Randomized Algorithms in Linear Algebra & the Column Subset Selection Problem

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To access my web page:

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Randomized algorithms & Linear Algebra

- **Randomized algorithms**
  - By (carefully) sampling rows/columns/entries of a matrix, we can construct new matrices (that have smaller dimensions or are sparse) and have bounded distance (in terms of some matrix norm) from the original matrix (with some failure probability).
  - By preprocessing the matrix using random projections (*), we can sample rows/columns/entries (?) much less carefully (uniformly at random) and still get nice bounds (with some failure probability).

(*) Alternatively, we can assume that the matrix is “well-behaved” and thus uniform sampling will work.
• **Randomized algorithms**
  
  • By (carefully) sampling rows/columns/entries of a matrix, we can construct new matrices (that have smaller dimensions or are sparse) and have bounded distance (in terms of some matrix norm) from the original matrix (with some failure probability).

  • By preprocessing the matrix using random projections, we can sample rows/columns/entries(?) much less carefully (uniformly at random) and still get nice bounds (with some failure probability).

• **Matrix perturbation theory**

  • The resulting smaller/sparser matrices behave similarly (in terms of singular values and singular vectors) to the original matrices thanks to the norm bounds.

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In this talk, I will illustrate some applications of the above ideas in the Column Subset Selection Problem and in approximating low-rank matrix approximations.
Interplay

(Data Mining) Applications
Biology & Medicine: population genetics (coming up...)
Electrical Engineering: testing of electronic circuits
Internet Data: recommendation systems, document-term data

Theoretical Computer Science
Randomized and approximation algorithms

Numerical Linear Algebra
Matrix computations and Linear Algebra (i.e., perturbation theory)
**Human genetics**

**Single Nucleotide Polymorphisms**: the most common type of genetic variation in the genome across different individuals.

They are known locations at the human genome where two alternate nucleotide bases (alleles) are observed (out of A, C, G, T).

SNPs

Matrices including thousands of individuals and hundreds of thousands if SNPs are available.
The Human Genome Diversity Panel (HGDP)

- 1,033 samples
- 7 geographic regions
- 52 populations

Rosenberg et al. (2002) Science
Li et al. (2008) Science
HGDP data
- 1,033 samples
- 7 geographic regions
- 52 populations

HapMap Phase 3 data
- 1,207 samples
- 11 populations

Rosenberg et al. (2002) Science
Li et al. (2008) Science
We will apply SVD/PCA on the (joint) HGDP and HapMap Phase 3 data.

**HGDP data**
- 1,033 samples
- 7 geographic regions
- 52 populations

**HapMap Phase 3 data**
- 1,207 samples
- 11 populations

**Matrix dimensions:**
- 2,240 subjects (rows)
- 447,143 SNPs (columns)

**Dense matrix:**
over one billion entries
Let the blue circles represent \( m \) data points in a 2-D Euclidean space.

Then, the SVD of the \( m \)-by-2 matrix of the data will return ...
Let the blue circles represent $m$ data points in a 2-D Euclidean space.

Then, the SVD of the $m$-by-$2$ matrix of the data will return ...

1st (right) singular vector:

direction of maximal variance,
Let the blue circles represent \( m \) data points in a 2-D Euclidean space.

Then, the SVD of the \( m \)-by-2 matrix of the data will return …

**1st (right) singular vector:**

direction of maximal variance,

**2nd (right) singular vector:**

direction of maximal variance, after removing the projection of the data along the first singular vector.
**Singular values**

σ₁: measures how much of the data variance is explained by the first singular vector.

σ₂: measures how much of the data variance is explained by the second singular vector.

Principal Components Analysis (PCA) is done via the computation of the Singular Value Decomposition (SVD) of a (mean-centered) covariance matrix.

Typically, a small constant number (say k) of the top singular vectors and values are kept.
HGDP data
- 1,033 samples
- 7 geographic regions
- 52 populations

HapMap Phase 3 data
- 1,207 samples
- 11 populations

Matrix dimensions:
2,240 subjects (rows)
447,143 SNPs (columns)

SVD/PCA returns...

Rosenberg et al. (2002) Science
Li et al. (2008) Science
• Top two Principal Components (PCs or eigenSNPs)
  (Lin and Altman (2005) *Am J Hum Genet*)

• The figure renders visual support to the “out-of-Africa” hypothesis.

• Mexican population seems out of place: we move to the top three PCs.
Not altogether satisfactory: the principal components are linear combinations of all SNPs, and - of course - can not be assayed!

Can we find actual SNPs that capture the information in the singular vectors? Formally: spanning the same subspace.
Issues

- **Computing large SVDs: computational time**
  - In commodity hardware (e.g., a 4GB RAM, dual-core laptop), using MatLab 7.0 (R14), the computation of the SVD of the dense 2,240-by-447,143 matrix $A$ takes about 12 minutes.
  - Computing this SVD is not a one-liner, since we cannot load the whole matrix in RAM (runs out-of-memory in MatLab).
  - We compute the eigendecomposition of $A A^T$.
  - In a similar experiment, we computed 1,200 SVDs on matrices of dimensions (approx.) 1,200-by-450,000 (roughly speaking a full leave-one-out cross-validation experiment).
    (Drineas, Lewis, & Paschou (2010) PLoS ONE)

- **Obviously, running time is a concern.**

- **We need efficient, easy to implement, methods.**
Issues (cont’d)

- Selecting good columns that “capture the structure” of the top PCs
  - Combinatorial optimization problem; hard even for small matrices.
  - Often called the Column Subset Selection Problem (CSSP).
  - Not clear that such columns even exist.
Issues (cont’d)

• Selecting good columns that “capture the structure” of the top PCs
  • Combinatorial optimization problem; hard even for small matrices.
  • Often called the Column Subset Selection Problem (CSSP).
  • Not clear that such columns even exist.

Such datasets will only continue to increase in size:

In collaboration with K. Kidd’s lab (Yale University, Department of Genetics) we are now analyzing:

• 4,000 samples from over 100 populations
• genotyped on over 500,000 SNPs.
Our perspective

The two issues are connected

• There exist “good” columns in any matrix that contain information about the top principal components.

• We can identify such columns via a simple statistic: the leverage scores.

• This does not immediately imply faster algorithms for the SVD, but, combined with random projections, it does!
SVD decomposes a matrix as...

\[
\begin{pmatrix}
A
\end{pmatrix}_{m \times n} \approx \begin{pmatrix}
U_k
\end{pmatrix}_{m \times k} \begin{pmatrix}
X
\end{pmatrix}_{k \times n}
\]

Top k left singular vectors

- It is easy to see that \( X = U_k^T A \).
- SVD has strong optimality properties.
- The columns of \( U_k \) are linear combinations of up to all columns of \( A \).
The CX decomposition

Drineas, Mahoney, & Muthukrishnan (2008) SIAM J Mat Anal Appl
Mahoney & Drineas (2009) PNAS

\[
\begin{pmatrix}
  m 	imes n \\
  A
\end{pmatrix} \approx \begin{pmatrix}
  m 	imes c \\
  C
\end{pmatrix} \begin{pmatrix}
  c 	imes n \\
  X
\end{pmatrix}
\]

Carefully chosen \( X \)

Goal: make (some norm) of \( A - CX \) small.

c columns of \( A \)

Why?

If \( A \) is an subject-SNP matrix, then selecting representative columns is equivalent to selecting representative SNPs to capture the same structure as the top eigenSNPs.

We want \( c \) as small as possible!
CX decomposition

\[
\begin{pmatrix} \begin{array}{c} m \times n \\ A \end{array} \end{pmatrix} \approx \begin{pmatrix} \begin{array}{c} m \times c \\ C \end{array} \end{pmatrix} \begin{pmatrix} \begin{array}{c} c \times n \\ X \end{array} \end{pmatrix}
\]

\(c\) columns of \(A\)

Easy to prove that optimal \(X = C^+A\). (\(C^+\) is the Moore-Penrose pseudoinverse of \(C\).)

Thus, the challenging part is to find good columns (SNPs) of \(A\) to include in \(C\).

From a mathematical perspective, this is a hard combinatorial problem, closely related to the so-called Column Subset Selection Problem (CSSP).

The CSSP has been heavily studied in Numerical Linear Algebra.
A much simpler statistic

(Frieze, Kannan, & Vempala FOCS 1998, Drineas, Frieze, Kannan, Vempala & Vinay SODA '99, Drineas, Kannan, & Mahoney SICOMP '06)

**Algorithm:** given an m-by-n matrix $A$, let $A^{(i)}$ be the i-th column of $A$.

- Sample $s$ columns of $A$ in i.i.d. trials (with replacement), where in each trial

  $$\Pr \text{[picking the } i\text{-th column]} = \frac{||A^{(i)}||_2^2}{||A||_F^2}$$

- Form the m-by-$s$ matrix $C$ by including $A^{(i)}$ as a column of $C$.

**Error bound:**

$$E \left[ \|A - CC^+A\|_F^2 \right] \leq \|A - A_k\|_F^2 + \sqrt{\frac{4k}{s}} \|A\|_F^2$$
Is this a good bound?

\[
\mathbb{E} \left[ \| A - CC^+ A \|_F^2 \right] \leq \| A - A_k \|_F^2 + \sqrt{\frac{4k}{s}} \| A \|_F^2
\]

**Problem 1:** If \( s = n \), we still do not get zero error.

That’s because of sampling with replacement.

(We know how to analyze uniform sampling without replacement, but we have no bounds on non-uniform sampling without replacement.)

**Problem 2:** If \( A \) had rank exactly \( k \), we would like a column selection procedure that drives the error down to zero when \( s = k \).

This can be done deterministically simply by selecting \( k \) linearly independent columns.

**Problem 3:** If \( A \) had *numerical rank* \( k \), we would like a bound that depends on the norm of \( A - A_k \) and not on the norm of \( A \).

A lot of prior work in the Numerical Linear Algebra community for the *spectral norm case* when \( s = k \); the resulting bounds depend (roughly) on \((k(n-k))^{1/2} \| A - A_k \|_2\)
Approximating singular vectors

1. Sample $s$ (=140) columns of the original matrix $A$ and rescale them appropriately to form a $512$-by-$c$ matrix $C$.

2. Project $A$ on $CC^*$ and show that $A-CC^*A$ is “small”.

($C^*$ is the pseudoinverse of $C$)
Approximating singular vectors

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2. Project $A$ on $CC^*$ and show that $A - CC^*A$ is “small”.

($C^*$ is the pseudoinverse of $C$)
Remark 1: Selecting the columns in this setting is trivial and can be implemented in a couple of (sequential) passes over the input matrix.

Remark 2: The proof is based on matrix perturbation theory and a probabilistic argument to bound $AA^T - \hat{C}\hat{C}^T$ (where $\hat{C}$ is a rescaled $C$).
Relative-error Frobenius norm bounds

Drineas, Mahoney, & Muthukrishnan (2008) SIAM J Mat Anal Appl

Given an $m$-by-$n$ matrix $A$, there exists an $O(mn^2)$ algorithm that picks

at most $O\left(\frac{k}{\epsilon^2} \log \frac{k}{\epsilon}\right)$ columns of $A$

such that with probability at least .9

$$\|A - CC^\dagger A\|_F \leq (1 + \epsilon) \|A - A_k\|_F$$
The algorithm

**Input:** m-by-n matrix $A$, 
$0 < \varepsilon < .5$, the desired accuracy

**Output:** $C$, the matrix consisting of the selected columns

**Sampling algorithm**

- Compute probabilities $p_j$ summing to 1.
- Let $c = O\left( \frac{k}{\varepsilon^2} \log \left( \frac{k}{\varepsilon} \right) \right)$.
- In $c$ i.i.d. trials pick columns of $A$, where in each trial the $j$-th column of $A$ is picked with probability $p_j$.
- Let $C$ be the matrix consisting of the chosen columns.
Subspace sampling (Frobenius norm)

\[
\begin{pmatrix}
A_k
\end{pmatrix}_{m \times n}
= \begin{pmatrix}
U_k
\end{pmatrix}_{m \times k} \cdot \begin{pmatrix}
\Sigma_k
\end{pmatrix}_{k \times k} \cdot \begin{pmatrix}
V_k^T
\end{pmatrix}_{k \times n}
\]

\(V_k\): orthogonal matrix containing the top \(k\) right singular vectors of \(A\).

\(\Sigma_k\): diagonal matrix containing the top \(k\) singular values of \(A\).

**Remark:** The rows of \(V_k^T\) are orthonormal vectors, but its columns \((V_k^T)^{(i)}\) are not.
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\end{pmatrix}
\cdot 
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V_k^T \\
k \times n
\end{pmatrix}
\]

\(V_k\): orthogonal matrix containing the top k right singular vectors of \(A\).
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**Remark:** The rows of \(V_k^T\) are orthonormal vectors, but its columns \((V_k^T)^{(i)}\) are not.

**Subspace sampling in** \(O(mn^2)\) **time**

\[
p_j = \frac{\left\| (V_k^T)^{(j)} \right\|_2^2}{k} \]

Normalization s.t. the \(p_j\) sum up to 1
Subspace sampling (Frobenius norm)

\[
\begin{pmatrix}
A_k \\
m \times n
\end{pmatrix}
= 
\begin{pmatrix}
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\end{pmatrix}
\cdot 
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\cdot 
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k \times n
\end{pmatrix}
\]

Remark: The rows of $V_k^T$ are orthonormal vectors, but its columns $(V_k^T)^{(i)}$ are not.

Subspace sampling in $O(mn^2)$ time

\[
p_j = \frac{\left\| (V_k^T)^{(j)} \right\|_2^2}{k}
\]

Leverage scores
(useful in statistics for outlier detection)

Normalization s.t. the $p_j$ sum up to 1
SNPs by chromosomal order

PCA

*top 30 PCA-correlated SNPs

Africa
Europe
Asia
America

SNPs by chromosomal order

Selecting PCA SNPs for individual assignment to four continents (Africa, Europe, Asia, America)

Leverage scores & effective resistances

Consider a weighted (positive weights only!) undirected graph $G$ and let $L$ be the Laplacian matrix of $G$.

Assuming $n$ vertices and $m > n$ edges, $L$ is an $n$-by-$n$ matrix, defined as follows:

\[
L = \begin{pmatrix} \ B^T \ \\ \ n \times m \end{pmatrix} \cdot \begin{pmatrix} \ W \ \\ \ m \times m \end{pmatrix} \cdot \begin{pmatrix} \ B \ \\ \ m \times n \end{pmatrix}
\]
Leverage scores & effective resistances

Consider a weighted (positive weights only!) undirected graph $G$ and let $L$ be the Laplacian matrix of $G$.

Assuming $n$ vertices and $m > n$ edges, $L$ is an $n$-by-$n$ matrix, defined as follows:

$$L = \left( B^T \right) \cdot \left( \begin{array}{c} 1 \end{array} \right) \cdot \left( \begin{array}{c} W^{1/2} \end{array} \right) \cdot \left( \begin{array}{c} B \end{array} \right).$$

Clearly, $L = (B^T W^{1/2})(W^{1/2}B) = (B^T W^{1/2})(B^TW^{1/2})^T$. 
Leverage scores & effective resistances

**Effective resistances:**

Let $G$ denote an electrical network, in which each edge $e$ corresponds to a resistor of resistance $1/w_e$.

The effective resistance $R_e$ between two vertices is equal to the potential difference induced between the two vertices when a unit of current is injected at one vertex and extracted at the other vertex.
Leverage scores & effective resistances

Effective resistances:

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The effective resistance $R_e$ between two vertices is equal to the potential difference induced between the two vertices when a unit of current is injected at one vertex and extracted at the other vertex.

Formally, the effective resistances are the diagonal entries of the $m$-by-$m$ matrix:

$$R = BL^T B^T = B(B^TW^2B)^+B^T$$

Lemma: The leverage scores of the $m$-by-$n$ matrix $W^{1/2}B$ are equal (up to a simple rescaling) to the effective resistances of the edges of $G$.

(Drineas & Mahoney, ArXiv '11)
Effective resistances are very important!

Very useful in graph sparsification (Spielman & Srivastava STOC ’08).

Graph sparsification is a critical step in solvers for Symmetric Diagonally Dominant (SDD) systems of linear equations (seminal work by Spielman and Teng).

Approximating effective resistances (Spielman & Srivastava STOC ’08)

They can be approximated using the SDD solver of Spielman and Teng.

Breakthrough by Koutis, Miller, & Peng (FOCS ’10, FOCS’11):

Low-stretch spanning trees provide a means to approximate effective resistances!

This observation (and a new, improved algorithm to approximate low-stretch spanning trees) led to almost optimal algorithms for solving SDD systems of linear equations.
Approximating leverage scores

Are leverage scores a viable alternative to approximate effective resistances?

Not yet! But, we now know the following:

**Theorem:** Given any $m$-by-$n$ matrix $A$ with $m > n$, we can approximate its leverage scores with relative error accuracy in $O(mn \log m)$ time,

as opposed to the - trivial - $O(mn^2)$ time.

*(Clarkson, Drineas, Mahoney, Magdon-Ismail, & Woodruff ArXiv ‘12)*
Approximating leverage scores

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$$O(mn \log m)$$

as opposed to the - trivial - $O(mn^2)$ time.

(Clarkson, Drineas, Mahoney, Magdon-Ismail, & Woodruff ArXiv ‘12)

**Not good enough for $W^{1/2}B$!**

This matrix is very sparse (2m non-zero entries). We must take advantage of the sparsity and approximate the leverage scores/effective resistances in $O(m \text{ polylog}(m))$ time.

Our algorithm will probably not do the trick, since it depends on random projections that “densify” the input matrix.
Problem
How many columns do we need to include in the matrix $C$ in order to get relative-error approximations?

Recall: with $O((k/\varepsilon^2) \log (k/\varepsilon))$ columns, we get (subject to a failure probability)

$$\| A - CC^\dagger A \|_F \leq (1 + \varepsilon) \| A - A_k \|_F$$

Deshpande & Rademacher (FOCS '10): with exactly $k$ columns, we get

$$\| A - CC^\dagger A \|_F \leq \sqrt{k} \| A - A_k \|_F$$

What about the range between $k$ and $O(k \log(k))$?
Selecting fewer columns (cont’d)

(Boutsidis, Drineas, & Magdon-Ismail, FOCS 2011)

**Question:**
What about the range between $k$ and $O(k \log(k))$?

**Answer:**
A relative-error bound is possible by selecting $s = \frac{3k}{\varepsilon}$ columns!

**Technical breakthrough:**
A combination of sampling strategies with a novel approach on column selection, inspired by the work of Batson, Spielman, & Srivastava (STOC ’09) on graph sparsifiers.

- The running time is $O((mnk+nk^3)\varepsilon^{-1})$.
- Simplicity is gone…
A two-phase algorithm

**Phase 1:**

Compute exactly (or, to improve speed, approximately) the top $k$ right singular vectors of $A$ and denote them by the $n$-by-$k$ matrix $\hat{V}_k$.

Construct an $n$-by-$r$ sampling-and-rescaling matrix $S$ such that

$$\sigma_k(\hat{V}_k^T S) > 1 - \sqrt{\frac{k}{r}}; \quad \| (A - A\hat{V}_k \hat{V}_k^T) S \|_F \leq \| A - A\hat{V}_k \hat{V}_k^T \|_F.$$
A two-phase algorithm

**Phase 1:**

Compute exactly (or, to improve speed, approximately) the top $k$ right singular vectors of $A$ and denote them by the $n$-by-$k$ matrix $\hat{V}_k$.

Construct an $n$-by-$r$ sampling-and-rescaling matrix $S$ such that

$$\sigma_k(\hat{V}_k^T S) > 1 - \sqrt{\frac{k}{r}}; \quad \|A - A\hat{V}_k \hat{V}_k^T S\|_F \leq \|A - A\hat{V}_k \hat{V}_k^T\|_F.$$

**Phase 2:**

Compute: $C_1 = AS; \; E = A - C_1 C_1^+ A$

Compute $p_i = \frac{\|E^{(i)}\|_2^2}{\|E\|_F^2}$ and sample $(s-r)$ columns with respect to the $p_i$'s.

**Output:** Return the columns of $A$ that correspond to the columns sampled in the phase 1 and phase 2.
The analysis

For simplicity, assume that we work with the exact top-k right singular vectors $V_k$.

**A structural result:**

\[
\|A - CC^+A\|_F^2 \leq \|A - A_k\|_F^2 + \|(A - A_k) S (V_k^T S)^+\|_F^2
\]
\[
\leq \|A - A_k\|_F^2 + \|(A - A_k) S\|_F^2 \left\| (V_k^T S)^+ \right\|_2^2
\]
\[
\leq \|A - A_k\|_F^2 + \|A - A_k\|_F^2 \left(1 - \sqrt{\frac{k}{r}}\right)^{-2}
\]

It is easy to see that setting $r = O(k/\varepsilon)$, we get a $(2+\varepsilon)$-multiplicative approximation.
The analysis

For simplicity, assume that we work with the exact top-k right singular vectors $V_k$.

**A structural result:**

$$
\| A - C C^+ A \|_F^2 \leq \| A - A_k \|_F^2 + \| (A - A_k) S (V_k^T S)^+ \|_F^2 \\
\leq \| A - A_k \|_F^2 + \| (A - A_k) S \|_F^2 \| (V_k^T S)^+ \|_2^2 \\
\leq \| A - A_k \|_F^2 + \| A - A_k \|_F^2 \left( 1 - \sqrt{\frac{k}{r}} \right)^{-2}
$$

It is easy to see that setting $r = O(k/\epsilon)$, we get a $(2+\epsilon)$-multiplicative approximation.

Phase 2 reduces this error to a $(1+\epsilon)$-multiplicative approximation; the analysis is similar to adaptive sampling.

(Deshpande, Rademacher, & Vempala SODA 2006).

Our full analysis accounts for approximate right singular vectors and works in expectation.
Spectral-Frobenius sparsification

Let $V$ be an $n$-by-$k$ matrix such that $V^TV=I$, with $k < n$, let $B$ be an $\ell n$-by-$n$ matrix, and let $r$ be a sampling parameter with $r > k$.

**Lemma**

There exists a deterministic algorithm which runs in time $O(rnk^2 + \ell n)$ and constructs a sampling matrix $S \in \mathbb{R}^{n \times r}$:

$$\sigma_k(V^TS) \geq 1 - \sqrt{\frac{k}{r}}$$

$$\|BS\|_F \leq \|B\|_F.$$

This lemma is inspired by the Spectral Sparsification result in (Batson, Spielman, & Srivastava, STOC 2009); there, it was used for graph sparsification.

Our generalization requires the use of a new barrier function which controls the Frobenius and spectral norm simultaneously.
Lower bounds and alternative approaches

**Deshpande & Vempala, RANDOM 2006**
A relative-error approximation necessitates at least $k/\varepsilon$ columns.

**Guruswami & Sinop, SODA 2012**
Alternative approaches, based on volume sampling, guarantee

$\frac{(r+1)}{(r+1-k)}$ relative error bounds.

This bound is asymptotically optimal (up to lower order terms).

The proposed deterministic algorithm runs in $O(rn^3 \log m)$ time, while the randomized algorithm runs in $O(rn^2)$ time and achieves the bound in expectation.

**Guruswami & Sinop, FOCS 2011**
Applications of column-based reconstruction in Quadratic Integer Programming.
For every set $S$ of $m$ points in $\mathbb{R}^n$ and every $\epsilon > 0$, there exists a mapping $f : \mathbb{R}^n \to \mathbb{R}^s$, where $s = O\left(\log m/\epsilon^2\right)$, such that for all points $u \in S$,

$$(1 - \epsilon) \|u\|_2 \leq \|f(u)\|_2 \leq (1 + \epsilon) \|u\|_2$$

holds with probability at least $1 - 1/m^2$.

Johnson & Lindenstrauss (1984)
Random projections: the JL lemma

For every set $S$ of $m$ points in $\mathbb{R}^n$ and every $\epsilon > 0$, there exists a mapping $f : \mathbb{R}^n \rightarrow \mathbb{R}^s$, where $s = O\left(\log m/\epsilon^2\right)$, such that for all points $u \in S$,

$$(1 - \epsilon) \|u\|_2 \leq \|f(u)\|_2 \leq (1 + \epsilon) \|u\|_2$$

holds with probability at least $1 - 1/m^2$.

Johnson & Lindenstrauss (1984)

• We can represent $S$ by an $m$-by-$n$ matrix $A$, whose rows correspond to points.
• We can represent all $f(u)$ by an $m$-by-$s$ $\tilde{A}$.
• The “mapping” corresponds to the construction of an $n$-by-$s$ matrix $R$ and computing $\tilde{A} = AR$

(The original JL lemma was proven by projecting the points of $S$ to a random $k$-dimensional subspace.)
Different constructions for $R$

- Frankl & Maehara (1988): random orthogonal matrix
- DasGupta & Gupta (1999): matrix with entries from $\mathcal{N}(0,1)$, normalized
- Indyk & Motwani (1998): matrix with entries from $\mathcal{N}(0,1)$
- **Achlioptas (2003)**: matrix with entries in $\{-1,0,+1\}$
- Alon (2003): optimal dependency on $n$, and almost optimal dependency on $\varepsilon$

Construct an $n$-by-$s$ matrix $R$ such that:

$$R_{ij} = \sqrt{3} \times \begin{cases} +1, \text{w.p. } 1/6 \\ 0, \text{w.p. } 2/3 \\ -1, \text{w.p. } 1/6 \end{cases}$$

Return: $\tilde{A} = \frac{1}{\sqrt{s}}AR \in \mathbb{R}^{m \times s}$

$O(mns) = O(mn \log m / \varepsilon^2)$ time computation
Fast JL transform

\[ P \in \mathbb{R}^{s \times n} \]
\[ s = O\left(\log m/\epsilon^2\right) \]

\[ H \in \mathbb{R}^{n \times n} \]

\[ D \in \mathbb{R}^{n \times n} \]

\[ R = (PHD)^T \in \mathbb{R}^{n \times s} \]

\[ P_{ij} = \sqrt{q} \times \begin{cases} +1, & \text{w.p. } q/2 \\ 0, & \text{w.p. } 1-q \\ -1, & \text{w.p. } q/2 \end{cases} \]

\[ q = O\left(\frac{\log^2 m}{n}\right) \]

Normalized Hadamard-Walsh transform matrix
(if \( n \) is not a power of 2, add all-zero columns to \( A \))

Diagonal matrix with \( D_{ii} \) set to +1 or -1 w.p. \( \frac{1}{2} \).

\[ \tilde{A} = \frac{1}{\sqrt{s}} AR \]
Applying PHD on a vector $u \in \mathbb{R}^n$ is fast, since:

- $Du : O(n)$, since $D$ is diagonal,

- $H(Du) : O(n \log n)$, using the Hadamard-Walsh algorithm,

- $P(H(Du)) : O(\log^3 m/\varepsilon^2)$, since $P$ has on average $O(\log^2 n)$ non-zeros per row (in expectation).
Let $A$ by an $m$-by-$n$ matrix whose SVD is:

$$A = U \Sigma V^T \in \mathbb{R}^{m \times n}$$

Apply the (HD) part of the (PHD) transform to $A$.

$$ADH = U \Sigma (V^T DH) \in \mathbb{R}^{m \times n}$$

**Observations:**

1. The left singular vectors of $ADH$ span the same space as the left singular vectors of $A$.
2. The matrix $ADH$ has (up to log $n$ factors) uniform leverage scores.
   (Thanks to $V^T DH$ having bounded entries - the proof closely follows JL-type proofs.)
3. We can approximate the left singular vectors of $ADH$ (and thus the left singular vectors of $A$) by uniformly sampling columns of $ADH$.  

 orthogonal matrix
Back to approximating singular vectors

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3. We can approximate the left singular vectors of $ADH$ (and thus the left singular vectors of $A$) by uniformly sampling columns of $ADH$.
4. The orthonormality of $HD$ and a version of our relative-error Frobenius norm bound (involving approximately optimal sampling probabilities) suffice to show that (w.h.p.)

$$\left\| A - \tilde{C} \tilde{C}^T A \right\|_F \leq (1 + \epsilon) \left\| A - A_k \right\|_F$$

Uniform sample of $s = O \left( \frac{k}{\epsilon^2 \log^{c_0} n} \right)$ columns of $ADH$. 
Running time

Let $A$ be an $m$-by-$n$ matrix whose SVD is:

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Apply the (HD) part of the (PHD) transform to $A$.

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**Orthogonal matrix**

**Running time:**

1. **Trivial analysis:** first, uniformly sample $s$ columns of $DH$ and then compute their product with $A$.
   Takes $O(mns) = O(mnk \text{ polylog}(n))$ time, already better than full SVD.

2. **Less trivial analysis:** take advantage of the fact that $H$ is a Hadamard-Walsh matrix.
   Improves the running time $O(mn \text{ polylog}(n) + mk^2 \text{ polylog}(n))$. 
Conclusions

- Randomization and sampling can be used to solve problems that are massive and/or computationally expensive.

- By (carefully) sampling rows/columns/entries of a matrix, we can construct new sparse/smaller matrices that behave like the original matrix.

  - Can entry-wise sampling be made competitive to column-sampling in terms of accuracy and speed? See Achlioptas and McSherry (2001) STOC, (2007) JACM.
  - We improved/generalized/simplified it. See Nguyen, Drineas, & Tran (2011), Drineas & Zouzias (2010).
  - Exact reconstruction possible using uniform sampling for constant-rank matrices that satisfy certain (strong) assumptions. See Candes & Recht (2008), Candes & Tao (2009), Recht (2009).

- By preprocessing the matrix using random projections, we can sample rows/columns much less carefully (even uniformly at random) and still get nice “behavior”.
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