Randomized algorithms for least squares and matrix approximation

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About me

I work on the design and analysis of practically fast and theoretically justified randomized algorithms for fundamental linear algebra tasks.

I like working with nearby communities (physics, theoretical computer science, optimization, computational science, etc.).

Academic history:

- Currently an Assistant Professor / Courant Instructor at New York University
 - Sponsor: Chris Musco
- PhD in Applied Math at University of Washington
 - Advisors: Anne Greenbaum and Tom Trogdon
- B.S. in Math and Physics at Tufts University, minor in Studio Art

Today

Stochastic Optimization¹

– First proof of $O(\sqrt{\kappa})$ convergence of minibatch stochastic gradient descent with heavy-ball momentum

Matrix approximation/operator learning²

- First matrix-vector product algorithm for near-optimal hierarchical matrix approximation
- Matrix-vector product algorithms for approximating fixed-sparsity matrices (no dimension dependence and matching lower bounds!)

¹Bollapragada, Chen, and Ward 2024.

²Amsel, Chen, Keles, Halikias, Musco, and Musco 2024; Chen, Keles, Halikias, Musco, Musco, and Persson 2024.

Linear least squares

Consider a consistent least squares problem

$$\min_{\mathbf{x} \in \mathbb{R}^d} f(\mathbf{x}), \qquad f(\mathbf{x}) = \frac{1}{2} \|\mathbf{A}\mathbf{x} - \mathbf{b}\|^2 = \sum_{i=1}^n \frac{1}{2} (\mathbf{a}_i^\mathsf{T} \mathbf{x} - b_i)^2.$$

The gradient is

$$\nabla f(\mathbf{x}) = \mathbf{A}^{\mathsf{T}}(\mathbf{A}\mathbf{x} - \mathbf{b}) = \sum_{i=1}^{n} \mathbf{a}_{i}(\mathbf{a}_{i}^{\mathsf{T}}\mathbf{x} - b_{i}).$$

We can then implement gradient descent

$$\mathbf{x}_{k+1} = \mathbf{x}_k - \alpha_k \nabla f(\mathbf{x}_k).$$

This requires one product with each A and A^T every iteration.

Improvements to gradient descent

Stochastic gradients: Often we don't want to read the whole data matrix each iteration. Instead, we sample a random index i with probability $p_i \propto \|\mathbf{a}_i\|^2$, and then use the stochastic gradient

$$\nabla f_i(\mathbf{x}) = \frac{1}{p_i} \mathbf{a}_i (\mathbf{a}_i^{\mathsf{T}} \mathbf{x} - b_i).$$

The stochastic gradient is equal to the true gradient on average:

$$\mathbb{E}[\nabla f_i(\mathbf{x})] = \sum_{i=1}^n p_i \cdot \frac{1}{p_i} \mathbf{a}_i (\mathbf{a}_i^\mathsf{T} \mathbf{x} - b_i) = \sum_{i=1}^n \mathbf{a}_i (\mathbf{a}_i^\mathsf{T} \mathbf{x} - b_i) = \nabla f(\mathbf{x}).$$

Momentum: Instead of just updating based on the gradient, we can take previous iterates into account.

L

Both momentum and stochastic gradient improve on classical gradient descent.

algorithm	iterations	cost/iter	formula
Gradient Descent	$\lambda_{ m max}/\lambda_{ m min}$	nd	$\mathbf{x}_{k+1} = \mathbf{x}_k - \alpha \nabla f(\mathbf{x}_k)$
Momentum	$\sqrt{\lambda_{\mathrm{max}}/\lambda_{\mathrm{min}}}$	nd	$\mathbf{x}_{k+1} = \mathbf{x}_k - \alpha \nabla f(\mathbf{x}_k) + \beta(\mathbf{x}_k - \mathbf{x}_{k-1})$
Stochastic Gradient	$n \cdot (\lambda_{\rm ave}/\lambda_{\rm min})$	d	$\mathbf{x}_{k+1} = \mathbf{x}_k - \alpha \nabla f_i(\mathbf{x}_k)$

However, the bounds are somewhat incomparable

- average condition number vs. square root of condition number
- the cost per iteration is not necessarily indicative of real-world costs
- sampling assumptions

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Key question:

What is the role of momentum in SGD?

Minibatch gradient descent

In practice: momentum is used with stochastic methods and plays a critical role in training deep learning models!

In theory: momentum cannot be used to accelerate SGD.³

Minibatching: Instead of sampling a single index, we can sample several *B* indices at once and put them into a list *S*. This gives the minibatch stochstic gradient

$$\nabla f_{S}(\mathbf{x}) = \frac{1}{B} \sum_{i \in S} \frac{1}{p_{i}} \mathbf{a}_{i} (\mathbf{a}_{i}^{\mathsf{T}} \mathbf{x} - b_{i}).$$

One minibatch evaluation costs O(Bd) operations.⁴

³Jain, Kakade, Kidambi, Netrapalli, and Sidford 2018.

⁴Due to parallelism, the real-world cost is often sort of independent of *B* when *B* is reasonably small.

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Minibatching+momentum

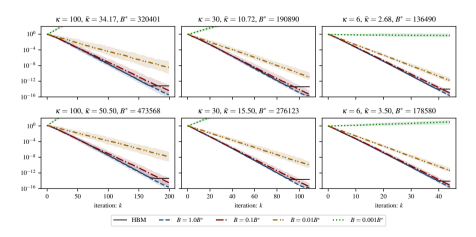
Theorem (Bollapragada, Chen, and Ward 2024). Minibatch heavy ball momentum converges in $\sqrt{\lambda_{\max}/\lambda_{\min}} \cdot \log(1/\varepsilon)$ iterations with minibatch size $B = O(d \log(d) \cdot (\lambda_{\text{ave}}/\lambda_{\min}) \cdot \sqrt{\lambda_{\max}/\lambda_{\min}})$.

Minibatch SGD can get the best of both worlds!

- same rate of convergence as full gradient methods
- cheaper gradients computations when n is large

Numerical example

Our theory might be precise enough to predict practical performance.



Let's first look at how to analyze the classical heavy ball momentum algorithm:

$$\mathbf{x}_{k+1} = \mathbf{x}_k - \alpha \underbrace{(\mathbf{A}^\mathsf{T} \mathbf{A} \mathbf{x}_k - \mathbf{A}^\mathsf{T} \mathbf{b})}_{\text{gradient}} + \beta \underbrace{(\mathbf{x}_k - \mathbf{x}_{k-1})}_{\text{momentum}}.$$

Assuming a consistent system, $\mathbf{b} = \mathbf{A}\mathbf{x}^*$, so we can rewrite the update as

$$\mathbf{x}_{k+1} = \mathbf{x}_k - \alpha \mathbf{A}^{\mathsf{T}} \mathbf{A} (\mathbf{x}_k - \mathbf{x}^*) + \beta (\mathbf{x}_k - \mathbf{x}_{k-1}).$$

Therefore,

$$\mathbf{x}_{k+1} - \mathbf{x}^* = (\mathbf{I} - \alpha \mathbf{A}^\mathsf{T} \mathbf{A})(\mathbf{x}_k - \mathbf{x}^*) + \beta(\mathbf{x}_k - \mathbf{x}^* + \mathbf{x}^* - \mathbf{x}_{k-1})$$
$$= ((1+\beta)\mathbf{I} - \alpha \mathbf{A}^\mathsf{T} \mathbf{A})(\mathbf{x}_k - \mathbf{x}^*) - \beta(\mathbf{x}_k - \mathbf{x}_{k-1}).$$

We can write this as a matrix iteration

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It's straightforward (but a bit tedious) to show that the eigenvalues of T are

$$z_j^\pm := rac{1}{2} \left(1 + eta - lpha \lambda_j \pm \sqrt{(1 + eta - lpha \lambda_j)^2 - 4eta} \,
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Finally, optimizing the choices of α and β , we can get a bound

$$\|\mathbf{T}^k\| \approx \|\mathbf{T}\|^k = \left(\frac{\sqrt{\lambda_{\max}/\lambda_{\min}} - 1}{\sqrt{\lambda_{\max}/\lambda_{\min}} + 1}\right)^k \leq \exp\left(-\frac{k}{2\sqrt{\lambda_{\max}/\lambda_{\min}}}\right)$$

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Proof overview

In minibatch-HBM, we replace the true gradient with a minibatch stochastic gradient

$$\nabla f_{S_k}(\mathbf{x}_k) = \frac{1}{B} \sum_{j \in S_k} \frac{1}{p_j} \mathbf{a}_j \mathbf{a}_j^{\mathsf{T}}(\mathbf{x}_k - \mathbf{x}^*).$$

Define the random matrix

$$\mathbf{M}_{S_k} = \frac{1}{B} \sum_{j \in S_k} \frac{1}{p_j} \mathbf{a}_j \mathbf{a}_j^{\mathsf{T}}$$

and observe that

$$\mathbb{E}[\mathbf{M}_{S_k}] = \frac{1}{B} \mathbb{E} \left[\sum_{j \in S_k} \frac{1}{p_j} \mathbf{a}_j \mathbf{a}_j^{\mathsf{T}} \right] = \mathbf{A}^{\mathsf{T}} \mathbf{A}.$$

A similar argument gives us a recurrence

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This gives an error bound

$$\left\| \begin{bmatrix} \mathbf{x}_{k+1} - \mathbf{x}^* \\ \mathbf{x}_k - \mathbf{x}^* \end{bmatrix} \right\| \le \left\| \mathbf{Y}_{S_k} \mathbf{Y}_{S_{k-1}} \cdots \mathbf{Y}_{S_1} \right\| \left\| \begin{bmatrix} \mathbf{x}_1 - \mathbf{x}^* \\ \mathbf{x}_0 - \mathbf{x}^* \end{bmatrix} \right\|.$$

We might hope that

$$\|\mathbf{Y}_{S_k}\mathbf{Y}_{S_{k-1}}\cdots\mathbf{Y}_{S_1}\|\approx \|\mathbb{E}[\mathbf{Y}_{S_k}]\mathbb{E}[\mathbf{Y}_{S_{k-1}}]\cdots\mathbb{E}[\mathbf{Y}_{S_1}]\|=\|\mathbf{T}^k\|.$$

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Products of random matrices

Theorem (Huang, Niles-Weed, Tropp, and Ward 2021). Consider an independent sequence of $d \times d$ random matrices $\mathbf{X}_1, \dots, \mathbf{X}_k$, and form the product

$$\mathbf{Z} = \mathbf{X}_k \mathbf{X}_{k-1} \cdots \mathbf{X}_1.$$

Assume $\|\mathbb{E}[\mathbf{X}_i]\| \le q_i$ and $\mathbb{E}[\|\mathbf{X}_i - \mathbb{E}\mathbf{X}_i\|^2]^{1/2} \le \sigma_i q_i$ for i = 1, ..., k. Let $Q = \prod_{i=1}^n q_i$ and $v = \sum_{i=1}^k \sigma_i^2$. Then

$$\mathbb{E}[\|\mathbf{Z}\|] \le Q \exp\left(\sqrt{2\nu \max\{2\nu, \log(d)\}}\right).$$

To determine q_i and σ_i we we use matrix concentration bounds for sums of matrices⁵.

⁵Tropp 2015.

Outlook

Lots of interesting potential follow up work:

- adaptive/automatic parameter selection
- beyond least squares

Key question:

What can we learn about matrices from matvecs?

In this model of computation, we are only given access to A via a black box that lets us perform matrix-vector products $\mathbf{x} \mapsto \mathbf{A}\mathbf{x}$ and $\mathbf{x} \mapsto \mathbf{A}^T\mathbf{x}$.

- $-\mathbf{A} = \mathbf{B}^{-1}$, and we only know \mathbf{B}
 - perform products by solving $\mathbf{B}\mathbf{y} = \mathbf{x}$
 - this generalizes to $\mathbf{A} = f(\mathbf{B})$
- $-\mathbf{A} = \exp(-\mathrm{i}t\mathbf{H})$
 - perform products by evolving according to Hamiltonian H for time t from initial state x
- A is the system matrix for a CT scanner
 - perform products by running the CT scanner on target x
- A is a regular matrix
 - perform products with highly optimized hardware/software (e.g. GPUs, crossbar array, MKL, etc.)

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 - perform products by solving $\mathbf{B}\mathbf{y} = \mathbf{x}$
 - this generalizes to $\mathbf{A} = f(\mathbf{B})$
- $-\mathbf{A} = \exp(-\mathrm{i}t\mathbf{H})$
 - perform products by evolving according to Hamiltonian H for time t from initial state x
- A is the system matrix for a CT scanner
 - perform products by running the CT scanner on target x
- A is a regular matrix
 - perform products with highly optimized hardware/software (e.g. GPUs, crossbar array, MKL, etc.)

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The computational complexity of linear algebra

Numerical linear algebra is one of the oldest fields of computing.

- Traditionally, the cost of algorithms is studied in terms of floating point operations, but flops are increasingly less relevant.
- Now, things like matrix products (or matrix loads) often dominate the cost of an algorithm.

If we take matrix-products as the basic unit of computation, we can also study the computational complexity of linear algebra problems.

– almost all linear algebra problems can be solved in $O(n^3)$ time, so classical complexity theory (e.g. P vs NP) is not very useful

Structrured matrices

Often, we can solved linear algebra problems faster when the matrices of interest are structured.

Examples:

- low-rank matrices
- hierarchical matrices
- sparse matrices
- banded matrices

Common framework: Approximate **A** with a structured matrix, then use the structured approximation.

Matrix approximation

Problem. Let S be some family of matrices and A an arbitrary matrix that can only be accessed by matrix-vector products.

Find a matrix $\widetilde{\mathbf{A}} \in S$ such that

$$\|\mathbf{A} - \widetilde{\mathbf{A}}\| \le (1 + \varepsilon) \min_{\mathbf{X} \in S} \|\mathbf{A} - \mathbf{X}\|.$$

This problem asks us to find a structured matrix approximation $\widetilde{\mathbf{A}}$ to \mathbf{A} competitive with the best possible approximation.

- If $A \in S$, then we require $\widetilde{A} = A$
- If **A** has a good approximation from *S*, then it might be okay for ε to be large

Wxample: low-rank approximation⁶

⁶Halko, Martinsson, and Tropp 2011; Musco and Musco 2015; Tropp and Webber 2023.

Hierarchical matrix motivation: kernel matrices

Consider data $\mathbf{x}_1, \dots, \mathbf{x}_n \in \mathbb{R}^d$ and a kernel function $k(\mathbf{x}_i, \mathbf{x}_i)$.

- examples:
$$k(\mathbf{x}, \mathbf{y}) = 1/\|\mathbf{x} - \mathbf{y}\|^2$$
, $k(\mathbf{y}) = \exp(-\|\mathbf{x} - \mathbf{y}\|^2/\sigma^2)$, $k(\mathbf{x}, \mathbf{y}) = (\mathbf{x}^{\mathsf{T}}\mathbf{y} + c)^q$

We can define a kernel matrix

$$\mathbf{K} = \begin{bmatrix} k(\mathbf{x}_1, \mathbf{x}_1) & \cdots & k(\mathbf{x}_1, \mathbf{x}_n) \\ \vdots & \ddots & \vdots \\ k(\mathbf{x}_n, \mathbf{x}_1) & \cdots & k(\mathbf{x}_n, \mathbf{x}_n) \end{bmatrix}.$$

Kernel matrices are widely used in data science and machine learning applications for task such as clustering and classification.⁷

⁷?.

Hierarchical matrix motivation: kernel matrices

For simplicity, consider 1 dimensional data:

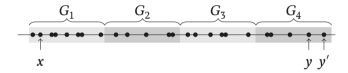


Observations:

- Interactions between points in non-adjacent groups are approximately low-rank.
- We can recursively treat interactions between adjacent groups or within a group at a finer scale.

Hierarchical matrix motivation: kernel matrices

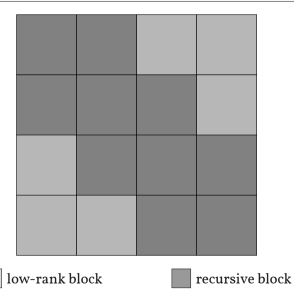
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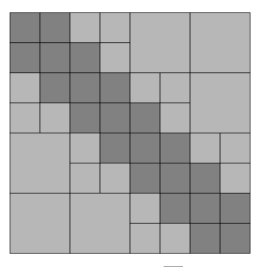
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Hierarchical matrices



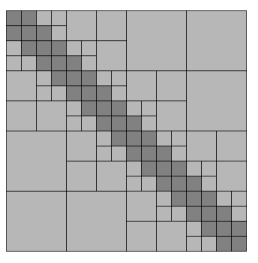
Hierarchical matrices



low-rank block

recursive block

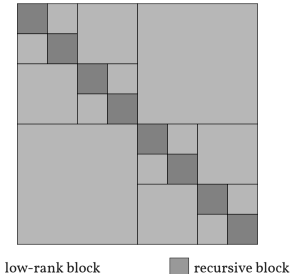
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HODLR matrices



The HODLR approximation problem

Problem. Given an $n \times n$ matrix **A**, accessible only by matrix-vector products, a rank parameter k, and an accuracy parameter ε , find a HODLR(k) matrix $\widetilde{\mathbf{A}}$ such that

$$\mathbb{E}\big[\| \mathbf{A} - \widetilde{\mathbf{A}} \|_{\mathsf{F}} \big] \leq \big(1 + \varepsilon \big) \min_{\mathbf{H} \in \mathrm{HODLR}(k)} \| \mathbf{A} - \mathbf{H} \|_{\mathsf{F}}.$$

The best HODLR approximation to **A** is obtained by applying a rank-k SVD to each low-rank block of **A**.

This is too expensive in the matrix-vector product model (n products)

In the special case that $A \in HODLR(k)$, then we require $\widetilde{A} = A$ (regardless of ε).

There are several matvec algorithms for this setting⁸

⁸Lin, Lu, and Ying 2011; Martinsson 2016; Levitt and Martinsson 2022; Halikias and Townsend 2023.

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Learning low-rank matrices from matrix-vector products

The Randomized SVD (RSVD) is a well-known algorithm for obtaining a low-rank approximation to a matrix ${\bf B}$:

- 1. Sample Gaussian matrix Ω
- 2. Form $\mathbf{Q} = \operatorname{orth}(\mathbf{B}\mathbf{\Omega})$
- 3. Compute $\mathbf{X} = \mathbf{Q}^{\mathsf{T}}\mathbf{B}$
- 4. Output $\mathbf{Q}[X]_k$

Theorem. If Ω has $O(k/\varepsilon)$ columns, then

$$\|\mathbf{B} - \mathbf{Q}[\![\mathbf{X}]\!]_k\|_{\mathsf{F}} \leq (1+\varepsilon) \min_{\mathrm{rank}(\mathbf{X}) \leq k} \|\mathbf{B} - \mathbf{X}\|_{\mathsf{F}}.$$

Corollary. If **B** is rank-k, then $\mathbb{Q}[\![X]\!]_k = \mathbb{B}$ (with probability one).

Peeling: an algorithm for the recovery problem⁹

The algorithm works from the top layer down.

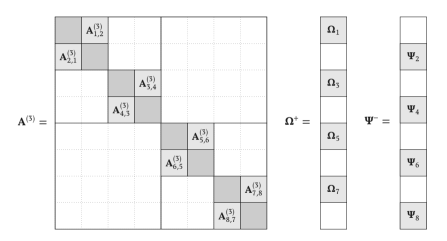
At each level, we simultaneosly apply the RSVD to the low-rank off-diagonal blocks.

We then "peel" off these blocks before proceeding to the next level

⁹Lin, Lu, and Ying 2011; Martinsson 2016.

2

Peeling: an algorithm for the recovery problem



Peeling: an algorithm for the recovery problem

At each level we use k matrix-vector products with \mathbf{A} and \mathbf{A}^{T} .

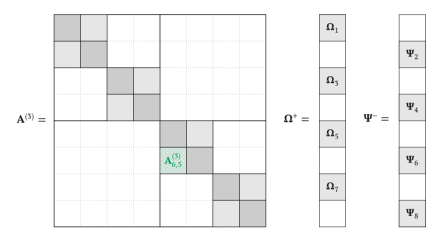
There are $\log_2(n/k) \le \log_2(n)$ levels until the blocks are of size k

- then we can directly recover them at once with k products

Theorem (Lin, Lu, and Ying 2011). There is an algorithm to recover a HODLR matrix using $O(k \log_2(n))$ products with **A**.

If **A** is not HODLR, we can still apply this algorithm, but we might worry about exponential blow-up in the error!

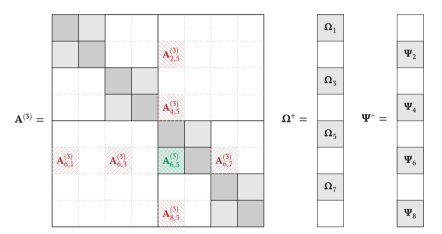
Peeling with error



We obtain perturbed sketches of the form $\mathbf{A}_{6,5}^{(3)} \mathbf{\Omega}_5 + \mathbf{A}_{6,1}^{(3)} \mathbf{\Omega}_1 + \mathbf{A}_{6,3}^{(3)} \mathbf{\Omega}_3 + \mathbf{A}_{6,7}^{(3)} \mathbf{\Omega}_7$ and $(\mathbf{A}_{6,5}^{(3)})^\mathsf{T} \mathbf{\Psi}_6 + (\mathbf{A}_{2,5}^{(3)})^\mathsf{T} \mathbf{\Psi}_2 + (\mathbf{A}_{4,5}^{(3)})^\mathsf{T} \mathbf{\Psi}_4 + (\mathbf{A}_{8,5}^{(3)})^\mathsf{T} \mathbf{\Psi}_8$.

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Peeling with error



We obtain perturbed sketches of the form $\mathbf{A}_{6,5}^{(3)} \mathbf{\Omega}_5 + \mathbf{A}_{6,1}^{(3)} \mathbf{\Omega}_1 + \mathbf{A}_{6,3}^{(3)} \mathbf{\Omega}_3 + \mathbf{A}_{6,7}^{(3)} \mathbf{\Omega}_7$ and $(\mathbf{A}_{6,5}^{(3)})^\mathsf{T} \mathbf{\Psi}_6 + (\mathbf{A}_{2,5}^{(3)})^\mathsf{T} \mathbf{\Psi}_2 + (\mathbf{A}_{4,5}^{(3)})^\mathsf{T} \mathbf{\Psi}_4 + (\mathbf{A}_{8,5}^{(3)})^\mathsf{T} \mathbf{\Psi}_8$.

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Generalized Nyström¹⁰

The RSVD tries to compute $\mathbf{Q}^{\mathsf{T}}\mathbf{B}$ directly; this is the solution to:

$$\min_{X}\|A-QX\|_{\text{F}}.$$

Instead, we can solve a sketched problem:

$$\min_{\boldsymbol{X}} \| \boldsymbol{\Psi}^{\mathsf{T}} \boldsymbol{A} - \boldsymbol{\Psi}^{\mathsf{T}} \boldsymbol{Q} \boldsymbol{X} \|_{\mathsf{F}}.$$

This means $\mathbf{X} = (\mathbf{\Psi}^{\mathsf{T}} \mathbf{Q})^{\dagger} \mathbf{\Psi}^{\mathsf{T}} \mathbf{A}$.

Observation. By adding columns to Ψ , we can average out certain errors in the product $\Psi^T A$.

The algorithm is also non-adaptive (we can do products with Ψ in advance)

¹⁰Clarkson and Woodruff 2009; Tropp, Yurtsever, Udell, and Cevher 2017; Nakatsukasa 2020.

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A perturbation bound for the RSVD

We prove a perturbation bound for the RSVD. This is likely of independent interest.

Theorem (Chen, Keles, Halikias, Musco, Musco, and Persson 2024). Let $\mathbf{Q} = \operatorname{orth}(\mathbf{B}\mathbf{\Omega} + \mathbf{E}_1)$ and $\mathbf{X} = \mathbf{Q}^T\mathbf{B} + \mathbf{E}_2$. Then

$$\|\mathbf{B} - \mathbf{Q} \|\mathbf{Q}^\mathsf{T} \mathbf{B} + \mathbf{E}_2\|_k\|_\mathsf{F} \leq \underbrace{\|\mathbf{E}_1 \mathbf{\Omega}_\mathsf{top}^\dagger\|_\mathsf{F} + 2\|\mathbf{E}_2\|_\mathsf{F}}_{\text{perturbations}} + \underbrace{\left(\|\mathbf{\Sigma}_\mathsf{bot}\|_\mathsf{F}^2 + \|\mathbf{\Sigma}_\mathsf{bot} \mathbf{\Omega}_\mathsf{bot}^\dagger \mathbf{\Omega}_\mathsf{top}^\dagger\|_\mathsf{F}^2\right)^{1/2}}_{\text{classical RSVD bound}}.$$

Takeaway: The pseudoinverse will help damp the perturbation \mathbf{E}_1 , but (unsurprisingly) all of the perturbation \mathbf{E}_2 can propagate.

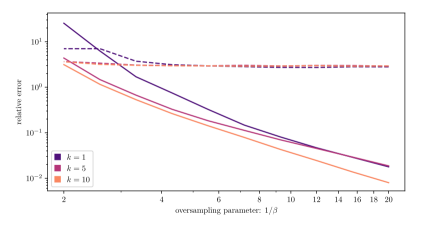
Matrix approximation

Theorem (Chen, Keles, Halikias, Musco, Musco, and Persson 2024). There exist matvec algorithms which use $O(k \log(n) \cdot \operatorname{poly}(1/\beta))$ products with **A** to obtain a HODLR(k) matrix $\widetilde{\mathbf{A}}$ satisfying

$$\|\mathbf{A} - \widetilde{\mathbf{A}}\|_{\mathsf{F}} \leq (1 + \beta)^{\log_2(n)} \min_{\mathbf{H} \in \mathrm{HODLR}(k)} \|\mathbf{A} - \mathbf{H}\|_{\mathsf{F}}.$$

 $\textbf{Corollary.} \ (1+\varepsilon) \text{-optimal approximation with } O\big(k\log(n) \cdot \operatorname{poly}(\log(n)/\varepsilon)\big) \ \text{matvecs}$

Corollary. $n^{0.01}$ -optimal approximation with $O(k \log(n))$ matvecs



 $solid = Nystr\"{o}m~based~algorithm, ~~dashed = RSVD~based~algorithm$

Outlook

- This is the first algorithm which can produce a near-optimal hierarchical approximation from matvecs.
- We studied HODLR, mostly for convenience. The same ideas extend to H^1 , but less clear about HSS or H^2 .
- A number of interesting questions about low-rank approximation via Nyström

The fixed-sparsity approximation problem

Fix a $n \times n$ binary matrix **S**, and consider the set of matrices whose nonzero pattern is given by **S**; i.e. $S = \{X : X \circ S = X\}$, where " \circ " is the entrywise product.

- examples: diagonal matrices, banded matrices

Problem. Given an $n \times n$ matrix **A**, accessible only by matrix-vector products, find a matrix $\widetilde{\mathbf{A}}$ with sparsity \mathbf{S} such that

$$\|\mathbf{A}-\widetilde{\mathbf{A}}\|_{\text{F}} \leq \left(1+\epsilon\right) \min_{\mathbf{X} \circ \mathbf{S} = \mathbf{X}} \|\mathbf{A} - \mathbf{X}\|_{\text{F}}.$$

The best sparse approximation to **A** is $\mathbf{A} \circ \mathbf{S}$.

- This is too expensive in the matrix-vector product model (n products)

Lots of past work for exact recovery¹¹ and diagonal estimation¹²

¹¹Coleman and Moré 1983; Coleman and Moré 1983.

¹²Baston and Nakatsukasa 2022; Hallman, Ipsen, and Saibaba 2023; Dharangutte and Musco 2023.

The fixed-sparsity approximation problem: an algorithm

One might try a simple sketching algorithm:

$$\widetilde{\mathbf{A}} = \mathop{\mathrm{argmin}}_{X \circ S = X} \|\mathbf{A} \mathbf{G} - \mathbf{X} \mathbf{G}\|_{\text{F}}.$$

Theorem (Amsel, Chen, Keles, Halikias, Musco, and Musco 2024). Suppose **S** has at most s nonzeros per row. There exists a matvec algorithms which use $O(s/\varepsilon)$ products with **A** to obtain a matrix $\widetilde{\mathbf{A}}$ with sparsity **S** satisfying

$$\mathbb{E}[\|\mathbf{A}-\widetilde{\mathbf{A}}\|_{\text{F}}] \leq \left(1+\epsilon\right) \min_{\mathbf{X} \circ \mathbf{S} = \mathbf{X}} \|\mathbf{A} - \mathbf{X}\|_{\text{F}}.$$

Unlike compressed-sensing type problems, no dimension dependence!

– in the compressed sensing setting, we have a necessary $\log(n)$ factor

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The fixed-sparsity approximation problem (hardness)

We also prove a lower-bound, which reveals that our algorithm is optimal, up to constant factors.

Theorem (Amsel, Chen, Keles, Halikias, Musco, and Musco 2024). For any **S** with $\Theta(s)$ nonzeros per row/column, there exist hard instances for which any algorithm producing $\widetilde{\mathbf{A}}$ with sparsity **S** must use $\Omega(s/\varepsilon)$ products with \mathbf{A} in order to guarantee

$$\mathbb{P}\bigg[\left\|\mathbf{A} - \widetilde{\mathbf{A}}\right\|_{\mathsf{F}} \le (1 + \varepsilon) \min_{\mathbf{X} \circ \mathbf{S} = \mathbf{X}} \left\|\mathbf{A} - \mathbf{X}\right\|_{\mathsf{F}}\bigg] > 1/2.$$

3

Lower bound case study: diagonal approximation

To illustrate the key ideas, let's consider the special case $\mathbf{S} = \mathbf{I}$; i.e. diagonal approximation.

Theorem. There exists a constant C > 0 such that the following holds:

For any $\varepsilon > 0$, there there exists a distribution ${\bf A}$ on $d \times d$ matrices such that, any algorithm which uses $m < C/\varepsilon$ queries cannot output an approximation ${\bf d}$ to diag(${\bf A}$) satisfying

$$\mathbb{P}\big[\|\operatorname{diag}(\mathbf{A}) - \mathbf{d}\|_2^2\big] \le \varepsilon \|\mathbf{A} - \mathbf{A} \circ \mathbf{I}\|_{\mathsf{F}}^2\big] > 1/2.$$

If $\widetilde{\mathbf{A}} = \operatorname{diag}(\mathbf{d})$ and the event in the probability holds,

$$\|\mathbf{A} - \widetilde{\mathbf{A}}\|_{\mathsf{F}}^2 = \|\mathbf{A} \circ \mathbf{S} - \widetilde{\mathbf{A}}\|_{\mathsf{F}}^2 + \|\mathbf{A} - \mathbf{A} \circ \mathbf{I}\|_{\mathsf{F}}^2 \le (1 + \varepsilon)\|\mathbf{A} - \mathbf{A} \circ \mathbf{I}\|_{\mathsf{F}}^2.$$

Since $\sqrt{1+\varepsilon} = 1 + O(\varepsilon)$, this is also a lower bound for the approximation problem.

Gaussian matrices after adaptive queries

Lemma. Let $\mathbf{A} \sim \operatorname{Gaussian}(d,d)$. Suppose we make a sequence of adaptive queries $\mathbf{x}_1,\ldots,\mathbf{x}_m$ with responses $\mathbf{y}_1=\mathbf{A}\mathbf{x}_1,\ldots,\mathbf{y}_m=\mathbf{A}\mathbf{x}_m$. Let $\mathbf{X}=[\mathbf{x}_1,\ldots,\mathbf{x}_m]$ and $\mathbf{Y}=[\mathbf{y}_1,\ldots,\mathbf{y}_m]$. Then, \mathbf{A} can be factored as

$$\mathbf{A} = \begin{bmatrix} \mathbf{Y} & \mathbf{G} \end{bmatrix} \begin{bmatrix} \mathbf{X}^\mathsf{T} \\ \mathbf{Z}^\mathsf{T} \end{bmatrix},$$

where $\mathbf{Z}^{\mathsf{T}}\mathbf{X} = \mathbf{0}$, $\mathbf{Z}^{\mathsf{T}}\mathbf{Z} = \mathbf{I}$, and $\mathbf{G} \sim \operatorname{Gaussian}(d, d - m)$, independently of \mathbf{X} and \mathbf{Y} .

Conditioned on the queries and responses, A still has a lot of randomness!

Analysis

We have that

$$\mathbf{A} = \mathbf{Y}\mathbf{X}^\mathsf{T} + \mathbf{G}\mathbf{Z}^\mathsf{T},$$

where $\mathbf{Z}^{\mathsf{T}}\mathbf{X} = \mathbf{0}$, $\mathbf{Z}^{\mathsf{T}}\mathbf{Z} = \mathbf{I}$, and $\mathbf{G} \sim \operatorname{Gaussian}(d, d - m)$, independently of \mathbf{X} and \mathbf{Y} .

Conditioned on the queries and responses,

$$\mathbf{A}_{i,i} = \mathbf{e}_i^\mathsf{T} \mathbf{Y} \mathbf{X}^\mathsf{T} \mathbf{e}_i + \mathbf{e}_i^\mathsf{T} \mathbf{G} \mathbf{Z}^\mathsf{T} \mathbf{e}_i = \mathrm{determistic} + \mathbf{g}_i^\mathsf{T} \mathbf{z}_i.$$

Observe that the \mathbf{g}_i are all independent Gaussian vectors of length d-m. Hence

$$d_i = \mathbf{g}_i^{\mathsf{T}} \mathbf{z}_i \sim N(0, \|\mathbf{z}_i\|^2)$$
 independently.

Suppose we try to output an approximation \mathbf{d} to diag(\mathbf{A}). Based on what we know (queries and responses), the best thing we can do is output $\mathbf{d} = \text{diag}(\mathbf{Y}\mathbf{X}^{\mathsf{T}})$. So then

$$\mathbb{E}[\|\operatorname{diag}(\mathbf{A}) - \mathbf{d}\|_{2}^{2}] = \sum_{i=1}^{n} \mathbb{E}[d_{i}^{2}] = \sum_{i=1}^{n} \|\mathbf{z}_{i}\|^{2} = \|\mathbf{Z}\|_{F}^{2} = d - m.$$

Choosing the parameters

Applying concentration inequality,

$$\mathbb{P}\big[\|\operatorname{diag}(\mathbf{A}) - \mathbf{d}\|_2^2 > 100(d - m)\big] \le \text{small.}$$

In addition,

$$\mathbb{P}\big[\|\mathbf{A} - \mathbf{A} \circ \mathbf{I}\|_{\mathsf{F}}^2 < d^2/100\big] \le \text{small}.$$

If we set $d = 5000/\varepsilon$ and $m < 2500/\varepsilon$,

$$100(d-m) > 100d/2 = \varepsilon d^2/100.$$

Therefore, by a union bound,

$$\mathbb{P}\big[\,\|\,\mathrm{diag}(\mathbf{A}) - \mathbf{d}\|_2^2 < \varepsilon\|\mathbf{A} - \mathbf{A} \circ \mathbf{I}\|_{\mathsf{F}}^2\big] \le \mathsf{small}.$$

Conclusion

We saw two broad directions in randomized linear algebra that I have worked on.

I'm also intereted in many other problems including (but certainly not limited to):

- (partial) trace approximation
- spectrum approximation
- classical Krylov subspace methods

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Quantum equilibrium thermodynamics

Consider a quantum system consisting of subsystems (s) and (b) with Hamiltonian

$$\mathbf{H} = \bar{\mathbf{H}}_{s} + \bar{\mathbf{H}}_{b} + \mathbf{H}_{sb}, \qquad \bar{\mathbf{H}}_{s} = \mathbf{H}_{s} \otimes \mathbf{I}_{b}, \quad \bar{\mathbf{H}}_{b} = \mathbf{I}_{s} \otimes \mathbf{H}_{b}. \tag{1}$$

In thermal equilibrium at interver temperature β , the state of the system is described by a density matrix

$$\rho_{t}(\beta) = \frac{\exp(-\beta \mathbf{H})}{Z_{t}(\beta)}, \qquad Z_{t}(\beta) = \operatorname{tr}(\exp(-\beta \mathbf{H}); \tag{2}$$

The denisty matrix for subsystem (s) is given by

$$\mathbf{\rho}^*(\beta) = \operatorname{tr}_{b}(\mathbf{\rho}_{t}(\beta)) = \frac{\operatorname{tr}_{b}(\exp(-\beta \mathbf{H}))}{\operatorname{tr}(\exp(-\beta \mathbf{H}))},\tag{3}$$

where $tr_h(\cdot)$ is the partial trace over subsystem (b).¹³

¹³Campisi, Zueco, and Talkner 2010; Ingold, Hänggi, and Talkner 2009; Talkner and Hänggi 2020.

von Neumann entropy of Heisenberg spin chains

The von Neumann entropy $-\operatorname{tr}(\boldsymbol{\rho}^*(\boldsymbol{\beta})\ln(\boldsymbol{\rho}^*(\boldsymbol{\beta})))$ is a measure of the entanglement between subsystems (s) and (b).

Understanding the von Neumann entropy for a range of a system with Hamiltonian $\mathbf{H}(\theta)$ at a range of parameter values θ and inverse temperatures β is of interest.

We will consider a special case

$$\mathbf{H} = \sum_{i,j} \left[J_{i,j}^{\mathbf{x}} \mathbf{\sigma}_{i}^{\mathbf{x}} \mathbf{\sigma}_{j}^{\mathbf{x}} + J_{i,j}^{\mathbf{y}} \mathbf{\sigma}_{i}^{\mathbf{y}} \mathbf{\sigma}_{j}^{\mathbf{y}} + J_{i,j}^{\mathbf{z}} \mathbf{\sigma}_{i}^{\mathbf{z}} \mathbf{\sigma}_{j}^{\mathbf{z}} \right] + \frac{h}{2} \sum_{i=1}^{N} \mathbf{\sigma}_{i}^{\mathbf{z}}.$$

where h is the magnetic field strength.

Subsystem (s) corresponds to i=1,2 and subsystem (b) corresponds to the rest of the spins.

Key question:

How to compute reduced density matrices numerically?

A starting point: stochastic trace estimation

If **b** is a standard Gaussian random vector:

$$\mathbb{E}[\mathbf{b}^{\mathsf{T}} f(\mathbf{A}) \mathbf{b}] = \operatorname{tr}(f(\mathbf{A})), \qquad \mathbb{V}[\mathbf{b}^{\mathsf{T}} f(\mathbf{A}) \mathbf{b}] = 2 \| f(\mathbf{A}) \|_{\mathsf{F}}^{2}.$$

It's standard to use a KSM to approximate products $\mathbf{b} \mapsto \mathbf{b}^{\mathsf{T}} f(\mathbf{A}) \mathbf{b}$.

Lots of work balancing the cost of the KSM with the variance of the estimator¹⁴.

¹⁴Han, Malioutov, Avron, and Shin 2017; Ubaru, Chen, and Saad 2017; Chen, Trogdon, and Ubaru 2021; Chen, Trogdon, and Ubaru 2022; Braverman, Krishnan, and Musco 2022.

Partial traces

Suppose **A** is a $d_s d_b \times d_s d_b$ matrix partitioned as:

$$\mathbf{A} = \begin{bmatrix} \mathbf{A}_{1,1} & \mathbf{A}_{1,2} & \cdots & \mathbf{A}_{1,d_s} \\ \mathbf{A}_{2,1} & \mathbf{A}_{2,2} & \cdots & \mathbf{A}_{2,d_s} \\ \vdots & \vdots & \ddots & \vdots \\ \mathbf{A}_{d_s,1} & \mathbf{A}_{d_s,2} & \cdots & \mathbf{A}_{d_s,d_s} \end{bmatrix},$$

Partial traces

Then the partial trace (wrt. this partitioning) is defined as:

$$\mathrm{tr}_{\mathrm{b}}(\mathbf{A}) = \begin{bmatrix} \mathrm{tr}(\mathbf{A}_{1,1}) & \mathrm{tr}(\mathbf{A}_{1,2}) & \cdots & \mathrm{tr}(\mathbf{A}_{1,d_{\mathrm{s}}}) \\ \mathrm{tr}(\mathbf{A}_{2,1}) & \mathrm{tr}(\mathbf{A}_{2,2}) & \cdots & \mathrm{tr}(\mathbf{A}_{2,d_{\mathrm{s}}}) \\ \vdots & \vdots & \ddots & \vdots \\ \mathrm{tr}(\mathbf{A}_{d_{\mathrm{s},1}}) & \mathrm{tr}(\mathbf{A}_{d_{\mathrm{s},2}}) & \cdots & \mathrm{tr}(\mathbf{A}_{d_{\mathrm{s},d_{\mathrm{s}}}}) \end{bmatrix}.$$

An algorithm for partial traces¹⁵

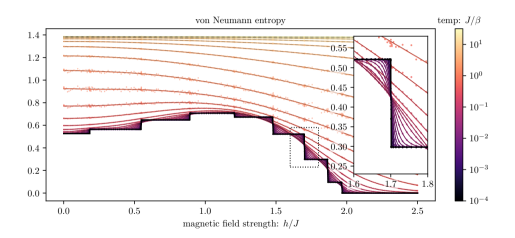
We can use a randomized estimator:

$$(\mathbf{I}_{d_{\mathrm{s}}} \otimes \mathbf{b})^{\mathsf{T}} \mathbf{A} (\mathbf{I}_{d_{\mathrm{s}}} \otimes \mathbf{b}) = \begin{bmatrix} \mathbf{b}^{\mathsf{T}} \mathbf{A}_{1,1} \mathbf{b} & \mathbf{b}^{\mathsf{T}} \mathbf{A}_{1,2} \mathbf{b} & \cdots & \mathbf{b}^{\mathsf{T}} \mathbf{A}_{1,d_{\mathrm{s}}} \mathbf{b} \\ \mathbf{b}^{\mathsf{T}} \mathbf{A}_{2,1} \mathbf{b} & \mathbf{b}^{\mathsf{T}} \mathbf{A}_{2,2} \mathbf{b} & \cdots & \mathbf{b}^{\mathsf{T}} \mathbf{A}_{2,d_{\mathrm{s}}} \mathbf{b} \\ \vdots & \vdots & \ddots & \vdots \\ \mathbf{b}^{\mathsf{T}} \mathbf{A}_{d_{\mathrm{s}},1} \mathbf{b} & \mathbf{b}^{\mathsf{T}} \mathbf{A}_{d_{\mathrm{s}},2} \mathbf{b} & \cdots & \mathbf{b}^{\mathsf{T}} \mathbf{A}_{d_{\mathrm{s}},d_{\mathrm{s}}} \mathbf{b} \end{bmatrix}.$$

Then use a KSM to approximate products with A = f(H).

¹⁵Chen and Cheng 2022.

von Neumann entropy



Spectrum approximation

Recall a symmetric matrix **A** has a real spectrum $\{\lambda_i\}$. We can encode the spectrum in the spectral density function

$$\varphi(x) = \sum_{i=1}^{n} \frac{1}{n} \delta(x - \lambda_i).$$

Here $\delta(x)$ is a Dirac delta point mass centered at zero.

Problem. Given only matrix-vector product access to **A**, approximate $\varphi(x)$.

Example application: high performance computing

State of the art parallel eigensolvers such as FEAST and EVSL work by splitting the spectrum of ${\bf A}$ into pieces, which can each be solved on different machines in parallel. 16



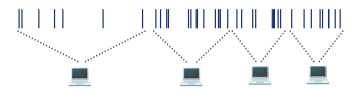
The spectral density tells us how many eigenvalues in a region

number of eigenvalues in
$$[a,b]=n\int_a^b \varphi(x)\mathrm{d}x$$

¹⁶Polizzi 2009; Li, Xi, Erlandson, and Saad 2019.

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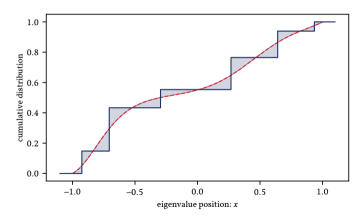
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¹⁶Polizzi 2009; Li, Xi, Erlandson, and Saad 2019.

Wasserstein distance

To measure the error in approximating $\varphi(x)$, we use the Wasserstein distance.



Stochatic Lanczos Quadrature

A widely used algorithm for approximating spectral densities Stocahstic Lanczos Quadrature (SLQ).

- used in ML to study the training of neural networks
- used in quantum physics for understanding many different systems

Until recently, no theoretical gurantees!

Theorem (Chen, Trogdon, and Ubaru 2021). For large matrices, ^a SLQ produces an ε accurate approximation to $\varphi(x)$ using $O(1/\varepsilon)$ products with **A**.

[&]quot;specifically, for $n \gg 1/\varepsilon^2$

Proof sketch

We use a general technique which actually applies to some other algorithms such as the Kernel Polynomial Method¹⁷.

- If two distributions have similar Chebyshev moments through degree O(1/ε), they are ε close in Wasserstein distance.
 - This follows from Jackson's Theorem from approximation theory and the dual representation of the Wasserstein distance in terms of 1-Lipshitz functions
- The Chebyshev moments of $\varphi(x)$ are $\int T_k(x)\varphi(x)\mathrm{d}x=n^{-1}\operatorname{tr}(T_k(\mathbf{A}))$
- We can estimate $tr(T_k(\mathbf{A}))$ using stochastic trace estimation
 - If \mathbf{x} is an isotropic random vector, $\mathbb{E}[\mathbf{x}^\mathsf{T} \mathbf{M} \mathbf{x}] = \mathrm{tr}(\mathbf{M})$ and $\mathbb{V}[\mathbf{x}^\mathsf{T} \mathbf{M} \mathbf{x}] \approx 2 \|\mathbf{M}\|_{\mathsf{F}}$.
- We can compute $\mathbf{x}^T T_k(\mathbf{A}) \mathbf{x}$ using O(k) products with \mathbf{A} .

¹⁷Chen, Trogdon, and Ubaru 2022.

Chebyshev moments

