \mathbb{C}^N to \mathbb{C}^N . This linear transform can be explicitly written out as a matrix, which I won't bother with here, look at 257 in [7].

This is a dense $N \times N$ matrix! The FT is in general hard to compute, hence the revolution of the FFT which can do it in log-linear time.

5.2 N-Dimensional DFT

The DFT takes a sequence of complex numbers $u_0, u_1, ..., u_{N-1}$ and transforms them into another sequence of complex numbers $\hat{u}_0, \hat{u}_1, ..., 0\hat{u}_{N-1}$, the forward and backwarf transfroms is defined as,

$$\hat{u}_k = \frac{1}{N} \sum_{j=0}^{N-1} u_j e^{-ikx_j}, \quad k = 0, 1, ..., N-1$$
(77)

$$u_k = \frac{1}{N} \sum_{j=0}^{N-1} \hat{u}_j e^{ikx_j}, \quad k = 0, 1, ..., N-1$$
 (78)

where $x_j = 2\pi j/N$. If instead the data is arranged in a multidimensional array, $u_{j_0,j_1,...,j_{d-1}}$ where there are d index sets $j_m = 0, 1, ..., N_{m-1}, m \in 0, 1, ..., d-1$ with $N_m = ||j_m||$ being the length of j_m . A forward d-dimensional DFT of the d-dimensional array will be computed as,

$$\hat{u}_{k_0,k_1,\dots,k_{d-1}} = \sum_{j_0=0}^{N_0-1} \left(\frac{\omega_0^{k_0j_0}}{N_0} \sum_{j_1=0}^{N_1-1} \left(\frac{\omega_1^{k_1j_1}}{N_1} \dots \sum_{j_d-1}^{N_{d-1}-1} \frac{\omega_{d-1}^{k_{d-1}j_{d-1}}}{N_{d-1}} u_{j_0,j_1,\dots,j_{d-1}} \right) \right)$$

$$(79)$$

where
$$w_j = e^{\frac{-2\pi i}{N_j}}$$

Normalisation in this context refers to the scaling of the output to be independent of input size. That is if you double the size of your input, e.g. via padding, the amplitude of the output frequencies should not change. This is done by dividing the output of the FFT by the length of the input array (or its square root depending on convention).

5.3 Method of Fundamental Solutions (MFS)

We can use MFS to approximate the field due to a set of discrete charges, by using the fundamental solutions as a basis, this is the approach taken by the authors of the KIFMM [9]. This note is based on exposition presented in [1], and adapted to the field approximations required in the KIFMM - our problem of interest.

Our goal during the FMM is to approximate either (a) the potential generated by particles inside a tree node in the exterior, (b) the evaluate the approximation of the potential generated by particles outside of a tree node inside its interior. The KIFMM achieves this using the method of fundamental solutions (MFS). Consider problem (a), sketched in figure 5.3.

The MFS approximation is to approximate the potential by using a linear cobination of fundamental solutions of the problem at hand,

$$u(\mathbf{x}) \approx u^N(\mathbf{x}) = \sum_{i \in I_v^B} G(\mathbf{x}, \mathbf{y}_i) \phi_i$$
 (80)

Here we use N points on an 'equivalent' surface, that encloses the node containing the points, specified by the index set of the points I_u^B the reason for denoting this with u in particular will become apparent later, ϕ_i are some unknown densities representing the charges contained in the node which we are to solve for and G(.,.) denotes the Green's function. In order to do this, we can calculate directly the potential generated by points contained in the node at the (blue) check surface,

$$\sum_{i \in I_s^B} G(\mathbf{x}, \mathbf{y}_i) \phi_i = q^{B, u} \tag{81}$$

which we call the check potential. The potential due to our charges is guaranteed to be equivalent to that generated by our equivalent charges due to the uniqueness of the exterior problem for our kernels of interest (e.g. Laplace).

We see that our MFS approximation is analogous to an approximation of a single-layer potential defined along the equivalent surface, Γ_e , with some continuous density ϕ

$$u(\mathbf{x}) \approx \int_{\Gamma_e} G(\mathbf{x}, \mathbf{y}) \phi(\mathbf{y}) d\mathbf{y}$$
 (82)

if we replace $\phi(\mathbf{y})$ with $\sum_{j=1}^{N} \phi_i \delta(\mathbf{y} - \mathbf{y}_j)$. Therefore the MFS approximates a first-kind integral equation, which is in general ill-conditioned [TODO - lookup in Kress].

We can solve this ill-conditioned system using Tikhonov regularisation to convert this first kind integral equation into a second-kind integral equation. Written in matrix form our problem becomes

$$K\phi = q \tag{83}$$

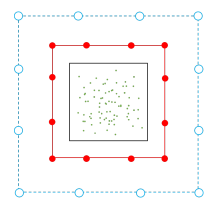


Figure 1: A set of particles contained in a tree node (green), enclosed by the tree node (black), an equivalent surface (red) and a check surface (blue check).

Where K is the discretisation of the matrix elements described by equation (82), ϕ is a vector containing the densities supported on the equivalent surface, and q is a vector of check potentials evaluated on the check surface. Applying Tikhonov regularisation,

$$\phi = (\alpha I + K^*K)^{-1}q \tag{84}$$

we can solve for q, where we choose the regularisation parameter experimentally.

The procedure described above amounts to the P2M operator in the FMM, the other FMM operators in the KIFMM are calculated in in a highly similar manner and we won't bother describing them here. The key thing to note here is that this approximation method relies only on evaluating the fundamental solutions (kernels) of the PDE, we don't need to actually write out an analytic expansion to approximate the potential in the exterior, instead we just need to find a set of equivalent densities and we can reconstruct the exterior field using the MFS. The literature shows that one can achieve exponential convergence with increasing N [1].

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