Randomized algorithms and fast direct solvers for elliptic PDEs

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Research support by:
Outline of talk:

• Randomized dimension reduction.

• Direct solvers for linear elliptic PDEs. (Sparse matrices.)

• Direct solvers for linear elliptic PDEs via integral equations. (Dense matrices.)

Slides: http://people.maths.ox.ac.uk/martinsson/main_talks.html
(or Google “Gunnar Martinsson”)
Randomized dimension reduction

Let \( \{ \mathbf{a}(j) \}_{j=1}^{n} \) be a set of points in \( \mathbb{R}^m \), where \( m \) is very large. Consider tasks such as:

- Suppose the points almost live on a linear subspace of (small) dimension \( k \).
  Find a basis for the “best” subspace.
- Suppose the points almost live on a low-dimensional nonlinear manifold.
  Find a parameterization of the manifold.
- Given \( k \), find the subset of \( k \) vectors with maximal spanning volume.
- Given \( k \), find for each vector \( \mathbf{a}(j) \) its \( k \) closest neighbors.
- Partition the points into clusters.

(Note: Some problems have well-defined solutions, some do not. The first can be solved with algorithms with complexity \( O(mn) \), some are combinatorially hard.)

Idea: Find a “nice” embedding \( f : \mathbb{R}^m \to \mathbb{R}^d \) for \( d \ll m \) that is almost isometric:

\[
\| f(\mathbf{a}(i)) - f(\mathbf{a}(j)) \| \approx \| \mathbf{a}(i) - \mathbf{a}(j) \|, \quad \forall i, j \in \{1, 2, 3, \ldots, n\}.
\]

Then solve the problems for the vectors \( \{ f(\mathbf{a}(j)) \}_{j=1}^{n} \) in \( \mathbb{R}^d \).

Lemma [Johnson-Lindenstrauss]: For \( d \sim \log(n) \), there exists an orthogonal projection that “approximately” preserves distances.
To be precise, we have:

**Lemma [Johnson-Lindenstrauss]:** Let $\varepsilon$ be a real number such that $\varepsilon \in (0, 1)$, let $n$ be a positive integer, and let $d$ be an integer such that

\[ d \geq \frac{4 \left( \frac{\varepsilon^2}{2} - \frac{\varepsilon^3}{3} \right)^{-1}}{\log(n)}. \]

Then for any set $V$ of $n$ points in $\mathbb{R}^m$, there is a map $f : \mathbb{R}^m \to \mathbb{R}^d$ such that

\[ (1 - \varepsilon) \|u - v\|^2 \leq \|f(u) - f(v)\| \leq (1 + \varepsilon) \|u - v\|^2, \quad \forall u, v \in V. \]

You can prove that if you pick $d$ as specified, and repeatedly draw a Gaussian random matrix $G$ of size $d \times m$, then the likelihood is larger than zero that the map

\[ f(x) = \frac{1}{\sqrt{d}} G x \]

satisfies the criteria.

**Practical problem:** You have two very bad choices:

1. Pick a small $\varepsilon$; then you get small distortions, but a huge $d$ since $d \sim \frac{8}{\varepsilon^2} \log(n)$.
2. Pick $\varepsilon$ that is not close to 0, then distortions are large.
Question: Is it possible to build algorithms that combine the powerful dimension reduction capability of randomized projections with the accuracy and robustness of classical deterministic methods?
**Question:** Is it possible to build algorithms that combine the powerful dimension reduction capability of randomized projections with the accuracy and robustness of classical deterministic methods?

**Putative answer:** Yes — use a two-stage approach:

(A) *Randomized pre-conditioner:*

In a pre-computation, random projections are used to create low-dimensional sketches of the high-dimensional data. These sketches are somewhat distorted, but approximately preserve key properties to very high probability.

(B) *Deterministic post-processing:*

Once a sketch of the data has been constructed in Stage A, classical deterministic techniques are used to compute desired quantities to very high accuracy, *starting directly from the original high-dimensional data.*
Example 1 of two-stage approach: Nearest neighbor search in $\mathbb{R}^m$

**Objective:** Suppose you are given $n$ points $\{a^{(j)}\}_{j=1}^n$ in $\mathbb{R}^m$. The coordinate matrix is

$$A = \begin{bmatrix} a^{(1)} & a^{(2)} & \ldots & a^{(n)} \end{bmatrix} \in \mathbb{R}^{m \times n}.$$

How do you find the $k$ nearest neighbors for every point?

If $m$ is “small” (say $m \leq 10$ or so), then you have several options; you can, e.g, sort the points into a tree based on hierarchically partitioning space (a “kd-tree”).

**Problem:** Classical techniques of this type get very expensive as $m$ grows.

**Simple idea:** Use a random map to project onto low-dimensional space. This “sort of” preserves distances. Execute a fast search there.

**Improved idea:** The output from a single random projection is unreliable. But, you can repeat the experiment several times, use these to generate a list of *candidates* for the nearest neighbors, and then compute exact distances to find the $k$ closest among the candidates.
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How do you find the $k$ nearest neighbors for every point?

(A) **Randomized probing of data:** Use a Johnson-Lindenstrauss random projection to map the $n$-particle problem in $\mathbb{R}^m$ (where $m$ is large) to an $n$-particle problem in $\mathbb{R}^d$ where $d \sim \log n$. Run a deterministic nearest-neighbor search in $\mathbb{R}^d$ and store a list of the $\ell$ nearest neighbors for each particle (for simplicity, one can set $\ell = k$). Then repeat the process several times. If for a given particle a previously undetected neighbor is discovered, then simply add it to a list of potential neighbors.

(B) **Deterministic post-processing:** The randomized probing will result in a list of putative neighbors that typically contains more than $k$ elements. But it is now easy to compute the pairwise distances in the original space $\mathbb{R}^m$ to judge which candidates in the list are the $k$ nearest neighbors.

*Jones, Osipov, Rokhlin, 2011*
**Example 2 of two-stage approach: Randomized SVD**

**Objective:** Given an $m \times n$ matrix $A$, find an approximate rank-$k$ partial SVD:

$$A \approx U D V^*$$

$m \times n$ $m \times k$ $k \times k$ $k \times n$

where $U$ and $V$ are orthonormal, and $D$ is diagonal.

---

(A) **Randomized pre-conditioner:**

Use randomized projection methods to form an approximate basis for the range of the matrix.

(B) **Deterministic post-processing:**

Restrict the matrix to the subspace determined in Stage A, and perform expensive but accurate computations on the resulting smaller matrix.

Observe that distortions in the randomized projections are fine, since all we need is a subspace that captures “the essential” part of the range. Pollution from unwanted singular modes is harmless, as long as we capture the dominant ones. The risk of missing the dominant ones is for practical purposes zero.
Example 2 of two-stage approach: Randomized SVD

**Objective:** Given an \( m \times n \) matrix \( A \), find an approximate rank-\( k \) partial SVD:

\[
A \approx U D V^* \\
\begin{array}{ccc}
m 	imes n & m \times k & k \times k \\
k \times n & &
\end{array}
\]

where \( U \) and \( V \) are orthonormal, and \( D \) is diagonal.

Fix an over-sampling parameter \( p \). Say \( p = 10 \).

(A) **Randomized pre-conditioner:**

A.1 Draw an \( n \times (k + p) \) Gaussian random matrix \( G \).

\[
G = \text{randn}(n, k+p)
\]

A.2 Form the \( m \times (k + p) \) sample matrix \( Y = AG \).

\[
Y = A \times G
\]

A.3 Form an \( m \times (k + p) \) orthonormal matrix \( Q \) such that \( Y = QR \).

\[
[Q, R] = \text{qr}(Y)
\]

(B) **Deterministic post-processing:**

B.1 Form the \((k + p) \times n\) matrix \( B = Q^* A \).

\[
B = Q' \times A
\]

B.2 Form SVD of the matrix \( B \): \( B = \hat{U} D V^* \).

\[
[\text{Uhat, Sigma, V}] = \text{svd}(B, \text{'econ'})
\]

B.3 Form the matrix \( U = Q \times \hat{U} \).

\[
U = Q \times \text{Uhat}
\]

(Truncate the last \( p \) terms in step B.2 to attain a factorization of precise rank \( k \).)
**Input:** An $m \times n$ matrix $A$, a target rank $k$, and an over-sampling parameter $p$ (say $p = 5$).

**Output:** Rank-$(k + p)$ factors $U$, $D$, and $V$ in an approximate SVD $A \approx UDV^*$.

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1. Draw an $n \times (k + p)$ random matrix $R$.
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3. Compute an ON matrix $Q$ s.t. $Y = QQ^*Y$.
4. Form the small matrix $B = Q^*A$.
5. Factor the small matrix $B = \hat{U}DV^*$.
6. Form $U = Q\hat{U}$.

- It is simple to adapt the scheme to the situation where the tolerance is given, and the rank has to be determined adaptively.

- Analogous schemes exist for computing a partial QR factorization, or a so called “interpolative decomposition” where a number of the columns/rows are chosen to serve as a basis for the column/row space.

  → Relaxed solution to “maximal spanning volume” problem on first slide.

- Accuracy of the basic scheme is good when the singular values decay reasonably fast. When they do not, the scheme can be combined with Krylov-type ideas:

  Taking one or two steps of subspace iteration vastly improves the accuracy.

  For instance, use the sampling matrix $Y = AA^*AG$ instead of $Y = AG$.

- We can reduce the flop count from $O(mnk)$ to $O(mn\log k)$ by using a so called “fast Johnson-Lindenstrauss” transform. Practical speed gain too!
**Input:** An $m \times n$ matrix $A$, a target rank $k$, and an over-sampling parameter $p$ (say $p = 5$).

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The output of RSVD is a random variable, as it depends on the draw of $R$. We have rigorous mathematical results describing the errors of the algorithm in expectation, as well as the risk of large deviations. Connections to random matrix theory.
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The perhaps most important feature of randomized algorithms is that they are very communication efficient. This makes them particularly competitive in strongly communication constrained environments (huge matrices stored out-of-core, distributed memory parallel computers, GPUs).

There exist *single-pass* versions of the RSVD that work even under the constraint that each matrix element can be viewed only once. (“Streaming algorithms.”)

Very recent result: Randomization can be used to greatly accelerate *full* rank-revealing factorizations such as the column pivoted QR factorization, or the UTV factorization. The gain is attained due to decreased communication, not fewer flops.
Speed-up of randomized column pivoted QR

**Speed-up attained by our randomized algorithm HQRRP** for computing a full column pivoted QR factorization of an $n \times n$ matrix. The speed-up is measured versus LAPACK's faster routine `dgeqp3` as implemented in Netlib (left) and Intel's MKL (right). Our implementation was done in C, and was executed on an Intel Xeon E5-2695. Joint work with G. Quintana-Ortí, N. Heavner, and R. van de Geijn. Available at: [https://github.com/flame/hqrrp/](https://github.com/flame/hqrrp/)
**Problem addressed:** The remainder of the talk concerns numerical methods for boundary value problems of the form

\[
\begin{aligned}
& A u(x) = g(x), \quad x \in \Omega, \\
& B u(x) = f(x), \quad x \in \Gamma,
\end{aligned}
\]

where \( \Omega \) is a domain with boundary \( \Gamma \), and where \( A \) is a linear elliptic differential operator (with possibly variable coefficients).

**Examples of problems we are interested in:**

- The equations of linear elasticity.
- Stokes’ equation.
- Helmholtz’ equation (at least at low and intermediate frequencies).
- Time-harmonic Maxwell (at least at low and intermediate frequencies).

**Archetypical example:** Poisson equation with Dirichlet boundary data:

\[
\begin{aligned}
& -\Delta u(x) = g(x), \quad x \in \Omega, \\
& u(x) = f(x), \quad x \in \Gamma,
\end{aligned}
\]

**Standard numerical recipe for (BVP):** (1) Discretize. (2) Solve linear system iteratively.
**Problem addressed:** The remainder of the talk concerns numerical methods for boundary value problems of the form

\[
\begin{cases}
Au(x) = g(x), & x \in \Omega, \\
Bu(x) = f(x), & x \in \Gamma,
\end{cases}
\]

where \( \Omega \) is a domain with boundary \( \Gamma \), and where \( A \) is a linear elliptic differential operator (with possibly variable coefficients).

**Observation:** The problem is *in principle* easy to solve! Simply integrate

\[
u(x) = \int_{\Omega} G(x, y) g(y) \, dy + \int_{\Gamma} F(x, y) f(y) \, dS(y), \quad x \in \Omega,
\]

where \( G \) and \( F \) are two kernel functions that depend on \( A, B, \) and \( \Omega \).

**Good:** The operators in (SLN) are friendly and nice.

\begin{center}
Bounded, smoothing, often fairly stable, etc.
\end{center}

**Problem:** The kernels \( G \) and \( F \) in (SLN) are generally *unknown.*

(Other than in trivial cases — constant coefficients and very simple domains.)

**Problem:** The operators in (SLN) are *global.*

\begin{center}
Dense matrices upon discretization. \( O(N^2) \) cost?
\end{center}
Greengard, Rokhlin (1985): The solution operator for the Poisson equation can be applied in $O(N)$ operations. “Fast Multipole Method.”


Hackbusch et al (1998): Explicit recipe for building the operators in $O(n \log^r n)$ operators for $r$ moderate. “$\mathcal{H}$-matrices.”
Recall that we are interested in solving the PDE
\[
\begin{align*}
Au(x) &= g(x), \quad x \in \Omega, \\
Bu(x) &= f(x), \quad x \in \Gamma.
\end{align*}
\] (BVP)

Explicit solution formula: 
\[
\begin{align*}
\mathbf{u}(\mathbf{x}) &= \int_{\Omega} G(\mathbf{x}, \mathbf{y}) g(\mathbf{y}) \, d\mathbf{y} + \int_{\Gamma} F(\mathbf{x}, \mathbf{y}) f(\mathbf{y}) \, dS(\mathbf{y}), \\
&\quad \mathbf{x} \in \Omega.
\end{align*}
\] (SLN)

**Recurring idea:** Upon discretization, (SLN) leads to a matrix with **off-diagonal blocks of low numerical rank.**

This property can be exploited to attain linear or close to linear complexity for operations such as matrix-vector multiply, matrix-matrix multiply, LU factorization, matrix inversion, forming of Schur complements, etc.

**Strong connections to Calderón-Zygmund theory for singular integral operators.**

**References:** Fast Multipole Method (Greengard, Rokhlin); Panel Clustering (Hackbusch); H- and H-matrices; Hierarchically Block Separable (HBS) matrices; Hierarchically Semi Separable (HSS) matrices; S-matrices, a.k.a. HODLR matrices;...
Recall that we are interested in solving the PDE
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\] (SLN)

**Question:** Why do the dense matrices resulting upon discretization of (SLN) typically have *off-diagonal blocks of low numerical rank*?

**(One) Answer:** It is a consequence of the *smoothing effect* of elliptic differential equations; it can be interpreted as a *loss of information*.

This effect has many well known physical consequences:

- Rapid convergence of *multipole expansions* when the region of sources is far away from the observation point.
- The *St Venant principle* in mechanics.
- The inaccuracy of imaging at sub-wavelength scales.
- The intractability of solving the heat equation backwards.

**Caveat:** High-frequency problems present difficulties — no loss of information for length-scales $> \lambda$. Extreme accuracy of optics, high-frequency imaging, *etc.*
A 1D model problem: Consider a simple 2-point BVP on the interval $[0, 1]$:

\[
-\frac{d^2 u}{dx^2} + p(x) \frac{du(x)}{dx} + q(x) u(x) = g(x), \quad x \in (0, 1),
\]

\[
\begin{align*}
  u(0) &= f_L, \\
  u(1) &= f_R.
\end{align*}
\]

Discretizing (BVP) using a standard second order finite difference scheme, we get

\[
Au = b,
\]

where $A$ is a sparse matrix of size, say, $n \times n$. Then $A^{-1}$ is dense.
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Sparsity pattern of $A$.

A is tridiagonal.

Sparsity pattern of $A^{-1}$.
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- **A** is tridiagonal.
- **A** is sparse.
- $A^{-1}$ is semi-separable.
A 1D model problem: Consider a simple 2-point BVP on the interval [0, 1]:

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Discretizing (BVP) using a standard second order finite difference scheme, we get

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Sparsity pattern of \(A\). \(A\) is tridiagonal. 

Sparsity pattern of \(A^{-1}\). \(A^{-1}\) is semi-separable. 

\(A\) is sparse. 

\(A^{-1}\) is data-sparse.
• Consider an elliptic PDE

\[
\begin{cases}
Au(x) = g(x), & x \in \Omega, \\
Bu(x) = f(x), & x \in \Gamma,
\end{cases}
\]

where \( \Omega \) is a domain in \( \mathbb{R}^2 \) or \( \mathbb{R}^3 \) with boundary \( \Gamma \).

• Discretize (BVP) using FEM / FD / \ldots to obtain a linear system

\[ Au = b. \]

The matrix \( A \) will be sparse.

• Given a computational tolerance \( \varepsilon \), we now seek a direct (that is, non-iterative) algorithm that builds a matrix \( S \) such that

\[ \| S - A^{-1} \| \leq \varepsilon. \]

The matrix \( S \) will be dense, but “data-sparse.”
A 2D model problem: Let $\Omega = [0, 1]^2$ and $\Gamma = \partial \Omega$. We seek to solve

\begin{align*}
\begin{cases}
-\Delta u(x) = g(x), & x \in \Omega, \\
u(x) = f(x), & x \in \Gamma.
\end{cases}
\end{align*}

We introduce an $n \times n$ grid on $\Omega$ with nodes $\{x_j\}_{j=1}^N$ where $N = n^2$, see Figure A. Letting $u = [u(j)]_{j=1}^N$ denote a vector of approximate solution values, $u(j) \approx u(x_j)$, and using the standard five-point stencil to discretize $-\Delta$, we end up with a sparse linear system

$$Au = b,$$

where $[Au](k) = \frac{1}{h^2}(4u(k) - u(k_s) - u(k_e) - u(k_n) - u(k_w))$, see Figure B.
Divide-and-conquer: Split the nodes in three groups as shown so that there are no connections between nodes in $\Omega_1$ and $\Omega_2$. Then $A$ has zero blocks as shown:

$$
\begin{array}{ccc}
\Omega_1 & \Omega_2 & \Omega_3 \\
\end{array}
$$

$$
A = \begin{bmatrix}
A_{11} & 0 & A_{13} \\
0 & A_{22} & A_{23} \\
A_{31} & A_{32} & A_{33}
\end{bmatrix}
$$

Notice the obvious recursion!
**Divide-and-conquer:** Split the nodes in three groups as shown so that there are no connections between nodes in $\Omega_1$ and $\Omega_2$. Then $A$ has zero blocks as shown:

$$A = \begin{bmatrix} A_{11} & 0 & A_{13} \\ 0 & A_{22} & A_{23} \\ A_{31} & A_{32} & A_{33} \end{bmatrix}$$

Now suppose that we can somehow construct $A_{11}^{-1}$ and $A_{22}^{-1}$. Then

$$A = \begin{bmatrix} I & 0 & 0 \\ 0 & I & 0 \\ A_{31}A_{11}^{-1} & A_{32}A_{22}^{-1} & I \end{bmatrix} \begin{bmatrix} A_{11} & 0 & 0 \\ 0 & A_{22} & 0 \\ 0 & 0 & S_{33} \end{bmatrix} \begin{bmatrix} I & 0 & A_{11}^{-1}A_{13} \\ 0 & I & A_{22}^{-1}A_{23} \\ 0 & 0 & I \end{bmatrix}$$

where $S_{33} = A_{33} - A_{31}A_{11}^{-1}A_{13} - A_{32}A_{22}^{-1}A_{23}$ is a Schur complement.
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In other words, in order to invert $A$, we need to execute three steps:

- Invert $A_{11}$ to form $A_{11}^{-1}$.  \( \text{size} \sim N/2 \times N/2 \)
- Invert $A_{22}$ to form $A_{22}^{-1}$.  \( \text{size} \sim N/2 \times N/2 \)
- Invert $S_{33} = A_{33} - A_{31}A_{11}^{-1}A_{13} - A_{32}A_{22}^{-1}A_{23}$.  \( \text{size} \sim \sqrt{N} \times \sqrt{N} \)

Notice the obvious recursion!
**Divide-and-conquer:** Split the nodes in three groups as shown so that there are no connections between nodes in \( \Omega_1 \) and \( \Omega_2 \). Then \( A \) has zero blocks as shown:

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A_{31} & A_{32} & A_{33}
\end{bmatrix}
\]

Now suppose that we can somehow construct \( A_{11}^{-1} \) and \( A_{22}^{-1} \). Then

\[
A = \begin{bmatrix}
I & 0 & 0 \\
0 & I & 0 \\
A_{31}A_{11}^{-1} & A_{32}A_{22}^{-1} & I
\end{bmatrix}
\begin{bmatrix}
A_{11} & 0 & 0 \\
0 & A_{22} & 0 \\
0 & 0 & S_{33}
\end{bmatrix}
\begin{bmatrix}
I & 0 & A_{11}^{-1}A_{13} \\
0 & I & A_{22}^{-1}A_{23} \\
0 & 0 & I
\end{bmatrix}
\]

where \( S_{33} = A_{33} - A_{31}A_{11}^{-1}A_{13} - A_{32}A_{22}^{-1}A_{23} \) is a **Schur complement**.

In other words, in order to invert \( A \), we need to execute three steps:

- Invert \( A_{11} \) to form \( A_{11}^{-1} \). \( \text{size} \sim N/2 \times N/2 \)
- Invert \( A_{22} \) to form \( A_{22}^{-1} \). \( \text{size} \sim N/2 \times N/2 \)
- Invert \( S_{33} = A_{33} - A_{31}A_{11}^{-1}A_{13} - A_{32}A_{22}^{-1}A_{23} \). \( \text{size} \sim \sqrt{N} \times \sqrt{N} \)

Notice the obvious recursion!
Divide-and-conquer: Split the nodes in three groups as shown so that there are no connections between nodes in $\Omega_1$ and $\Omega_2$. Then $A$ has zero blocks as shown:

\[
A = \begin{bmatrix}
A_{11} & 0 & A_{13} \\
0 & A_{22} & A_{23} \\
A_{31} & A_{32} & A_{33}
\end{bmatrix}
\]

Now suppose that we can somehow factor $A_{11} = L_{11}U_{11}$ and $A_{22} = L_{22}U_{22}$. Then

\[
A = \begin{bmatrix}
L_{11} & 0 & A_{13} \\
0 & L_{22} & A_{23} \\
A_{31}U_{11}^{-1} & A_{32}U_{22}^{-1} & I
\end{bmatrix}
\begin{bmatrix}
I & 0 & 0 \\
0 & I & 0 \\
0 & 0 & S_{33}
\end{bmatrix}
\begin{bmatrix}
U_{11} & 0 & L_{11}^{-1}A_{13} \\
0 & U_{22} & L_{22}^{-1}A_{23} \\
0 & 0 & I
\end{bmatrix}
\]

where $S_{33} = A_{33} - A_{31}U_{11}^{-1}L_{11}^{-1}A_{13} - A_{32}U_{22}^{-1}L_{22}^{-1}A_{23}$ is a Schur complement.

In other words, in order to invert $A$, we need to execute three steps:

- Factor $A_{11}$ to form $A_{11} = L_{11}U_{11}$. \hspace{1cm} \text{size} \sim N/2 \times N/2
- Factor $A_{22}$ to form $A_{22} = L_{22}U_{22}$. \hspace{1cm} \text{size} \sim N/2 \times N/2
- Factor $S_{33} = A_{33} - A_{31}U_{11}^{-1}L_{11}^{-1}A_{13} - A_{32}U_{22}^{-1}L_{22}^{-1}A_{23}$. \hspace{1cm} \text{size} \sim \sqrt{N} \times \sqrt{N}

Notice the obvious recursion!
Upwards pass — build all solution operators:


2. Downwards pass — solve for a particular data function (very fast!):

Upwards pass — build all solution operators:

The original grid. Leaves reduced. After merge. After merge.

Downwards pass — solve for a particular data function (very fast!):


Alan George  Iain Duff  Tim Davis
The direct solver described works very well for moderate problem sizes. But problems arise as the number of discretization points increases . . .
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Consider a regular grid in 2D with $N = n \times n$ total nodes. The top level merge requires inversion of a matrix representing interactions between the red nodes:

![Diagram of a grid with red nodes](image)

Since this dense matrix is of size $n \times n$, the cost for the merge is

$$\text{COST} \sim n^3 \sim (N^{1/2})^3 \sim N^{3/2}.$$
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**Problem:** 3D is much worse!
The direct solver described works very well for moderate problem sizes. But problems arise as the number of discretization points increases . . .

Consider a regular grid in 3D with $N = n \times n \times n$ total nodes. The top level merge requires inversion of a matrix representing interactions between the red nodes:

The merge requires factorization of a dense matrix of size $n^2 \times n^2$. Consequently:

$$\text{COST} \sim (N^{1/3})^6 \sim N^2.$$
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Consider a regular grid in 3D with $N = n \times n \times n$ total nodes. The top level merge requires inversion of a matrix representing interactions between the red nodes:

![Diagram of a 3D grid](image)

The merge requires factorization of a dense matrix of size $n^2 \times n^2$. Consequently:

$$\text{COST} \sim (N^{1/3})^6 \sim N^2.$$  

**Assertion:** The dense matrix very often behaves like a discretized integral operator. (E.g. Dirichlet-to-Neumann.)

It is rank-structured, and is amenable to “fast” matrix algebra.

We can reduce the complexity of the top level solve from $O(N^2)$ down to $O(N)$, and sometimes even $O(N^{2/3})$. 
Exploiting the assertion on the previous page, we have in the last 10 years managed to reduce the asymptotic complexity of direct solvers for elliptic PDEs dramatically:

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<thead>
<tr>
<th>Build stage</th>
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**Key idea:** Represent dense matrices using rank-structured formats (such as $\mathcal{H}$-matrices).


*O(N) direct solvers for integral equations* were developed by Martinsson & Rokhlin (2005), Greengard, Gueyffier, Martinsson, & Rokhlin (2009), Gillman, Young, & Martinsson (2012), Ho & Greengard (2012), Ho & Ying (2015). Work in 1990’s Y. Chen, P. Starr, V. Rokhlin, L. Greengard, E. Michielssen. Related to work on $\mathcal{H}$ and $\mathcal{H}^2$ matrix methods (1998 and forwards) by Bebendorf, Börm, Hackbusch, Khoromskij, Sauter, etc.

**Note:** Complexity is not $O(N)$ if the nr. of “points-per-wavelength” is fixed as $N \to \infty$. This limits direct solvers to problems of size a couple hundreds of wave-lengths or so.
Key selling point: Better parallelism

Let us consider the flop counts of various parts of the computation:

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**Observations:**

- While the dominant cost of the old scheme is processing dense matrices of size $O(N^{2/3}) \times O(N^{2/3})$, the dominant cost of the new scheme is processing the leaves.
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Observations:

- While the dominant cost of the old scheme is processing dense matrices of size $O(N^{2/3}) \times O(N^{2/3})$, the dominant cost of the new scheme is processing the leaves.

  *The leaf computations are very easy to parallelize!*

- Parallel implementations of structured matrix algebra require hard work (J. Poulson’s dissertation; S. Li at LBNL; G. Biros; R. Kriemann; P. Amestoy & A. Buttari; G. Turkiyyah & D. Keyes; J. Xia; etc).

- For intermediate size problems, the structured matrices of size $O(N^{2/3}) \times O(N^{2/3})$ often fit on one machine.

- The methodology need not be all-or-nothing. Direct solvers can be used locally to handle areas with mesh refinement etc.
Claim: Direct solvers are ideal for combining with high order discretization.

- Direct solvers use a lot of memory per degree of freedom.
  → You want to maximize the oomph per DOF.

- Direct solvers are particularly well suited for “high” frequency wave problems.
  → Need high accuracy due to ill-conditioned physics.

- High order methods sometimes lead to more ill-conditioned systems.
  → Can be hard to get iterative solvers to converge.

Problem: If you combine “nested dissection” with traditional discretization techniques (FD, FEM, etc), then the performance plummets as the order is increased.

Solution: Derive a new (or at least newish) discretization scheme that is directly tailored to work with fast direct solvers.
The Hierarchical Poincaré-Steklov Method

A direct solver based on a multidomain spectral collocation discretization

For simplicity, let us consider a “variable wave speed” Helmholtz problem in 2D: Given $f$, $g$, and $b$, find $u$ such that

\[
\begin{align*}
-\Delta u(x) - b(x) u(x) &= g(x), & x &\in \Omega, \\
u(x) &= f(x), & x &\in \Gamma,
\end{align*}
\]

where $\Omega = [0, 1]^2$ is the unit square and $\Gamma = \partial \Omega$.

We assume $u$ is smooth.

The unknown function $u$ is represented as a vector holding approximations to its point-wise values at the grid points (collocation). Across domain boundaries, we enforce continuity of potentials and normal derivatives.

A global solution operator will be built using a nested-dissection type solver.

Prior work: The discretization scheme is similar to existing composite (or “multi-domain”) spectral collocation methods by Hesthaven and others. In particular: Pfeiffer, Kidder, Scheel, Teukolsky, (2003). Connections to domain decomposition and “reduction to interface” methods (Khoromskij & Wittum, etc.).
The Hierarchical Poincaré-Steklov Method

Model problem: Given $f$, $g$, and $b$, find $u$ such that

\[
\begin{cases}
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where $\Omega = [0, 1]^2$ is the unit square and $\Gamma = \partial \Omega$. We assume $u$ is smooth.

Process leaves: Eliminate the interior (blue) nodes. ("Static condensation.") Technically, we compute the Dirichlet-to-Neumann operator via a local spectral computation.
Model problem: Given \( f, g, \) and \( b, \) find \( u \) such that
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Upwards sweep: Merge boxes by pairs and eliminate the interior (blue) nodes. To do this, use the computed DtN operators to enforce continuity of $u$ and $du/dn$ across interior boundaries. Compute the DtN operator for the larger box.
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Top level solve: Invert the DtN operator for the top level box.
The Hierarchical Poincaré-Steklov Method

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\]

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Downwards sweep: We know $u$ on the red nodes. We can use the computed DtN operators to reconstruct $u$ on the blue nodes.
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Hierarchical Poincaré-Steklov Method:

- In contrast to prior schemes, the speed of the solver does not deteriorate as the order is increased.

- Very high order can be used (say $20 \times 20$ local mesh).
  
  → Overall errors close to machine precision for problems with smooth solutions.

- Capable of solving 2D problems on domains that are several hundred wave-lengths across in minutes on a laptop.

- Exploiting internal structure in the boundary-to-boundary operators, we can attain $O(N)$ complexity and handle $N \sim 10^8$ on a desktop. (For 2D problems; 3D is harder.)

- Extension to 3D is under way. (Joint work with A. Gillman, G. Turkiyyah, D. Keyes.)
Hierarchical Poincaré-Steklov Method — rank deficiencies in the DtN operator:

Recall that at the top level, we need to invert a dense matrix that is defined on the nodes of the interface highlighted in red and blue below. This matrix holds restrictions of the Dirichlet-to-Neumann (DtN) operators for the two blocks. We have claimed that this matrix is rank-structured. *But what are the ranks?*

Let $T$ denote the restriction of the DtN matrix mapping Dirichlet data on $\Gamma_1$ to Neumann data on $\Gamma_2$ for a 1089 $\times$ 1089 grid. Then $T$ is of size 512 $\times$ 512.

![Diagram of the interface with nodes highlighted in red and blue](image-url)
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![Singular values of $\mathbf{T}$.](image)

**Laplace problem.**

**Rank=22.**
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**Singular values of $\mathbf{T}$.**

Helmholtz problem.
Domain is $110\lambda \times 110\lambda$.
Rank=85.
Hierarchical Poincaré-Steklov Method: numerical results

Set $\Omega = [0, 1]^2$ and $\Gamma = \partial \Omega$. Consider the problem

$$
\begin{align*}
-\Delta u(x) - \kappa^2 u(x) &= 0, \quad x \in \Omega, \\
u(x) &= f(x), \quad x \in \Gamma.
\end{align*}
$$

We pick $f$ as the restriction of a wave from a point source, $x \mapsto Y_0(\kappa |x - \hat{x}|)$. We then know the exact solution, $u_{\text{exact}}(x) = Y_0(\kappa |x - \hat{x}|)$. 

Approximate solution. ntot=1681 pts-per-wave=12.00
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The spectral computation on a leaf involves $21 \times 21$ points.

$\kappa$ is chosen so that there are 12 points per wave-length.

<table>
<thead>
<tr>
<th>$p$</th>
<th>$N$</th>
<th>$N_{\text{wave}}$</th>
<th>$t_{\text{build}}$ (sec)</th>
<th>$t_{\text{solve}}$ (sec)</th>
<th>$E_{\text{pot}}$</th>
<th>$E_{\text{grad}}$</th>
<th>$M$ (MB)</th>
<th>$M/N$ (reals/DOF)</th>
</tr>
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<tbody>
<tr>
<td>21</td>
<td>6561</td>
<td>6.7</td>
<td>0.23</td>
<td>0.0011</td>
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<td>21</td>
<td>1640961</td>
<td>106.7</td>
<td>99.20</td>
<td>0.2965</td>
<td>1.90502e-09</td>
<td>1.24859e-06</td>
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<td>117.2</td>
</tr>
<tr>
<td>21</td>
<td>6558721</td>
<td>213.3</td>
<td>551.32</td>
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<td>3.74616e-06</td>
<td>6218.7</td>
<td>124.3</td>
</tr>
</tbody>
</table>

Error is measured in sup-norm: $e = \max_{x \in \Omega} |u(x) - u_{\text{exact}}(x)|$.

**Note 1:** Translation invariance is *not* exploited.

**Note 2:** The times refer to a simple Matlab implementation executed on a $1k$ laptop.

**Note 3:** Keeping a fixed number of points per wave-length works well for this scheme!
Hierarchical Poincaré-Steklov Method: numerical results

Set $\Omega = [0, 1]^2$ and $\Gamma = \partial \Omega$. Consider the problem

\[
\begin{aligned}
-\Delta u(x) - \kappa^2 u(x) &= 0, & x &\in \Omega, \\
 u(x) &= f(x), & x &\in \Gamma.
\end{aligned}
\]

We pick $f$ as the restriction of a wave from a point source, $x \mapsto Y_0(\kappa|x - \hat{x}|)$.

We then know the exact solution, $u_{\text{exact}}(x) = Y_0(\kappa|x - \hat{x}|)$.

The spectral computation on a leaf involves $41 \times 41$ points.

$\kappa$ is chosen so that there are 12 points per wave-length.

<table>
<thead>
<tr>
<th>$p$</th>
<th>$N$</th>
<th>$N_{\text{wave}}$</th>
<th>$t_{\text{build}}$ (sec)</th>
<th>$t_{\text{solve}}$ (sec)</th>
<th>$E_{\text{pot}}$</th>
<th>$E_{\text{grad}}$</th>
<th>$M$ (MB)</th>
<th>$M/N$ (reals/DOF)</th>
</tr>
</thead>
<tbody>
<tr>
<td>41</td>
<td>6561</td>
<td>6.7</td>
<td>1.50</td>
<td>0.0025</td>
<td>9.88931e-14</td>
<td>3.46762e-12</td>
<td>7.9</td>
<td>157.5</td>
</tr>
<tr>
<td>41</td>
<td>25921</td>
<td>13.3</td>
<td>4.81</td>
<td>0.0041</td>
<td>1.58873e-13</td>
<td>1.12883e-11</td>
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<td>166.4</td>
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<tr>
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<td>103041</td>
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<td>18.34</td>
<td>0.0162</td>
<td>3.95531e-13</td>
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<tr>
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</table>

Error is measured in sup-norm: $e = \max_{x \in \Omega} |u(x) - u_{\text{exact}}(x)|$.

**Note 1:** Translation invariance is \textit{not} exploited.

**Note 2:** The times refer to a simple Matlab implementation executed on a $1k$ laptop.

**Note 3:** Keeping a fixed number of points per wave-length works well for this scheme!
The line $t_{\text{solve}}$ scales perfectly linearly (until memory problems kick in), as expected.

**Interesting:** The line $t_{\text{build}}$ also scales almost linearly. (Unexpectedly?) It turns out that $t_{\text{build}}$ is dominated by the leaf computation; we have not yet hit the $O(N^{1.5})$ asymptotic.
Hierarchical Poincaré-Steklov Method: numerical results — variable coefficients

Now consider the variable coefficient problem

\[-\Delta u(x) - \kappa^2 \left( 1 - b(x) \right) u(x) = 0 \quad x \in \Omega,\]

\[u(x) = f(x) \quad x \in \Gamma,\]

where \( \Omega = [0, 1]^2 \), where \( \Gamma = \partial \Omega \), and where \( b(x) = \left( \sin(4\pi x_1) \sin(4\pi x_2) \right)^2 \).

The Helmholtz parameter was kept fixed at \( \kappa = 80 \), corresponding to a domain size of \( 12.7 \times 12.7 \) wave lengths. The boundary data was given by \( f(x) = \cos(8x_1) \left( 1 - 2x_2 \right) \).

The error estimator \( E^\text{int}_N = u_N(\hat{x}) - u_{4N}(\hat{x}) \) where \( \hat{x} = (0.75, 0.25) \) is reported below:

<table>
<thead>
<tr>
<th>( p )</th>
<th>( N )</th>
<th>pts per wave</th>
<th>( u_N(\hat{x}) )</th>
<th>( E^\text{int}_N )</th>
<th>( w_N(\hat{y}) )</th>
<th>( E^\text{bnd}_N )</th>
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<tr>
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<tr>
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<td>-5.463e-06</td>
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<tr>
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<td>-2.792e-05</td>
</tr>
<tr>
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<td>-33231.6177808723</td>
<td>-4.668e-05</td>
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</tr>
<tr>
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<td>-33231.6178008533</td>
<td>-4.668e-05</td>
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<td>-33231.6177541722</td>
<td>-2.792e-05</td>
<td></td>
</tr>
</tbody>
</table>
Consider a curved domain $\Psi$ as shown above and the equation

\begin{align*}
\begin{cases}
-\Delta u(y) - \kappa^2 u(y) &= 0 & y \in \Psi, \\
u(y) &= f(y) & y \in \partial \Psi.
\end{cases}
\end{align*}

The reparameterization is $y_1 = x_1$ and $y_2 = \psi(y_1) y_2$, and so the Helmholtz equation (9) takes the form

\[
\frac{\partial^2 u}{\partial x_1^2} + \frac{2\psi'(x_1)}{\psi(x_1)} \frac{x_2}{\partial x_1} \frac{\partial^2 u}{\partial x_2} + \left( \frac{x_2^2 \psi'(x_1)^2}{\psi(x_1)^2} + \psi(x_1)^2 \right) \frac{\partial^2 u}{\partial x_2^2} + \frac{x_2 \psi''(x_1)}{\psi(x_1)} \frac{\partial u}{\partial x_2} + k^2 u = 0, \quad x \in \Omega.
\]
Numerical results for curved domain

The equation is (constant coefficient) Helmholtz on a domain of size $35 \times 50$ wavelengths.

<table>
<thead>
<tr>
<th>$N$</th>
<th>$E_{\text{pot}}$</th>
<th>$E_{\text{grad}}$</th>
<th>$E_{N}^{(1)}$</th>
<th>$E_{N}^{(2)}$</th>
<th>$E_{N}^{(3)}$</th>
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<tbody>
<tr>
<td>25921</td>
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<td>2.26204e-07</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>
Hierarchical Poincaré-Steklov Method: “FEM-BEM coupling”

Consider the free space acoustic scattering problem

\[
\begin{cases}
-\Delta u(x) - \kappa^2 (1 - b(x)) u(x) = -\kappa^2 b(x) v(x), & x \in \mathbb{R}^2 \\
\lim_{|x| \to \infty} \sqrt{|x|} (\partial_x u(x) - i\kappa u(x)) = 0,
\end{cases}
\]

where

- \( b \) is a smooth scattering potential with compact support, where
- \( v \) is a given “incoming potential” and where
- \( u \) is the sought “outgoing potential.”

Joint work with A. Barnett and A. Gillman.
Hierarchical Poincaré-Steklov Method: “FEM-BEM coupling”

Consider the free space acoustic scattering problem

\[
\begin{cases}
-\Delta u(x) - \kappa^2 (1 - b(x)) u(x) = -\kappa^2 b(x) v(x), & x \in \mathbb{R}^2 \\
\lim_{|x|\to\infty} \sqrt{|x|} (\partial_x u(x)) = 0,
\end{cases}
\]

where

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- \( v \) is a given “incoming potential” and where
- \( u \) is the sought “outgoing potential.”

Introduce an artificial box \( \Omega \) such that \( \text{support}(b) \subseteq \Omega \).

\( -\Delta u - \kappa^2 (1 - b) u = -\kappa^2 b v \)

\( \text{support}(b) \)

\( -\Delta u - \kappa^2 u = 0 \) on \( \partial \mathbb{R}^2 \)

Joint work with A. Barnett and A. Gillman.
Hierarchical Poincaré-Steklov Method: “FEM-BEM coupling”

Consider the free space acoustic scattering problem

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\begin{cases}
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\lim_{|x| \to \infty} \sqrt{|x|} (\partial_{|x|} u(x) - i\kappa u(x)) = 0,
\end{cases}
\]

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On \(\Omega\):
- Variable coefficient PDE.

On \(\Omega^c\):
- Constant coefficient PDE.

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Hierarchical Poincaré-Steklov Method: “FEM-BEM coupling”

Consider the free space acoustic scattering problem

\[
\begin{align*}
-\Delta u(x) - \kappa^2 (1 - b(x)) u(x) &= - \kappa^2 b(x) v(x), \quad x \in \mathbb{R}^2 \\
\lim_{|x| \to \infty} \sqrt{|x|} (\partial_x u(x) - i\kappa u(x)) &= 0,
\end{align*}
\]

where

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On \( \Omega \):
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- Use HPS.

On \( \Omega^c \):
- Constant coefficient PDE.
- Use BIE.

Joint work with A. Barnett and A. Gillman.
Hierarchical Poincaré-Steklov Method: “FEM-BEM coupling”

Consider the free space acoustic scattering problem

\[
\begin{cases}
-\Delta u(x) - \kappa^2 (1 - b(x)) u(x) = -\kappa^2 b(x) v(x), & x \in \mathbb{R}^2 \\
\lim_{|x| \to \infty} \sqrt{|x|} ((\partial_x u(x) - i\kappa u(x)) = 0,
\end{cases}
\]

where

- \(b\) is a smooth scattering potential with compact support, where
- \(v\) is a given “incoming potential” and where
- \(u\) is the sought “outgoing potential.”

Introduce an artificial box \(\Omega\) such that support\((b) \subseteq \Omega\).

\[
\begin{align*}
-\Delta u - \kappa^2 (1-b) u &= -\kappa^2 b v \\
\text{support}(b) \quad \Omega
\end{align*}
\]

On \(\Omega\):
- Variable coefficient PDE.
- Use HPS.
- Build DtN for \(\partial\Omega\).

On \(\Omega^c\):
- Constant coefficient PDE.
- Use BIE.
- Build DtN for \(\partial\Omega^c\).

Joint work with A. Barnett and A. Gillman.
Hierarchical Poincaré-Steklov Method: “FEM-BEM coupling”

Consider the free space acoustic scattering problem

\[
\begin{cases}
-\Delta u(x) - \kappa^2 (1 - b(x)) u(x) = -\kappa^2 b(x) v(x), & x \in \mathbb{R}^2 \\
\lim_{|x| \to \infty} \sqrt{|x|} (\partial_x u(x) - i\kappa u(x)) = 0,
\end{cases}
\]

where

- \(b\) is a smooth scattering potential with compact support, where
- \(v\) is a given “incoming potential” and where
- \(u\) is the sought “outgoing potential.”

Introduce an artificial box \(\Omega\) such that \(\text{support}(b) \subseteq \Omega\).

On \(\Omega\):
- Variable coefficient PDE.
- Use HPS.
- Build DtN for \(\partial\Omega\).

On \(\Omega^c\):
- Constant coefficient PDE.
- Use BIE.
- Build DtN for \(\partial\Omega^c\).

\[ -\Delta u - \kappa^2 (1-b) u = -\kappa^2 b v \]

\[ -\Delta u - \kappa^2 u = 0 \text{ on } \Omega^c \]

\[ \text{\textbullet Merge using fast operator algebra!} \]

Joint work with A. Barnett and A. Gillman.
The scattering potential $b$
Example: Free space scattering

\[
\begin{aligned}
-\Delta u_{\text{out}}(\mathbf{x}) - \kappa^2 (1 - b(\mathbf{x})) u_{\text{out}}(\mathbf{x}) &= -\kappa^2 b(\mathbf{x}) u_{\text{in}}(\mathbf{x}) \\
\lim_{|\mathbf{x}| \to \infty} \sqrt{|\mathbf{x}|} (\partial_{|\mathbf{x}|} u_{\text{out}}(\mathbf{x}) - i\kappa u_{\text{out}}(\mathbf{x})) &= 0
\end{aligned}
\]

The outgoing field \( u_{\text{out}} \) (resulting from an incoming plane wave \( u_{\text{in}}(\mathbf{x}) = \cos(\kappa x_1) \))

\[ N = 231361 \quad T_{\text{build}} = 7.2 \text{ sec} \quad T_{\text{solve}} = 0.06 \text{ sec} \quad E \approx 10^{-7} \text{ (estimated)} \]
Example: Free space scattering

\[
\begin{align*}
-\Delta u_{\text{out}}(x) - \kappa^2 (1 - b(x)) u_{\text{out}}(x) &= -\kappa^2 b(x) u_{\text{in}}(x) \\
\lim_{|x| \to \infty} \sqrt{|x|} (\partial_{|x|} u_{\text{out}}(x) - i\kappa u_{\text{out}}(x)) &= 0
\end{align*}
\]

The outgoing field \( u_{\text{out}} \) (resulting from an incoming plane wave \( u_{\text{in}}(x) = \cos(\kappa x_1) \))

\[
N = 231\,361 \quad T_{\text{build}} = 7.2 \text{ sec} \quad T_{\text{solve}} = 0.06 \text{ sec} \quad E \approx 10^{-7} \text{ (estimated)}
\]
Example: Free space scattering

\[
\begin{cases}
-\Delta u_{\text{out}}(x) - \kappa^2 (1 - b(x)) u_{\text{out}}(x) = -\kappa^2 b(x) u_{\text{in}}(x) \\
\lim_{|x| \to \infty} \sqrt{|x|} (\frac{\partial}{\partial |x|} u_{\text{out}}(x) - i\kappa u_{\text{out}}(x)) = 0
\end{cases}
\]

*The scattering potential $b$ — now a photonic crystal with a wave guide.*

\[ N = 231361 \quad T_{\text{build}} = 7.2 \text{ sec} \quad T_{\text{solve}} = 0.06 \text{ sec} \quad E \approx 10^{-6} \text{ (estimated)} \]
Example: Free space scattering

\[
\begin{aligned}
-\Delta u_{\text{out}}(x) - \kappa^2 (1 - b(x)) u_{\text{out}}(x) &= -\kappa^2 b(x) u_{\text{in}}(x) \\
\lim_{|x| \to \infty} \sqrt{|x|} (\partial_x |u_{\text{out}}(x)| - i\kappa u_{\text{out}}(x)) &= 0
\end{aligned}
\]

The total field \( u = u_{\text{in}} + u_{\text{out}} \) (resulting from an incoming plane wave \( u_{\text{in}}(x) = \cos(\kappa x_1) \)).
Recall: The method as presented relies on a hierarchical construction of Dirichlet-to-Neumann operators for every box in a hierarchical tree.

Problem! The interior Helmholtz equation may encounter resonances — even for zero Dirichlet data, there may be non-trivial solutions.

Conceptual problem: The DtN operator does not always exist.

Practical problem: The numerical DtN operator can be very ill-conditioned.

Joint work with Alexander Barnett (Dartmouth) and Adrianna Gillman (Rice).
Recall: The method as presented relies on a hierarchical construction of Dirichlet-to-Neumann operators for every box in a hierarchical tree.

Problem! The interior Helmholtz equation may encounter resonances — even for zero Dirichlet data, there may be non-trivial solutions.

Conceptual problem: The DtN operator does not always exist.

Practical problem: The numerical DtN operator can be very ill-conditioned.

Solution: Rather than the \textit{Dirichlet-to-Neumann map}\n
$$ T : \ u|_{\Gamma} \mapsto \frac{\partial u}{\partial n}|_{\Gamma} $$

consider the \textit{impedance map}\n
$$ E : \ u|_{\Gamma} + i \frac{\partial u}{\partial n}|_{\Gamma} \mapsto u|_{\Gamma} - i \frac{\partial u}{\partial n}|_{\Gamma} $$

The impedance map exists for every wave-number, and is a unitary map.

Joint work with Alexander Barnett (Dartmouth) and Adrianna Gillman (Rice).
The build stage can be accelerated to optimal $O(N)$ complexity:

Consider the merge of two patches $\Omega^{(\alpha)}$ and $\Omega^{(\beta)}$ with boundaries $\Gamma_1$, $\Gamma_2$, $\Gamma_3$:

In the composite spectral method we have

$$T = \begin{bmatrix} T_{1,1}^{(\alpha)} & 0 \\ 0 & T_{2,2}^{(\beta)} \end{bmatrix} + \begin{bmatrix} T_{1,3}^{(\alpha)} \\ T_{2,3}^{(\beta)} \end{bmatrix} (T_{3,3}^{(\alpha)} - T_{3,3}^{(\beta)})^{-1} \begin{bmatrix} -T_{3,1}^{(\alpha)} | T_{3,2}^{(\beta)} \end{bmatrix}.$$

low rank update!
The build stage can be accelerated to optimal $O(N)$ complexity:

Consider the merge of two patches $\Omega^{(\alpha)}$ and $\Omega^{(\beta)}$ with boundaries $\Gamma_1, \Gamma_2, \Gamma_3$:

In the composite spectral method we have

$$T = \begin{bmatrix} T^{(\alpha)}_{1,1} & 0 \\ 0 & T^{(\beta)}_{2,2} \end{bmatrix} + \begin{bmatrix} T^{(\alpha)}_{1,3} \\ T^{(\beta)}_{2,3} \end{bmatrix} \left( T^{(\alpha)}_{3,3} - T^{(\beta)}_{3,3} \right)^{-1} \begin{bmatrix} -T^{(\alpha)}_{3,1} \\ T^{(\beta)}_{3,2} \end{bmatrix}.$$ 

low rank update!

There is more structure!
The build stage can be accelerated to optimal $O(N)$ complexity:

Consider the merge of two patches $\Omega^{(\alpha)}$ and $\Omega^{(\beta)}$ with boundaries $\Gamma_1, \Gamma_2, \Gamma_3$:

In the composite spectral method we have

$$
\mathbf{T} = \begin{bmatrix}
\mathbf{T}^{(\alpha)}_{1,1} & 0 \\
0 & \mathbf{T}^{(\beta)}_{2,2}
\end{bmatrix} + \begin{bmatrix}
\mathbf{T}^{(\alpha)}_{1,3} \\
\mathbf{T}^{(\beta)}_{2,3}
\end{bmatrix} \left( \mathbf{T}^{(\alpha)}_{3,3} - \mathbf{T}^{(\beta)}_{3,3} \right)^{-1} \left[ -\mathbf{T}^{(\alpha)}_{3,1} \mid \mathbf{T}^{(\beta)}_{3,2} \right].
$$

There is more structure:

- The blue terms are of low numerical rank (say rank 40 to precision $10^{-10}$).
- The red terms are “hierarchically block separable” matrices.
  (Their off-diagonal blocks have low rank, cf. $\mathcal{H}$-matrices, etc).

The bottom line is that *the solution operators can be built in optimal $O(N)$ time*.
(Not true when $N$ is scaled to the wave-length for Helmholtz-type problems.)

*Joint work with Adrianna Gillman.*
Claim: Matrices with low-rank off-diagonal blocks can be inverted/multiplied/... rapidly.

As an example, consider a $2 \times 2$ blocked matrix of size $2n \times 2n$

$$A = \begin{bmatrix} A_{11} & A_{12} \\ A_{21} & A_{22} \end{bmatrix}.$$ 

Suppose the off-diagonal blocks are rank-deficient

$$A_{12} = U_1 \tilde{A}_{12} V_2^*$$
and

$$A_{21} = U_2 \tilde{A}_{21} V_1^*,$$

where $k \ll n$. We can then write $A$ as follows

$$A = \begin{bmatrix} A_{11} & 0 \\ 0 & A_{22} \end{bmatrix} + \begin{bmatrix} U_1 & 0 \\ 0 & U_2 \end{bmatrix} \begin{bmatrix} 0 & \tilde{A}_{12} \\ \tilde{A}_{21} & 0 \end{bmatrix} \begin{bmatrix} V_1^* & 0 \\ 0 & V_2^* \end{bmatrix}.$$ 

Recall the Woodbury formula

$$(D + U\tilde{A}V^*)^{-1} = D^{-1} - D^{-1}U(\tilde{A} + V^*D^{-1}U)^{-1}V^*D^{-1}.$$ 

Applying the Woodbury formula, we find, with $S_{11} = V_1^*A_{11}^{-1}U_1$ and $S_2 = V_2^*A_{22}^{-1}U_2$,

$$A^{-1} = \begin{bmatrix} A_{11}^{-1} & 0 \\ 0 & A_{22}^{-1} \end{bmatrix} + \begin{bmatrix} A_{11}^{-1}U_1 & 0 \\ 0 & A_{22}^{-1}U_2 \end{bmatrix} \begin{bmatrix} S_1 & \tilde{A}_{12} \\ \tilde{A}_{21} & S_2 \end{bmatrix}^{-1} \begin{bmatrix} V_1^*A_{11}^{-1} & 0 \\ 0 & V_2^*A_{22}^{-1} \end{bmatrix}.$$ 

Now suppose $A_{11}$ and $A_{22}$ have the same structure, and recurse.
Hierarchical Poincaré-Steklov Method: numerical results — $O(N)$ version

<table>
<thead>
<tr>
<th>Problem</th>
<th>$N$</th>
<th>$T_{\text{build}}$</th>
<th>$T_{\text{solve}}$</th>
<th>MB</th>
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</table>

(About six accurate digits in solution.)

Thanks to A. Barnett for use of a work-station!
Hierarchical Poincaré-Steklov Method: numerical results — $O(N)$ version

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<thead>
<tr>
<th>Problem</th>
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<th>$\epsilon = 10^{-12}$</th>
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Hierarchical Poincaré-Steklov Method: numerical results — $O(N)$ version

\[ \begin{aligned}
-\Delta u(x) - c_1(x) \partial_1 u(x) - c_2(x) \partial_2 u(x) - c(x) u(x) &= 0, \quad x \in \Omega, \\
u(x) &= f(x), \quad x \in \Gamma,
\end{aligned} \]  

(10)

**Laplace**  Let $c_1(x) = c_2(x) = c(x) = 0$ in (10).

**Helmholtz I**  Let $c_1(x) = c_2(x) = 0$, and $c(x) = \kappa^2$ where $\kappa = 80$ in (10). This represents a vibration problem on a domain $\Omega$ of size roughly $12 \times 12$ wave-lengths. (Recall that the wave-length is given by $\lambda = \frac{2\pi}{\kappa}$.)

**Helmholtz II**  Let $c_1(x) = c_2(x) = 0$, and $c(x) = \kappa^2$ where $\kappa = 640$ in (10). This corresponds to a domain of size roughly $102 \times 102$ wave-lengths.

**Helmholtz III**  We again set $c_1(x) = c_2(x) = 0$, and $c(x) = \kappa^2$ in (10), but now we let $\kappa$ grow as the number of discretization points grows to maintain a constant 12 points per wavelength.
Before showing the results from 3D ... some programming notes ...

- These results are very tentative ... code recently completed ...
- Timings for the BUILD stage are very bad ... can be greatly improved ... I think ...
- Memory requirements are bad (by current standards). Can be improved some.
- Solve time is excellent! And can be improved!
Hierarchical Poincaré-Steklov Method: numerical results — \( O(N) \) version in 3D

Set \( \Omega = [0, 1]^3 \) and \( \Gamma = \partial \Omega \). Consider the problem

\[
\begin{aligned}
\begin{cases}
-\Delta u(x) = 0, & x \in \Omega, \\
u(x) = f(x), & x \in \Gamma.
\end{cases}
\end{aligned}
\]

We pick \( f \) as the restriction of a field from a point source, \( x \mapsto |x - \hat{x}|^{-1} \).

We then know the exact solution, \( u_{\text{exact}}(x) = |x - \hat{x}|^{-1} \).

<table>
<thead>
<tr>
<th>( N_{\text{tot}} )</th>
<th>R (GB)</th>
<th>( T_{\text{build}} ) (sec)</th>
<th>( T_{\text{solve}} ) (sec)</th>
<th>( E^\infty )</th>
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<td>0.004</td>
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<td>6.51e-09</td>
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</tr>
</tbody>
</table>
Hierarchical Poincaré-Steklov Method: numerical results — $O(N)$ version in 3D

Set $\Omega = [0, 1]^3$ and $\Gamma = \partial \Omega$. Consider the problem

$$
\begin{cases}
-\Delta u(x) - \kappa^2 u(x) = 0, & x \in \Omega, \\
u(x) = f(x), & x \in \Gamma.
\end{cases}
$$

We pick $f$ as the restriction of a wave from a point source, $x \mapsto Y_0(\kappa|x - \hat{x}|)$.

We then know the exact solution, $u_{\text{exact}}(x) = Y_0(\kappa|x - \hat{x}|)$.

<table>
<thead>
<tr>
<th>$N_{\text{tot}}$</th>
<th>$N_{\text{Gauss}}$</th>
<th>Memory (GB)</th>
<th>$T_{\text{build}}$ (sec)</th>
<th>$T_{\text{solve}}$ (sec)</th>
<th>$E^\infty$</th>
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<td>1.1</td>
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<td>978.7</td>
<td>5.26e-05</td>
<td>1.48e-03</td>
</tr>
</tbody>
</table>

Results for solving Helmholtz equation with compression parameter $\epsilon = 10^{-5}$ with $20 \times 20 \times 20$ wavelength across the domain.
Note: In all cases, application of the solution operator is extremely fast.

Observation 1: The direct solver can be used to accelerate implicit time-stepping schemes for parabolic PDEs. As a toy example, consider:

\[
\begin{align*}
-\frac{\partial u(x, t)}{\partial t} &= -\Delta u, \quad x \in \Omega, \\
    u(x, t) &= f(x, t), \quad x \in \Gamma, \\
    u(x, 0) &= h(x), \quad x \in \Omega.
\end{align*}
\]

Say, for simplicity, that we use backwards Euler to discretize in time, with

\[
\frac{\partial u^n}{\partial t} \approx \frac{1}{k} (u^n - u^{n-1}).
\]

Then for each time-step we need to solve

\[
\begin{align*}
-\Delta u^n + \frac{1}{k} u^n &= \frac{1}{k} u^{n-1}, \quad \Omega, \\
    u^n &= f^n, \quad \Gamma.
\end{align*}
\]

This is very well suited for our direct solver.

Current work: Investigate stability with better time-stepping schemes (specifically ESDIRK). Numerical experiments are very promising. Extension to Stokes, low Reynolds number Navier-Stokes, etc.
Example: Consider the convection-diffusion problem
\[
\frac{\partial u}{\partial t} - \Delta u + 30 \frac{\partial u}{\partial x_1} = 0,
\]
defined on the domain \( \Omega \) shown below:

Zero Neumann condition on blue boundaries. Periodic BC on red boundaries.

The following mesh is used (observe corner refinement!):
**Observation 2:** The direct solver can be used to explicitly build time-evolution operators for hyperbolic problems. Consider, for instance,

\[
\begin{cases}
\frac{\partial u(x, t)}{\partial t} = Bu(x, t), & x \in \Omega, \ t > 0 \\
u(x, 0) = f(x) & x \in \Omega,
\end{cases}
\]

where \(B\) is a skew-Hermitian operator (e.g. \(B = \sqrt{\Delta}\) with Dirichlet/Neumann BC). The solution is

\[u(x, t) = \left[\exp(tB)f\right](x),\]

where \(\exp(tB)\) is the time-evolution operator. Now suppose that we can approximate the oscillatory function \(x \mapsto \exp(ix)\) by a rational function

\[R_M(ix) = \sum_{m=-M}^{M} \frac{b_m}{ix - \alpha_m},\]

where \(\{b_m\}\) and \(\{\alpha_m\}\) are some complex numbers such that \(|R_M(ix)| \leq 1\) for \(x \in \mathbb{R}\). We require that

\[|e^{ix} - R_M(ix)| \leq \delta, \quad x \in [-\tau \Lambda, \tau \Lambda],\]

where \(\tau\) is a time step, and where \(\Lambda\) is a “band-width” — in other words, we accurately resolve the parts of \(B\) whose spectrum fall in the interval \([-i\Lambda, i\Lambda]\). **Very high accuracy can be attained** — say \(\delta = 10^{-10}\) for about 5 – 10 points per wavelength [Beylkin, Haut]. Then approximate

\[\exp(\tau B) \approx \sum_{m=-M}^{M} b_m (B - \alpha_m)^{-1}.\]

**Notes:** The time-step \(\tau\) can be large. Application of \(\exp(\tau B)\) is almost instantaneous. Quite high memory demands, but distributed memory is fine. **Parallel in time!**

**Current project:** Shallow water equations on cubed sphere at LANL.
Direct solvers for integral equations

Recall that many boundary value problems can advantageously be recast as boundary integral equations. Consider, e.g., (sound-soft) acoustic scattering from a finite body:

\[
\begin{cases}
-\Delta u(x) - \kappa^2 u(x) = 0 & x \in \mathbb{R}^3 \setminus \Omega \\
u(x) = v(x) & x \in \partial \Omega \\
\lim_{|x| \to \infty} |x| (\partial_n u(x) - i\kappa u(x)) = 0.
\end{cases}
\]

(11)

The BVP (11) is in many ways equivalent to the BIE

\[
-\pi i \sigma(x) + \int_{\partial \Omega} \left( \left( \partial_n(y) + i\kappa \right) \frac{e^{i\kappa|x-y|}}{|x-y|} \right) \sigma(y) \, dS(y) = f(x), \quad x \in \partial \Omega.
\]

(12)

The integral equation (12) has several advantages over the PDE (11), including:

- The domain of computation \(\partial \Omega\) is finite.
- The domain of computation \(\partial \Omega\) is 2D, while \(\mathbb{R}^3 \setminus \Omega\) is 3D.
- Equation (12) is inherently well-conditioned (as a “2nd kind Fredholm equation”).

A serious drawback of integral equations is that they lead to dense coefficient matrices. Since we are interested in constructing inverses anyway, this is unproblematic for us!
Direct solvers for integral equations

It is possible to construct direct solvers that follow the same template as before.

Upwards pass — build all solution operators:

1. The original grid.
2. Leaves reduced.
3. After merge.
4. After merge.

Downwards pass — solve for a particular data function (very fast!):

5. Full solution.
5. Solve.
4. Top level solve.

Our “solution operators” will be (conceptually) scattering matrices instead of Poincaré-Steklov operators.

The operators will no longer be pure boundary operators.
**Example:** BIE on a surface in $\mathbb{R}^3$:

Let $A$ denote an $N \times N$ matrix arising upon discretizing a boundary integral operator

$$[Aq](x) = q(x) + \int_{\Gamma} \frac{1}{|x - y|} q(y) \, dA(y), \quad x \in \Gamma,$$

where $\Gamma$ is the “torus-like” domain shown (it is deformed to avoid rotational symmetry).
The reduced matrix represents a Nyström discretization supported on the panels shown.
The reduced matrix represents a Nyström discretization supported on the panels shown.
The reduced matrix represents a Nyström discretization supported on the panels shown.
The reduced matrix represents a Nyström discretization supported on the panels shown.
The reduced matrix represents a Nyström discretization supported on the panels shown.
The reduced matrix represents a Nyström discretization supported on the panels shown.
Example: Multibody scattering from a domain with multiple cavities

Consider scattering from some multibody domain involving cavities.
Example: Multibody scattering from a domain with multiple cavities

There are lots of discretization nodes involved. Very computationally intense!
Example: Multibody scattering from a domain with multiple cavities

After local compression of each scatter, the problem is much more tractable.
**Example:** Multibody scattering from a domain with multiple cavities.

Acoustic scattering on the exterior domain.

Each bowl is about $5\lambda$.

A hybrid direct/iterative solver is used (a highly accurate scattering matrix is computed for each body).

On an office desktop, we achieved an accuracy of $10^{-5}$, in about 6h (essentially all the time is spent in applying the inter-body interactions via the Fast Multipole Method). Accuracy $10^{-7}$ took 27h.
Example: BIEs on rotationally symmetric bodies (2014, with S. Hao and P. Young)

<table>
<thead>
<tr>
<th>$N$</th>
<th>$N_{\text{body}}$</th>
<th>$T_{\text{fmm}}$ (precond / no precond)</th>
<th>$I_{\text{GMRES}}$ (precond / no precond)</th>
<th>$T_{\text{total}}$ (precond / no precond)</th>
<th>$E_{\infty}$</th>
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<td>22 / 112</td>
<td>1.25e+03 / 5.84e+03</td>
<td>3.035e-06</td>
</tr>
</tbody>
</table>

*Exterior Laplace problem solved on the multibody bowl domain with and without preconditioner.*
Example: BIEs on rotationally symmetric bodies (2014, with S. Hao and P. Young)

<table>
<thead>
<tr>
<th>$N$</th>
<th>$N_{\text{body}}$</th>
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<th>$I_{\text{GMRES}}$</th>
<th>$T_{\text{solve}}$</th>
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<td>65</td>
<td>9.75e+04</td>
<td>6.884e-07</td>
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</tbody>
</table>

Exterior Helmholtz problem solved on multibody bowl domain.

Each bowl is 5 wavelength in diameter.

We do not give timings for standard iterative methods since in this example, they typically did not converge at all (even though the BIE is a 2nd kind Fredholm equation).
Numerical example — BIE on surfaces in 3D (2013, with J. Bremer and A. Gillman)

Consider sound-soft scattering from a multi-body scatterer of size 4 wave-lengths:

The global scattering matrix is computed using the hierarchical direct solver described. (The ellipsoids are not rotationally symmetric.)
Numerical example — BIE on surfaces in 3D (2013, with J. Bremer and A. Gillman)

The local truncation error is set to $10^{-3}$.

<table>
<thead>
<tr>
<th>Grid dimensions</th>
<th>$N$</th>
<th>$T$</th>
<th>$E$</th>
<th>Ratio</th>
<th>Predicted</th>
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<td>-</td>
<td>-</td>
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</tr>
<tr>
<td>$4 \times 4 \times 4$</td>
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<td>$1.57 \times 10^{-04}$</td>
<td>2.3</td>
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<td>$8 \times 8 \times 8$</td>
<td>786432</td>
<td>$6.70 \times 10^2$</td>
<td>$4.70 \times 10^{-04}$</td>
<td>2.3</td>
<td>3.7</td>
</tr>
<tr>
<td>$10 \times 10 \times 10$</td>
<td>1536000</td>
<td>$2.46 \times 10^3$</td>
<td>$3.53 \times 10^{-04}$</td>
<td>3.7</td>
<td>2.7</td>
</tr>
</tbody>
</table>
Numerical example — BIE on surfaces in 3D (2013, with J. Bremer and A. Gillman)

The local truncation error is set to $10^{-3}$.

<table>
<thead>
<tr>
<th>Grid dimensions</th>
<th>$N$</th>
<th>$T$</th>
<th>$E$</th>
<th>Ratio</th>
<th>Predicted</th>
</tr>
</thead>
<tbody>
<tr>
<td>$2 \times 2 \times 2$</td>
<td>12288</td>
<td>$1.02 \times 10^{+1}$</td>
<td>$3.37 \times 10^{-04}$</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>$3 \times 3 \times 3$</td>
<td>41472</td>
<td>$3.43 \times 10^{+1}$</td>
<td>$4.81 \times 10^{-04}$</td>
<td>3.4</td>
<td>6.2</td>
</tr>
<tr>
<td>$4 \times 4 \times 4$</td>
<td>98304</td>
<td>$7.92 \times 10^{+1}$</td>
<td>$1.57 \times 10^{-04}$</td>
<td>2.3</td>
<td>3.7</td>
</tr>
<tr>
<td>$6 \times 6 \times 6$</td>
<td>331776</td>
<td>$2.96 \times 10^{+2}$</td>
<td>$7.03 \times 10^{-04}$</td>
<td>3.7</td>
<td>6.2</td>
</tr>
<tr>
<td>$8 \times 8 \times 8$</td>
<td>786432</td>
<td>$6.70 \times 10^{+2}$</td>
<td>$4.70 \times 10^{-04}$</td>
<td>2.3</td>
<td>3.7</td>
</tr>
<tr>
<td>$10 \times 10 \times 10$</td>
<td>1536000</td>
<td>$2.46 \times 10^{+3}$</td>
<td>$3.53 \times 10^{-04}$</td>
<td>3.7</td>
<td>2.7</td>
</tr>
</tbody>
</table>

Increasing the accuracy is possible, but comes at a cost.

Now the local truncation error is set to $10^{-6}$.

<table>
<thead>
<tr>
<th>Grid dimensions</th>
<th>$N$</th>
<th>$T$</th>
<th>$E$</th>
<th>Ratio</th>
<th>Predicted</th>
</tr>
</thead>
<tbody>
<tr>
<td>$2 \times 2 \times 2$</td>
<td>49152</td>
<td>$1.61 \times 10^{+2}$</td>
<td>$1.22 \times 10^{-07}$</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>$3 \times 3 \times 3$</td>
<td>165888</td>
<td>$6.87 \times 10^{+2}$</td>
<td>$4.92 \times 10^{-07}$</td>
<td>4.3</td>
<td>6.2</td>
</tr>
<tr>
<td>$4 \times 4 \times 4$</td>
<td>393216</td>
<td>$1.68 \times 10^{+3}$</td>
<td>$5.31 \times 10^{-07}$</td>
<td>2.4</td>
<td>3.6</td>
</tr>
<tr>
<td>$6 \times 6 \times 6$</td>
<td>1327104</td>
<td>$6.66 \times 10^{+3}$</td>
<td>$4.60 \times 10^{-06}$</td>
<td>4.0</td>
<td>6.2</td>
</tr>
<tr>
<td>$8 \times 8 \times 8$</td>
<td>3145728</td>
<td>$1.59 \times 10^{+4}$</td>
<td>$2.30 \times 10^{-07}$</td>
<td>2.4</td>
<td>3.6</td>
</tr>
</tbody>
</table>
**Example:** Acoustic scattering from a “deformed torus” (with J. Bremer and A. Gillman)

The domain is roughly $2 \times 2 \times 0.7$ wave-lengths in size.

<table>
<thead>
<tr>
<th>$N_{\text{triangles}}$</th>
<th>$N$</th>
<th>$T$</th>
<th>$E$</th>
</tr>
</thead>
<tbody>
<tr>
<td>32</td>
<td>1 664</td>
<td>$7.16 \times 10^{+00}$</td>
<td>$3.51 \times 10^{-02}$</td>
</tr>
<tr>
<td>128</td>
<td>6 656</td>
<td>$6.29 \times 10^{+01}$</td>
<td>$4.41 \times 10^{-03}$</td>
</tr>
<tr>
<td>512</td>
<td>26 624</td>
<td>$2.81 \times 10^{+02}$</td>
<td>$4.08 \times 10^{-05}$</td>
</tr>
<tr>
<td>2 048</td>
<td>106 496</td>
<td>$2.60 \times 10^{+03}$</td>
<td>$7.80 \times 10^{-07}$</td>
</tr>
<tr>
<td>8 192</td>
<td>425 984</td>
<td>$1.47 \times 10^{+04}$</td>
<td>$3.25 \times 10^{-08}$</td>
</tr>
</tbody>
</table>

(Note: Laplace problems are much faster.)
A surface $\Gamma$ with corners and edges.

The grid has been refined to attain high accuracy.

Computing scattering matrices for the corners is conceptually easy (but laborious). The direct solver eliminates “extra” DOFs.

Compressing the edges takes effort!

<table>
<thead>
<tr>
<th>$N_{\text{tris}}$</th>
<th>$N$</th>
<th>$E$</th>
<th>$T$</th>
<th>$N_{\text{out}} \times N_{\text{in}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>192</td>
<td>21 504</td>
<td>$2.60 \times 10^{-08}$</td>
<td>$6.11 \times 10^{+02}$</td>
<td>$617 \times 712$</td>
</tr>
<tr>
<td>432</td>
<td>48 384</td>
<td>$2.13 \times 10^{-09}$</td>
<td>$1.65 \times 10^{+03}$</td>
<td>$620 \times 694$</td>
</tr>
<tr>
<td>768</td>
<td>86 016</td>
<td>$3.13 \times 10^{-10}$</td>
<td>$3.58 \times 10^{+03}$</td>
<td>$612 \times 685$</td>
</tr>
</tbody>
</table>

Results from a Helmholtz problem (acoustic scattering) on the domain exterior to the “edgy” cube. The domain is about 3.5 wave-lengths in diameter.
**Note:** We compress patches that are *directly adjacent*.

This is in contrast to, e.g., the Fast Multipole Methods, $\mathcal{H}$- and $\mathcal{H}^2$-matrix methods, etc.

**Advantages:** Easier data structures, more efficient inversion, better localization of data (leading to algorithms that are easier to parallelize).

**Disadvantages:** Ranks are higher, sometimes much higher.

*Numerical compression is required.*
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**Additional machinery required to attain $O(N)$ complexity in 3D:**

- Use Nested hierarchies — the dense blocks themselves have structure.

- Use multiple, staggered, grids.
Numerical example — Volume int. eq. in 2D (2013, with E. Corona and D. Zorin)

Consider a volume integral equation in the plane:

\[ q(x) + \int_{\Omega} b(x) \log |x - y| q(y) \, dy = f(x), \quad x \in \Omega, \]

where \( \Omega = [0, 1]^2 \), and where

\[ b(x) = 1 + 0.5e^{-(x_1 - 0.3)^2 - (x_2 - 0.6)^2}. \]

The domain is discretized on a uniform grid, with simplistic quadrature. By exploiting internal structure (HBS structure) in the scattering matrices, we have built a direct solver with optimal \( O(N) \) complexity for every step.
Execution times in Matlab, on an Intel Xeon X5650 (6 core) 2.67 GHz.

For a computed approximate inverse $B \approx A^{-1}$, the error reported is

$$\text{Error} = \max_i \frac{\|v^{(i)} - ABv^{(i)}\|}{\|v^{(i)}\|}$$

where $\{v^{(i)}\}_{i=1}^{10}$ is a collection of random vectors.
Key points on direct solvers for elliptic PDEs:

**Long range interactions are low rank.**
- Cf. St Venant principle, multipole expansions, etc.
- Numerical compression is essential.
- Wave problems with small $\lambda$ remain challenging.

![Graphs showing the behavior of solutions for Laplace and Helmholtz equations.](image)
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**Rank-structured matrix algebra. “Data Sparsity.”**
- Linear complexity operations for many global operators.
- Randomized compression is very helpful.

*All gray blocks have low rank.*
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**Hierarchical Divide-and-Conquer.**
- Generalizations of nested dissection.
- Using *double* hierarchies, we attain $O(N)$ complexity in most standard environments (including BIE).
- For BIEs, single hierarchy is often good enough.

**Current state of affairs:**
- We can solve certain problems that are intractable to iterative methods.
- For many direct solvers, every intermediate step is numerically benign.
- Direct solvers are memory hogs. (But distributed memory is OK.)
- We see major acceleration in certain environments (multiple solves).
Future directions within direct solvers:

- Accelerate build stage in 3D. (Already $O(N)$, but constants need improvement.)
- **Exploit communication efficiency in direct solvers!**
  - Low-hanging fruit for mid-size problems where the structured matrix algebra on matrices of size $O(N^{2/3}) \times O(N^{2/3})$ can be done on one node.
  - In the medium term, hybrid solvers that use direct solves for subproblems (e.g. near refined points in a mesh) will be highly useful.
  - The development of fully parallel codes where every component is executed using distributed memory will require hard work. Potentially huge rewards, however.
- Mathematical theory for direct solvers is sorely lacking. “Signing the blueprint.” Efficient and accurate à posteriori error estimation techniques do exist.
- Extension to parabolic and hyperbolic equations. *Parallel in time.*
- Real applications with complex geometries. Acoustics, sonar, imaging, etc.
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- **Full factorizations** — CPQR, UTV. Eigenvalue and singular value decompositions?
- Optimize for different architectures — GPU, distributed memory, out-of-core, etc.
- Randomized methods for sparse matrices — tighter connection with Krylov methods.
- High-dimensional geometry — clustering, non-linear manifolds, etc.
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Important: Availability of new powerful tools $\implies$ plenty of opportunities!