A Kernel-independent Adaptive Fast Multipole Method

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Joint work with George Biros and Denis Zorin
Problem Statement

Given

- $G$ an elliptic PDE kernel, e.g. $G(x, x') = \frac{1}{|x-x'|}$
- $\{x_i\}$ points in $\mathbb{R}^d$ ($d = 2, 3$)
- $\{\phi_i\}$ charges

Evaluate potential

$$u_i = \sum_{j=1}^{N} G(x_i, x_j) \phi_j, \ i = 1, \ldots, N$$

- Direct evaluation: $\mathcal{O}(N^2)$
- Barnes-Hut algorithm: $\mathcal{O}(N \log N)$
- Fast Multipole Method (FMM): $\mathcal{O}(N)$
  - Greengard and Rokhlin (JCP, 1987)
Contribution

New kernel-independent FMM
- Use only kernel evaluations
- Extend FMM to general kernels
- Efficiency and accurate

MPI based parallel implementation
- Scale up to 3000 processors
- 1.6 Tflops peak / 1.13Tflops sustained performance on 2.1 billion unknowns

3D high-order BIE solver
- Laplace and Stokes equation
- Smooth boundary
Applications

Electrostatics

Gravitation

Fluid dynamics

Molecular dynamics
Outline

Classical FMM

New algorithm

Parallel algorithm

Boundary integral equation solver
Idea of Classical FMM

Direct evaluation

Barnes-Hut

FMM, 2-level

FMM, one-level

Well separated
2 efficient representation
3 efficient translation
Domain Partitioning

Quadtree / Octree

- Leaf box contains at most $s$ charges

For each box $B$, define

- Near range
- Far range
- Interaction list

$B$ and its far range are well separated
Efficient Representations

Multipole expansion (ME)
\{\phi_j\} at \{z_j\} : |z_j - z_C| < r, for |z - z_C| > R
\[ u(z) \approx a_0 \log(z - z_C) + \sum_{k=1}^{p} \frac{a_k}{(z - z_C)^k} \]
\[ a_0 = \sum_{j=1}^{m} \phi_j \quad a_k = \sum_{j=1}^{m} \frac{-\phi_i(z_i - z_C)^k}{k} \]

Local expansion (LE)
\{\phi_j\} at \{z_j\} : |z_j - z_C| > R, for |z - z_C| < r
\[ u(z) \approx \sum_{k=0}^{p} c_k (z - z_C)^k \]
\[ c_0 = \sum_{j=1}^{m} \phi_j \log(z_C - z_j) \quad c_l = \sum_{j=1}^{m} \frac{-\phi_j}{l \cdot (z_j - z_C)^l} \]

m is a constant → Both expansions are efficient
Efficient Translations

M2M: multipole to multipole
- From child to parent box

M2L: multipole to local
- From B’s interaction list to B

L2L: local to local
- From parent to child box

All translations are efficient
Algorithm

1. Compute multipole expansion (ME) for each box
   - If leaf box, compute from exact sources in that box
   - If non-leaf box, compute from its children’s ME using M2M

2. Compute local expansion (LE) for each box
   - Use M2L to translate the ME of the boxes in the interaction list and add to the LE of the current box
   - Use L2L to translate the LE of the parent box and add to the LE of the current box

   Invariance: LE represents the potential from far field accurately

3. For each box, compute potential at each of its point by combining
   - Contribution from LE of this box
   - Potential from points in near range of this box
Kernel Dependency

Expansions and translations depend on underlying PDE

3D implementation only available for Laplace and Helmholtz

10 years to get efficient implementation for Laplace kernel

FMM can be extremely difficult for kernels look like this:

\[
D(x, y) = A ((r \cdot n)I + n \otimes r) + B(r \otimes n) + C(r \cdot n)(r \otimes r)
\]

\[
A = -\frac{f_{rrr}}{r} - (d-3)\frac{f_{rr}}{r^2} + (d-3)\frac{f_r}{r^3}
B = -p + 2\frac{f_{rr}}{r^2} - 2\frac{f_r}{r^3}
C = 2\frac{f_{rrr}}{r^3} - 6\frac{f_{rr}}{r^4} + 6\frac{f_r}{r^5}
\]

\[
f = \begin{cases} 
\frac{1}{2\pi\lambda^2} (\ln(\frac{1}{r}) - k_0(\lambda r)) & (2D) \\
\frac{1}{4\pi\lambda^2} (\frac{1}{r} - \frac{1}{r} e^{-\lambda r}) & (3D)
\end{cases}
\]

\[
p = \begin{cases} 
\frac{1}{2\pi r^2} & (2D) \\
\frac{1}{4\pi r^4} & (3D)
\end{cases}
\]
Outline

Classical FMM

New algorithm

Parallel algorithm

Boundary integral equation solver
Previous Work

FMM without multipole
- Anderson ’92
- Not fully kernel independent

FFT-based methods
- Phillips, White ’97, pre-corrected FFT
- Bruno, Kunyansky ’01, improved version

Wavelet-based methods
- BCR ’91 and Alpert et al. ’93
Idea of New Algorithm

An electrostatic example
- Represent sources with equivalent densities on the circle
- Match check potential on the boundary of the far field

Extend this idea to FMM
- Multipole expansion—upwards equivalent density
- Local expansion—downs equivalent density
- M2M, M2L and L2L translation—linear solution for the matching process
Efficient Representations

Upwards equivalent density

\[
\int_{y_B,u} G(x,y) \phi_{B,u} \, dy = \sum_{i \in I^B} G(x,y_i) \phi_i = u^{B,u}
\]
for any \( x \in x^{B,u} \).

Downwards equivalent density

\[
\int_{y_B,d} G(x,y) \phi_{B,d} \, dy = \sum_{i \in I^B} G(x,y_i) \phi_i = u^{B,d}
\]
for any \( x \in x^{B,d} \).
Efficient Translations

M2M \( \int_{y_{B,u}} G(x, y) \phi_{B,u}^B(y) \, dy = \int_{y_{A,u}} G(x, y) \phi_{A,u}^A(y) \, dy \)
- From child to parent

for all \( x \in x^{B,u} \)

M2L \( \int_{y_{B,d}} G(x, y) \phi_{B,d}^B(y) \, dy = \int_{y_{A,u}} G(x, y) \phi_{A,u}^A(y) \, dy \)
- From interaction list to box

for all \( x \in x^{B,d} \)

L2L \( \int_{y_{B,d}} G(x, y) \phi_{B,d}^B(y) \, dy = \int_{y_{A,d}} G(x, y) \phi_{A,d}^A(y) \, dy \)
- From parent to child

for all \( x \in x^{B,d} \)
Discretization

3D case

- Equivalent and check surfaces are cubes
- Boundary points of Cartesian grid as discretization points

Discretized translations (M2M, M2L and L2L) are small matrices

- Independent of the position of the box involved
- Pre-compute and store them
Acceleration of M2L

2D
- Store the SVD of the M2L matrix

3D
- Pad interior Cartesian nodes with zeros
- Kernel translation invariant
- M2L translation = small 3D discrete convolution
- Use 3D FFT to compute it
Eq. Density Approximation

2D

Single-layer Navier.

3D

Single-layer Laplacian.
# 3D Laplace Kernel

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The particles are distributed on the surface of a sphere.

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The particles are uniformly distributed inside a cube.
## 3D Navier Kernel

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The particles are uniformly distributed in a cube.
Outline

Classical FMM

New algorithm

Parallel algorithm

Boundary integral equation solver
Motivation

Real applications solve huge N-body problem
- E.g. astrophysics, molecular dynamics
- New algorithm can still be slow

Parallel linear solvers are available
- E.g. PetSc
- FMM (often serves as matvec) better be parallel as well

FMM algorithm can be nicely parallelized
- Most operations are local
- E.g. parent to child, box to its interaction list …
Two Problems

Octree can be huge
- Needs to be generated and stored in a distributed way
- Each processor keeps its Local Essential Tree (LET)
- Most work is done on LET

Synchronization due to data dependence
- Parent node’s ME depends on child node’s ME, but two nodes can reside in different processors
- Use linearity to remove data dependence
  - All translations are linear operator
Results

Tests setup

- Laplace, Stokes, Navier and modified versions
- Uniform and non-uniform distributions in box $[-1, 1]^3$

Software

- C++, uses PETSC, FFTW

Hardware

- TCS1 HP Alpha cluster at Pittsburgh supercomputing center
  - 750 compute nodes
  - 4 1GHz processors and 4GB memory per node
  - 187GB/s quadrics interconnection network
Fixed Size Scalability

Laplace equation, uniform distribution

3.2 million points in total

![Graphs showing CPU cycles per particle and MFlops efficiency vs. number of processors.]
Isogranular Scalability

Laplace equation, uniform distribution
0.2 million points per processor
3000 Processor Run

Stokes equation, uniform distribution

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Summary

- Fixed size tests, 80% efficiency up to 256 processors
- Isogranular tests, good efficiency up to 3000 processors
- 1.6 Tflops peak / 1.13 Tflops sustained for 2.1 billion unknowns
- Tree construction bottleneck for 1024+ processors
- Some load imbalance in non-uniform distribution
Outline

Classical FMM

New algorithm

Parallel algorithm

Boundary integral equation solver
3D BIE Solver

Properties

- Uses second-kind integral formulation
- Works for Laplace, Stokes equations
- General smooth domain boundary
- Based on Nyström discretization
- High-order accurate
- Efficient
Components

High-order accuracy $\rightarrow$ high-order boundary representation

- Manifold-based boundary representation method
  (ACM Transaction of Graphics 23(3), ’04)

Smooth (dense) part of the integral operator

- Kernel independent FMM algorithm

Singular (diagonal) part of the integral operator

- Use local partition of unity to isolate singularity
- Local high-order integration in polar coordinates
- Similar to (Bruno and Kunyansky ’01)
Summary

New kernel-independent FMM
- Use only kernel evaluations
- Extend FMM to general kernels
- Efficiency and accurate

MPI based parallel implementation
- Scale up to 3000 processors
- 1.6 Tflops peak / 1.13Tflops sustained performance on 2.1 billion unknowns

3D high-order BIE solver
- Laplace and Stokes equation
- Smooth boundary
Thank you