Randomized algorithms for large scale linear algebra

Per-Gunnar Martinsson
Dept. of Mathematics & Oden Institute for Computational Sciences and Engineering
University of Texas at Austin

Students, postdocs, collaborators: Ke Chen, Yijun Dong, Robert van de Geijn, Abinand Gopal, Nathan Halko, Nathan Heavner, James Levitt, Gregorio Quintana-Ortí, Joel Tropp, Sergey Voronin, Bowei Wu, Anna Yesypenko.

Slides: http://users.ices.utexas.edu/~pgm/main_talks.html

Research support by:
Randomized dimension reduction

Let \( \{a^{(j)}\}_{j=1}^n \) 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.

- Given \( k \), find the subset of \( k \) vectors with **maximal spanning volume**.

- Suppose the points almost live on a low-dimensional **nonlinear manifold**.
  
  Find a parameterization of the manifold.

- Given \( k \), find for each vector \( a^{(j)} \) 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 moderate complexity; some are combinatorially hard.)

**Idea:** Find an embedding \( f : \mathbb{R}^m \to \mathbb{R}^d \) for \( d \ll m \) that is almost isometric in the sense

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

Then solve the problems for the vectors \( \{f(a^{(j)})\}_{j=1}^n \) in \( \mathbb{R}^d \).
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  Find a basis for the “best” subspace.
- Given \( k \), find the subset of \( k \) vectors with maximal spanning volume.
- Suppose the points almost live on a low-dimensional nonlinear manifold.
  Find a parameterization of the manifold.
- Given \( k \), find for each vector \( a^{(j)} \) its \( k \) closest neighbors.
- Partition the points into clusters.

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

**Lemma [Johnson-Lindenstrauss]:** For \( d \sim \log(n) \), there exists a well-behaved embedding \( f \) 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

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

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

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

You can prove that if you pick \( d \) as specified, and draw a Gaussian random matrix \( R \) of size \( d \times m \), then there is a positive likelihood that the map

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

satisfies the criteria.

**Practical problem:** You have two 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?
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**Putative answer:** Yes — use a two-stage approach:

(A) *Randomized sketching:*

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

\[
\mathbf{A} = \begin{bmatrix}
\mathbf{a}^{(1)} & \mathbf{a}^{(2)} & \cdots & \mathbf{a}^{(n)}
\end{bmatrix} \in \mathbb{R}^{m \times n}.
\]

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.

*Jones, Osipov, Rokhlin, 2011*
Outline of talk

(1) Randomized dimension reduction — introduction.

(2) Computing low rank approximations (“randomized singular value decomposition”).

(3) Solving $Ax = b$. 
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Randomized SVD:

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

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

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

where $U$ and $V$ are orthonormal, and $D$ is diagonal. (We assume $k \ll \min(m, n)$.)
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(A) Randomized sketching:
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.
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**Objective:** Given an \( m \times n \) matrix \( A \), find an approximate rank-\( k \) partial SVD:

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A.1 Draw an \( n \times k \) Gaussian random matrix \( R \).

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R = \text{randn}(n,k)
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A.2 Form the \( m \times k \) sample matrix \( Y = AR \).

\[
Y = A \times R
\]

A.3 Form an \( m \times k \) orthonormal matrix \( Q \) such that \( \text{ran}(Q) = \text{ran}(Y) \).

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

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

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

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

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[\hat{U}, \Sigma, \hat{V}] = \text{svd}(B,'econ')
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B.3 Form the matrix \( U = Q \hat{U} \).

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B.3 Form the matrix $U = Q \text{Uhat}$.  
$U = Q \ast \text{Uhat}$

The objective of Stage A is to compute an ON-basis that approximately spans the column space of $A$. The matrix $Q$ holds these basis vectors and $A \approx QQ^*A$. 
**Randomized SVD:**

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

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The objective of Stage A is to compute an ON-basis that approximately spans the column space of \( A \). The matrix \( Q \) holds these basis vectors and \( A \approx QQ^*A \).

Stage B is exact:

\[
\|A - QQ^*A\|_{B} = \|A - Q\hat{U}DV^*\| = \|A - UdV^*\| = \|A - UDV^*\|
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B.1 Form the \( k \times n \) matrix \( B = Q^* A \).
B.2 Form SVD of the matrix \( B: B = \hat{U}D\hat{V}^* \).
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How does it work? To develop intuition, it helps to first consider the case \( \text{rank}(A) = k \).
Then \( \text{ran}(Y) = \text{ran}(A) \) holds with probability 1, so the output is exactly the SVD of \( A \).
In the general case, contributions from the singular modes beyond the first \( k \) will shift \( \text{ran}(Y) \) away from the desired space spanned by the dominant \( k \) left singular vectors.
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**Objective:** Given an $m \times n$ matrix $A$, find an approximate rank-$k$ partial SVD:

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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. By drawing $p$ extra samples (for, say, $p = 5$ or $p = 10$), we make the risk of missing anything important essentially zero.
Randomized SVD:

**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 $A \approx UDV^\ast$.

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- 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 “structure preserving” factorizations where a number of the columns/rows are chosen to serve as a basis for the column/row space. “Interpolative decomposition” / “CUR decomposition” / “skeletonization” / … → *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 $Y = AG$. 
Randomized SVD:

The plot shows the errors from the randomized range finder. To be precise, we plot

\[ e_k = \| A - P_k A \|, \]

where \( P_k \) is the orthogonal projection onto the first \( k \) columns of

\[ Y = (AA^*)^q AG, \]

and where \( G \) is a Gaussian random matrix. (Recall that \( P_k A = U_k D_k V_k^* \).

The matrix \( A \) is an approximation to a scattering operator for a Helmholtz problem.
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and where \( G \) is a Gaussian random matrix. (Recall that \( P_k A = U_k D_k V_k^* \).) The matrix \( A \) now has singular values that decay slowly.
Randomized SVD: The same plot as before, but now showing 100 instantiations.

The darker lines show the mean errors across the 100 experiments.
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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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**Theorem:** Let $A$ be an $m \times n$ matrix with singular values $\{\sigma_j\}_{j=1}^{\min(m,n)}$. Let $k$ be a target rank, and let $p$ be an over-sampling parameter such that $p \geq 2$ and $k + p \leq \min(m, n)$. Let $R$ be a Gaussian random matrix of size $n \times (k + p)$ and set $Q = \text{orth}(AR)$. Then the average error satisfies

$$
\mathbb{E}\left[\|A - QQ^*A\|_{Fro}\right] \leq \left(1 + \frac{k}{p - 1}\right)^{1/2} \left(\sum_{j=k+1}^{\min(m,n)} \sigma_j^2\right)^{1/2},
$$

$$
\mathbb{E}\left[\|A - QQ^*A\|\right] \leq \left(1 + \sqrt{\frac{k}{p - 1}}\right) \sigma_{k+1} + \frac{e\sqrt{k + p}}{p} \left(\sum_{j=k+1}^{\min(m,n)} \sigma_j^2\right)^{1/2}.
$$
**Randomized SVD:**

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We can reduce the flop count from $O(mnk)$ to $O(mn\log k)$ by using a so called “fast Johnson-Lindenstrauss” transform. A popular choice is the *subsampled random Fourier Transform (SRFT)* which can be applied using a variation of the FFT. Many other options: sub-sampled Hadamard transform, chains of Givens rotations, sparse projections, ...
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Example: The SRFT takes the form

$$R = D \ F \ S.$$ 

$\begin{array}{cccc}
\text{n \times k} & \text{n \times n} & \text{n \times n} & \text{n \times k} \\
\end{array}$

- $D$ is a diagonal matrix whose entries are i.i.d. random variables drawn from a uniform distribution on the unit circle in $\mathbb{C}$.
- $F$ is the discrete Fourier transform, $F_{pq} = \frac{1}{\sqrt{n}} e^{-2\pi i (p-1)(q-1)/n}$.
- $S$ is a matrix whose entries are all zeros except for a single, randomly placed 1 in each column. (So the action of $S$ is to draw $k$ columns at random from $DF$.)
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- The algorithm must be modified a bit beside replacing the random matrix.
- The SRFT leads to large speed-ups for moderate matrix sizes. For instance, for $m = n = 4000$, and $k \sim 10^2$, we observe about $\times 5$ speedup.
- In practice, accuracy is similar to what you get from Gaussian random matrices.
- Theory is still quite weak.

**References:** Ailon and Chazelle (2006); Liberty, Rokhlin, Tygert, and Woolfe (2006). Halko, Martinsson, Tropp (2011). Much subsequent work ...
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The perhaps most important feature of randomized algorithms in linear algebra 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.”)
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(2) Computing low rank approximations (“randomized singular value decomposition”).

(3) Solving \( Ax = b \) where \( A \in \mathbb{R}^{n\times n} \). (Focus is on the case where \( A \) is dense.)
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- Faster than $O(n^3)$ methods for general coefficient matrices?
- Linear complexity methods for “special” coefficient matrices.
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Observation: When $A$ is well-conditioned, iterative methods converge rapidly. Worst case complexity for solving $Ax = b$ to precision $\varepsilon$ is then $O(n^2 \log(1/\varepsilon))$. The challenge concerns matrices that are ill-conditioned. (Or, to be more precise, whose spectra are not clustered.)
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

Suppose $A$ is a dense ill-conditioned matrix of moderate size. In such a case, it is natural to look to $O(n^3)$ methods that compute a full factorization of the matrix.

Standard options (all with complexity $O(n^3)$) include:

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Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

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Not always stable.  
Fast.  

Always stable.  
Slow.

The “robust” factorizations to the right all depend on algorithms that proceed through a sequence of rank-one updates to the matrix. This makes them slow when executed on modern hardware (even on a single core).
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

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Computational time to factorize matrix

![Graph showing computational time for CPQR and QR methods](image-url)
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

The culprit preventing us from attaining high performance is *pivoting* since it relies on a sequence of rank-one updates.

Further refinements — Demmel, Dumitriu, Holtz, Grigori, Dongarra, etc.
Randomized methods for solving $\mathbf{Ax} = \mathbf{b}$: $O(n^3)$ complexity methods

The culprit preventing us from attaining high performance is *pivoting* since it relies on a sequence of rank-one updates.

**Randomization to the rescue!** D. Stott Parker (1995) proposed an elegant solution:

1. Randomly mix the columns by right multiplying $\mathbf{A}$ by a random unitary matrix $\mathbf{V}$:
   \[ \mathbf{A}_{\text{rand}} = \mathbf{AV}. \]

2. Perform unpivoted QR on the new matrix $\mathbf{A}_{\text{rand}} = \mathbf{UR}$

The resulting factorization
   \[ \mathbf{A} = \mathbf{A}_{\text{rand}} \mathbf{V}^* = \mathbf{URV}^* \]

is provably “rank-revealing” and leads to stable linear solves.

For computational efficiency, Parker introduced a random structured matrix (a bit ahead of the times) called a “random butterfly transform”.

Further refinements — Demmel, Dumitriu, Holtz, Grigori, Dongarra, etc.
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

Improved URV factorization: Do $q$ steps of power iteration (for $q = 1$ or $q = 2$, say):

1. Draw a Gaussian random matrix $G$ and form $Y = (AA^*)^q G$.
2. Perform unpivoted QR on $Y$ so that $Y = VR_{\text{trash}}$.
3. Perform unpivoted QR on $AV$ so that $AV = UR$.

This results in a factorization

$$A = (AV)V^* = URV^*$$

that is excellent at revealing the rank of $A$. Faster than CPQR, despite far more flops.
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The method is extremely simple to code:

```matlab
G = randn(n);
for j = 1:q
    G = A*(A'*G);
end
[V,~] = qr(G);
[U,R] = qr(A*V);
```
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

Given a dense $n \times n$ matrix $A$, compute a column pivoted QR factorization

$$A \sim Q \cdot R,$$

where, as usual, $Q$ should be ON, $P$ is a permutation, and $R$ is upper triangular.

The technique proposed is based on a *blocked* version of classical Householder QR:

$$A_0 = A \quad A_1 = Q_1^*A_0P_1 \quad A_2 = Q_2^*A_1P_2 \quad A_3 = Q_3^*A_2P_3 \quad A_4 = Q_4^*A_3P_4$$

Each $P_j$ is a permutation matrix computed via randomized sampling. Each $Q_j$ is a product of Householder reflectors.

The key challenge has been to find good permutation matrices. We seek $P_j$ so that the set of $b$ chosen columns has maximal spanning volume.

Perfect for randomized sampling! The likelihood that any block of columns is “hit” by the random vectors is directly proportional to its volume. Perfect optimality is not required.
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

How to do block pivoting using randomization:

Let $A$ be of size $m \times n$, and let $b$ be a block size.

\[ A \rightarrow Q^*AP \]

$Q$ is a product of $b$ Householder reflectors.

$P$ is a permutation matrix that moves $b$ “pivot” columns to the leftmost slots.

We seek $P$ so that the set of chosen columns has maximal spanning volume.

Draw a Gaussian random matrix $G$ of size $b \times m$ and form

\[ Y = \begin{bmatrix} G & A \\ b \times n & b \times m \end{bmatrix} \]

The rows of $Y$ are random linear combinations of the rows of $A$.

Then compute the pivot matrix $P$ for the first block by executing traditional column pivoting on the small matrix $Y$:

\[ YP = \begin{bmatrix} Q_{\text{trash}} & R_{\text{trash}} \\ b \times n & n \times n & b \times b & b \times n \end{bmatrix} \]
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

Speedup 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/
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

Given a dense $n \times n$ matrix $A$, compute a factorization

$$A = U \ T \ V^*,\quad n \times n \ n \times n \ n \times n \ n \times n \ n \times n$$

where $T$ is upper triangular, $U$ and $V$ are unitary.

Observe: More general than CPQR since we used to insist that $V$ be a permutation.

The technique proposed is based on a blocked version of classical Householder QR:

$$A_0 = A \quad A_1 = U_1^* A_0 V_1 \quad A_2 = U_2^* A_1 V_2 \quad A_3 = U_3^* A_2 V_3 \quad A_4 = U_4^* A_3 V_4$$

Both $U_j$ and $V_j$ are (mostly...) products of $b$ Householder reflectors.

Our objective is in each step to find an approximation to the linear subspace spanned by the $b$ dominant singular vectors of a matrix. The randomized range finder is perfect for this, especially when a small number of power iterations are performed. Easier and more natural than choosing pivoting vectors.
Rank-k approximation errors for the matrix “Fast Decay” of size $4000 \times 4000$. The black lines mark the theoretically minimal errors. The block size was $b = 100$ and the green vertical lines mark block limits.
Rank-k approximation errors for the matrix “BIE” of size $4000 \times 4000$. The black lines mark the theoretically minimal errors. The block size was $b = 100$ and the green vertical lines mark block limits.
Rank-k approximation errors for $k \leq 300$ for the matrix “Gap” of size $4000 \times 4000$. The black lines mark the theoretically minimal errors. The block size was $b = 100$ and the green vertical lines mark block limits.
The diagonal entries of the $T$-matrix in the UTV decomposition (red) provide excellent approximations to the true singular values (black).
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

For the task of computing low-rank approximations to matrices, the classical choice is between SVD and column pivoted QR (CPQR). SVD is slow, and CPQR is inaccurate:

Accuracy

Optimal

Ok

Speed

Slow

Fast

SVD

CPQR
Randomized methods for solving $Ax = b$: $O(n^3)$ complexity methods

For the task of computing low-rank approximations to matrices, the classical choice is between SVD and column pivoted QR (CPQR). SVD is slow, and CPQR is inaccurate:

Accuracy

Optimal  SVD

Very good  randUTV

Ok  CPQR  randCPQR

Speed

Slow  Fast  Faster!

The randomized algorithm **randUTV** combines the best properties of both factorizations. Additionally, **randUTV** parallelizes better, and allows the computation of partial factorizations (like CPQR, but unlike SVD).
Randomized methods for solving $Ax = b$: Strassen-type methods

The essential feature of the randomized methods described is that they enable us to expend almost all flops on the matrix-matrix computation, which is much faster per flop than other matrix operations.

Alternatively, use asymptotically faster methods for the matrix-matrix multiplication:

- **Strassen**: $O(n^{2.83})$. Stable. Reasonable breakeven point.

- **Coppersmith-Winograd etc.**: $O(n^{2.37})$. Unstable. Unreasonable breakeven point.

**Observation:**

Randomization allows you to use “fast” matrix-matrix multiplication algorithms to compute rank-revealing factorizations in a numerically stable way. In particular:

$$\text{fast+stable matrix-matrix multiplication} \implies \text{fast+stable linear system solve}$$

Randomized methods for solving $Ax = b$: Randomized pre-conditioning

Let us consider

$$Ax = b$$

for $A \in \mathbb{R}^{m \times n}$ with $m \gg n$. Complexity of standard solvers: $O(mn^2)$
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Method proposed by Rokhlin and Tygert (PNAS 2008): Form a “sketched equation”

$$X^*Ax = X^*b$$

where $X$ is an $m \times \ell$ SRFT. Compute QR factorization of the new coefficient matrix

$$X^*A = QRP^*.$$ 

Form a preconditioner

$$M = RP^*.$$ 

Solve the preconditioned linear system

$$\begin{pmatrix} AM^{-1} \end{pmatrix} \begin{pmatrix} Mx \end{pmatrix} = b$$

for the new unknown $y$. Complexity of randomized solver: $O((\log(n) + \log(1/\varepsilon))mn + n^3)$.

Later improvements include BLENDENPIK by Avron, Maymounkov, Toledo (2010).
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Randomized methods for solving $Ax = b$: Randomized Kaczmarz

The classical Kaczmarz algorithm:

With $A \in \mathbb{R}^{m \times n}$, we seek to solve $Ax = b$ through an iterative procedure. Given an approximate solution $x_{\text{old}}$, compute an improved solution $x_{\text{new}}$ as follows:

1. Pick a row index $i \in \{1, 2, \ldots, m\}$.
2. Require that $x_{\text{new}}$ is picked so that row $i$ of the system is satisfied exactly.
3. Within the hyperplane defined by (2), pick $x_{\text{new}}$ as the point closest to $x_{\text{old}}$.

The resulting formula is

$$x_{\text{new}} = x_{\text{old}} + \frac{b(i) - A(i,:) \cdot x_{\text{old}}}{\|A(i,:)\|_2^2} A(i,:).$$
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**Question:** How do you pick the row index $i$ in step (1)?
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**Strohmer & Vershynin (2009):** Draw $i$ with probability proportional to $\|A(i,:)\|$.

**Theorem:** Let $x_\star$ denote the exact solution to $Ax = b$, and let $x_k$ denote the $k$'th iterate of the S&V randomized Kaczmarz method. Then
\[
\mathbb{E}[\|x_k - x_\star\|] \leq \left(1 - \frac{1}{\kappa(A)^2}\right)^k \|x_0 - x_\star\|,
\]
where $\kappa(A)$ is the “scaled” condition number $\kappa(A) = \|A\|_F \|A^{-1}\|_2$. 

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Question: How do you pick the row index $i$ in step (1)?

Gower & Richtarik (2015): Draw an $m \times \ell$ random map $X$

$$x_{\text{new}} = \arg\min \{ \|y - x_{\text{old}}\|: y \text{ satisfies } X^*Ay = X^*b \}.$$ 

Leads to stronger analysis, and a much richer set of dimension reducing maps. In particular, it improves practical performance since it enables blocking.

Note: An ideal weight for a group of rows would be their spanning volume ...
Randomized methods for solving $Ax = b$: Randomized Newton-Schulz

Classical Newton-Schulz for computing $A^{-1}$: With $A \in \mathbb{R}^{n \times n}$, we build $B = A^{-1}$ through an iterative scheme. Given an approximation $B_{\text{old}}$, the improved one is

$$B_{\text{new}} = B_{\text{old}} - AB_{\text{old}}A.$$

Converges rapidly from a good initial guess. But basin of convergence is not large.

Gower & Richtarik (2019): Find $B = A^{-1}$ by solving the equation

$$A^* = A^*AB. $$

Equation (3) is solved through sketching + iteration: Draw an $m \times \ell$ random map $X$

$$B_{\text{new}} = \arg\min \{\|M - B_{\text{old}}\| : M \text{ satisfies } X^*A^* = X^*A^*AM\}.$$

Equivalent to iteration

$$B_{\text{new}} = B_{\text{old}} - A^*AX (X^*A^*AA^*AX)^\dagger X^*A^*(AB_{\text{old}} - I).$$

Detailed error analysis exists. For instance:

*The expectation of the error converges exponentially fast, regardless of starting point.*
Randomized iterative solvers is a very active area: J. Weare & L.-H. Lim article in SIAM Review (2017); recent and current work by H. Avron, P. Drineas, M. Mahoney, D. Needell, V. Rokhlin, S. Toledo, J. Tropp, R. Ward, and many more.
Randomized methods for solving $Ax = b$: Graph Laplacians

Let us consider a linear system
\[
Ax = b
\]
involving a coefficient matrix that is a graph Laplacian with $n$ nodes and $m$ edges.

- $A = A^* \in \mathbb{R}^{n \times n}$.
- $A(i, j) \leq 0$ when $i \neq j$.
- $A(i, i) = -\sum_{j \neq i} A(i, j)$

We assume that the underlying graph is connected, in which case $A$ has a 1-dimensional nullspace. We enforce that $\sum_i x(i) = 0$ and $\sum_i b(i) = 0$ in everything that follows.

(a) A graph with $n = 5$ vertices, and $m = 6$ edges. The conductivities of each edge is marked with a Greek letter.

(b) The $5 \times 5$ graph Laplacian matrix associated with the graph shown in (a).
Randomized methods for solving $\mathbf{Ax} = \mathbf{b}$: Graph Laplacians

Let us consider a linear system

$$\mathbf{Ax} = \mathbf{b}$$

involving a coefficient matrix that is a graph Laplacian with $n$ nodes and $m$ edges.

**Standard solution techniques:**

- **Multigrid:** Works great for certain classes of matrices.

- **Cholesky:** Compute a decomposition

  $$\mathbf{A} = \mathbf{CC}^*,$$

  with $\mathbf{C}$ lower triangular. Always works. Numerically stable (when pivoting is used). Can be expensive since the factor $\mathbf{C}$ typically has far more non-zero entries than $\mathbf{A}$.

- **Incomplete Cholesky:** Compute an approximate factorization

  $$\mathbf{A} \approx \mathbf{CC}^*,$$

  where $\mathbf{C}$ is constrained to be as sparse as $\mathbf{A}$ (typically the same pattern). Then use CG to solve a system with the preconditioned coefficient matrix $\mathbf{C}^{-1}\mathbf{AC}^{-*}$. Can work very well, hard to analyze.
Randomized methods for solving $Ax = b$: Graph Laplacians

Let us consider a linear system

$$Ax = b$$

involving a coefficient matrix that is a graph Laplacian with $n$ nodes and $m$ edges.

**Randomized solution techniques:**

- **Spielman-Teng (2004):** Complexity $O(m \text{ poly}(\log n) \log(1/\varepsilon))$.
  
  Relies on graph theoretical constructs (low-stretch trees, graph sparsification, explicit expander graphs, ...). Important theoretical results.

- **Kyng-Lee-Sachdeva-Spielman (2016):** $O(m (\log n)^2)$.
  
  Relies on local sampling only. Much closer to a realistic algorithm.

The idea is to build an approximate sparse Cholesky factor that is accurate with high probability. For instance, the 2016 paper proposes to build factors for which

$$\frac{1}{2} A \preceq CC^* \preceq \frac{3}{2} A.$$

When this bound holds, CG converges as $O(\gamma^n)$ with $\gamma = \frac{\sqrt{3}-1}{\sqrt{3}+1} \approx 0.27$.

Sparsity is maintained by performing inexact rank-1 updates in the Cholesky procedure. As a group of edges in the graph is removed, a set of randomly drawn new edges are added, in a way that is correct *in expectation*.
Randomized approximation of “rank-structured” matrices

Many matrices in applications have off-diagonal blocks that are of low rank:
- Matrices approximating integral equations associated with elliptic PDEs. (Essentially, discretized Calderón-Zygmund operators.)
- Scattering matrices in acoustic and electro-magnetic scattering.
- Inverses of (sparse) matrices arising upon FEM discretization of elliptic PDEs.
- Buzzwords: $\mathcal{H}$-matrices, HSS-matrices, HBS matrices, …

Using randomized algorithms, we have developed $O(N)$-complexity methods for performing algebraic operations on dense matrices of this type. This leads to:
- Accelerated direct solvers for elliptic PDEs.
- $O(N)$ complexity in many situations.

A representative tessellation of a rank-structured matrix. Each off-diagonal block (gray) has low numerical rank. The diagonal blocks (red) are full rank, but are small in size. Matrices of this type allow efficient matrix-vector multiplication, matrix inversion, etc.
Existing randomized compression schemes for rank-structured matrices:

Let $\mathbf{A}$ be a rank-structured matrix, for which we can rapidly evaluate $\mathbf{x} \mapsto \mathbf{A}\mathbf{x}$ and $\mathbf{x} \mapsto \mathbf{A}^*\mathbf{x}$.

**Case 1:** Suppose that in addition to matvec, we can also evaluate individual entries of $\mathbf{A}$. Then an HBS (a.k.a. HSS) representation can be computed in $O(N)$ operations. Very computationally efficient in practice — requires only $O(k)$ matvecs.

- P.G. Martinsson, SIMAX, **32**(4), 2011.
- Later improvements by Jianlin Xia, Sherry Li, etc.

**Case 2:** If all we have is the matvec, then we can still compute a rank-structured representation of $\mathbf{A}$ using so called “peeling” algorithms. The price we have to pay is that we now need $O(k \times \log N)$ matvecs involving $\mathbf{A}$ and $\mathbf{A}^*$.

The method is still fast in many situations, and does save messy coding work. For instance, without this black-box method, implementing the matrix-matrix multiplication, or changing the partition tree, are quite hard to implement efficiently.

“Hierarchically Off-Diagonal Low Rank (HODLR)” matrices

This is a very simple structured matrix format.

Informally, it means that you tessellate a matrix “like this”:

Then \( \mathbf{A} \) is HODLR if each off-diagonal block (in blue in the figure) has “low rank”.

**Note:** A.k.a. “S-matrix” (“S” for simple), \( \mathcal{H} \)-matrix with weak permissibility condition, etc.
To more formally define the HODLR format, we need to introduce a *tree structure* on the index set $I = [1, 2, 3, \ldots, N]$.

Let $A$ be an $N \times N$ matrix.

Suppose $T$ is a binary tree on the index vector $I = [1, 2, 3, \ldots, N]$.

For a node $\tau$ in the tree, let $I_\tau$ denote the corresponding index vector.

For nodes $\sigma$ and $\tau$ on the same level, set $A_{\sigma,\tau} = A(I_\sigma, I_\tau)$.
Formal definition of HODLR format

For every sibling pair \( \{\alpha, \beta\} \) set
\[
A_{\alpha,\beta} = A(I_\alpha, I_\beta).
\]
Fix a bound \( k \) on the rank, and a tolerance \( \varepsilon \). We then require that each off-diagonal block (in blue in the figure) have \( \varepsilon \)-rank at most \( k \). Define factors
\[
A_{\alpha,\beta} = U_\alpha \tilde{A}_{\alpha,\beta} V_\beta^*.
\]
\[
n_\alpha \times n_\beta \quad n_\alpha \times k \quad k \times k \quad k \times n_\beta
\]
For every sibling pair \{\alpha, \beta\} set
\[
A_{\alpha, \beta} = A(l_\alpha, l_\beta).
\]

Fix a bound \(k\) on the rank, and a tolerance \(\varepsilon\). We then require that each off-diagonal block (in blue in the figure) have \(\varepsilon\)-rank at most \(k\). Define factors
\[
A_{\alpha, \beta} = U_\alpha \tilde{A}_{\alpha, \beta} V_\beta^*.
\]

The cost of performing a matvec is then
\[
2 \times \frac{N}{2} k + 4 \times \frac{N}{4} k + 8 \times \frac{N}{8} k + \cdots \sim N \log(N) k.
\]
At level 0, we seek to factorize the red blocks.

Green blocks are random matrices, yellow blocks are sample matrices $Y_\tau$.

Now: $Y_2 = A_{2,3}R_3$ and $Y_3 = A_{3,2}R_2$.

The columns of $Y_\tau$ span $A(l_\tau, l_\sigma)$, where $\sigma$ is the sibling of $\tau$, so $U_\tau = qr(Y_\tau)$. 

At level 0, we seek to factorize the red blocks.

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How determine $\tilde{A}_{2,3}$, $\tilde{A}_{3,2}$, $\mathcal{V}_2$ and $\mathcal{V}_3$?
We know $\mathcal{U}_2$ and $\mathcal{U}_3$, and seek to determine $\tilde{A}_{2,3}$, $\tilde{A}_{3,2}$, $\mathcal{V}_2$ and $\mathcal{V}_3$.

Put $\mathcal{U}_2$ and $\mathcal{U}_3$ into the probing matrix.

Then: $Z_2 = A_{3,2}^* \mathcal{U}_3 = \mathcal{V}_2 A_{3,2}^*$ and $Z_3 = A_{2,3}^* \mathcal{U}_2 = \mathcal{V}_3 A_{2,3}^*$.

We then get $[\mathcal{V}_2, \tilde{A}_{3,2}^*] = qr(Z_2)$ and $[\mathcal{V}_3, \tilde{A}_{2,3}^*] = qr(Z_3)$. 
At level 1, we seek to determine the red blocks.

Green blocks are random matrices, yellow blocks are sample matrices $Y_\tau$.

*Subtract the contributions from the known gray blocks to get new sample matrices $Z_\tau$. The columns of $Z_\tau$ span $A(l_\tau, l_\sigma)$, where $\sigma$ is the sibling of $\tau$, so $U_\tau = \text{orth}(Y_\tau, \varepsilon)$. 
Put the matrices \( \{U_\tau\}_\tau \) is on level 1 into the probing matrix.

Then, e.g., \( Z_4 = A_{5,4}^* U_5 = \mathcal{V}_4 A_{5,4}^* \).

We get, e.g., \( [\mathcal{V}_4, \tilde{A}_{5,4}^*] = qr(Z_4, 0) \).
At level 2, we seek to determine the red blocks.

Green blocks are random matrices, yellow blocks are sample matrices $Y_\tau$.

*Subtract the contributions from the known gray blocks to get new sample matrices $Z_\tau$.*

The columns of $Y_\tau$ span $A(I_\tau, I_\sigma)$, where $\sigma$ is the sibling of $\tau$, so $U_\tau = \text{orth}(Y_\tau, \varepsilon)$.

By sampling $A^*$, we determine $V_\tau$ and $\tilde{A}_{\tau, \sigma}$. 

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**The “Peeling algorithm” — level 2**
Once you reach the finest level, simply use small identity matrices as probing matrices.

The diagonal blocks $D_\tau$ are obtained by subtracting contributions from the off-diagonal blocks from $Y_\tau$. 
Build compressed representations of all off-diagonal blocks.

**Loop** over levels $\ell = 0 : (L - 1)$

**Build the random matrices** $R_1$ and $R_2$.

$R_1 = \text{zeros}(n, r)$

$R_2 = \text{zeros}(n, r)$

**Loop** over boxes $\tau$ on level $\ell$

Let $\{\alpha, \beta\}$ denote the children of box $\tau$.

$R_1(l_\alpha, :) = \text{randn}(n_\alpha, r)$

$R_2(l_\beta, :) = \text{randn}(n_\beta, r)$

**End loop**

Apply $A$ to build the samples for the incoming basis matrices.

$Y_1 = AR_2 - A^{(l)}R_2$

$Y_2 = AR_1 - A^{(l)}R_1$

**Orthonormalize the sample matrices to build the incoming basis matrices.**

**Loop** over boxes $\tau$ on level $\ell$

Let $\{\alpha, \beta\}$ denote the children of box $\tau$.

$U_\alpha = \text{qr}(Y_1(l_\alpha, :))$.

$U_\beta = \text{qr}(Y_2(l_\beta, :))$.

**End loop**

**Apply $A^*$ to build the samples for the outgoing basis matrices.**

$Z_1 = A^*R_2 - (A^{(l)})^*R_2$

$Z_2 = A^*R_1 - (A^{(l)})^*R_1$

**Take local SVDs to build incoming basis matrices and sibling interaction matrices.**

**Loop** over boxes $\tau$ on level $\ell$

Let $\{\alpha, \beta\}$ denote the children of box $\tau$.

$[\mathbf{V}_\alpha, \mathbf{B}_{\alpha\beta}, \hat{\mathbf{U}}_\beta] = \text{svd}(Z_1(l_\alpha, :), \varepsilon)$.

$[\mathbf{V}_\beta, \mathbf{B}_{\alpha\beta}, \hat{\mathbf{V}}_\alpha] = \text{svd}(Z_2(l_\beta, :), \varepsilon)$.

$U_\beta \leftarrow U_\beta \hat{U}_\beta$.

$U_\alpha \leftarrow U_\alpha \hat{U}_\alpha$.

**End loop**

**Extract the diagonal matrices.**

$n_{\text{max}} = \max \{n_\tau : \tau \text{ is a leaf}\}$

$R = \text{zeros}(N, n_{\text{max}})$

**Loop** over leaf boxes $\tau$

$R(l_\tau, 1 : n_\tau) = \text{eye}(n_\tau)$.

**End loop**

$Y = AR - A^{(L)}R$

**Loop** over leaf boxes $\tau$

$D_\tau = Y(l_\tau, 1 : n_\tau)$.

**End loop**

The matrix $A^{(\ell)}$ consists of all blocks on level $\ell$ or coarser.
Key points on “peeling” algorithm for compressing rank-structured matrices:

- Entirely black-box — very easy to use.
- Complexity $O(N \log N)$.
- “Reasonable” practical speed.

**Question:** Is $O(N)$ complexity (without log factors) attainable?
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Answer: Yes, but currently with many caveats. Most importantly, current methods require the ability to evaluate a small number of individual entries of the matrix.
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**Research objectives:**

- Accelerate randomized compression algorithms for rank-structured matrices.
- Attempt to build $O(N)$ methods that do not require individual entries. This appears to be non-trivial …
- Improve performance for non-uniform trees.
- Improve accuracy by incorporating power iteration, if needed.
- Eliminate tuning parameters.
Key points:

- Randomized low-rank approximation ("randomized SVD").
  - Superior performance in many regards, in particular for very large problems.
  - For a fixed number of matrix-vector multiplies, Krylov methods are more accurate.

- Essential benefit of randomization in linear algebra: Reduces communication.
  - Enables processing of huge data sets. (Out-of-core / streaming / cloud computing / . . .)
  - Very fast on GPUs, distributed memory machines, etc.

- Recommended approach: Sketch first, then return to original data.
  (As opposed to: Sketch, then solve using only the sketched data.)

- Even though the algorithms are randomized, the output can be trusted.
  The probability of failure is extremely low (say $10^{-10}$).
  In most situations, you can explicitly compute the residual error.
  Cf. Monte Carlo vs. Las Vegas methods.
Future and ongoing work:

1. **Accelerate full factorizations of matrices.**
   - New randomized column pivoted QR algorithm is much faster than LAPACK.
   - New “UTV” factorization method is almost as accurate as SVD and much faster.

2. **Randomized algorithms for structured matrices.**
   - Use randomization to accelerate key numerical solvers for PDEs, for simulating Gaussian processes, etc.

3. **[High risk/high reward] Accelerate linear solvers for “general” systems $Ax = b$.**
   - The goal is methods with complexity $O(n^\gamma)$ for $\gamma < 3$. Crucially, we seek methods that retain stability, and have high practical efficiency for realistic problem sizes.

4. **Use randomized projections to accelerate non-linear algebraic tasks.**
   - Faster nearest neighbor search, faster clustering algorithms, etc. The idea is to use randomized projections for *sketching* to develop a rough map of a large data set. Then use high-accuracy deterministic methods for the actual computation.
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Great potential for new discoveries in linear algebra!
Papers (see also http://users.ices.utexas.edu/~pgm/main_publications.html):


Tutorials, summer schools, etc:

- 2014: CBMS summer school at Dartmouth College. 10 lectures on YouTube.

Software packages:

- Column pivoted QR: https://github.com/flame/hqrrp (much faster than LAPACK!)
- Randomized UTV: https://github.com/flame/randutv
- RSVDPACK: https://github.com/sergeyvoronin
- ID: http://tygert.com/software.html