STAT 9910-303: Large-Scale Optimization for Data Science

Nonconvex Optimization for High-Dimensional Estimation (Part 2)



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A case study: solving quadratic systems of equations

Solving quadratic systems of equations



Recover $\boldsymbol{x}^{\star} \in \mathbb{R}^n$ from m random quadratic measurements

$$y_k = (\boldsymbol{a}_k^\top \boldsymbol{x}^\star)^2, \qquad k = 1, \dots, m$$

assume w.l.o.g. $\| oldsymbol{x}^\star \|_2 = 1$

Motivation: phase retrieval

Detectors record intensities of diffracted rays

• electric field $x(t_1, t_2) \longrightarrow$ Fourier transform $\widehat{x}(f_1, f_2)$

figure credit: Stanford SLAC



intensity of electrical field: $|\hat{x}(f_1, f_2)|^2 = \left| \int x(t_1, t_2) e^{-i2\pi(f_1t_1 + f_2t_2)} dt_1 dt_2 \right|^2$

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Phase retrieval: recover signal $x(t_1, t_2)$ from intensity $|\hat{x}(f_1, f_2)|^2$

Motivation: learning neural nets with quadratic activation

— Soltanolkotabi, Javanmard, Lee '17, Li, Ma, Zhang '17



input layer

input features: *a*;

res:
$$a$$
; weights: $X^{\star} = [x_1^{\star}, \cdots, x_r^{\star}]$
output: $y = \sum_{i=1}^r \sigma(a^{\top} x_i^{\star})$

Motivation: learning neural nets with quadratic activation

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We consider simplest model when r = 1 (higher r is similar)

An equivalent view: low-rank factorization



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Introduce $\boldsymbol{X} = \boldsymbol{x} \boldsymbol{x}^{\top}$ to linearize constraints $y_k = (\boldsymbol{a}_k^{\top} \boldsymbol{x})^2 = \boldsymbol{a}_k^{\top} (\boldsymbol{x} \boldsymbol{x}^{\top}) \boldsymbol{a} \implies y_k = \boldsymbol{a}_k^{\top} \boldsymbol{X} \boldsymbol{a}_k$





An equivalent view: low-rank factorization

Introduce $X = xx^{\top}$ to linearize constraints $y_k = (a_k^{\top}x)^2 = a_k^{\top}(xx^{\top})a \implies y_k = a_k^{\top}Xa_k$



Solving quadratic systems is essentially low-rank matrix completion

given:
$$y_k = (\boldsymbol{a}_k^{\top} \boldsymbol{x}^{\star})^2, \quad 1 \le k \le m$$

$$\downarrow$$
minimize $_{\boldsymbol{x} \in \mathbb{R}^n} \quad f(\boldsymbol{x}) = \frac{1}{4m} \sum_{k=1}^m \left[(\boldsymbol{a}_k^{\top} \boldsymbol{x})^2 - y_k \right]^2$

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- pros: often exact as long as sample size is sufficiently large
- cons: $f(\cdot)$ is highly nonconvex \longrightarrow computationally challenging!

Wirtinger flow (Candès, Li, Soltanolkotabi '14)

minimize_{*x*}
$$f(\boldsymbol{x}) = \frac{1}{4m} \sum_{k=1}^{m} \left[\left(\boldsymbol{a}_{k}^{\top} \boldsymbol{x} \right)^{2} - y_{k} \right]^{2}$$

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- spectral initialization: $x^0 \leftarrow$ leading eigenvector of certain data matrix
- gradient descent:

$$\boldsymbol{x}^{t+1} = \boldsymbol{x}^t - \eta \nabla f(\boldsymbol{x}^t), \qquad t = 0, 1, \cdots$$

 $x^0 \leftarrow$ leading eigenvector of

$$oldsymbol{Y} := rac{1}{m} \sum_{k=1}^m y_k oldsymbol{a}_k oldsymbol{a}_k^ op$$

Rationale: under random Gaussian design $a_i \stackrel{\text{ind.}}{\sim} \mathcal{N}(\mathbf{0}, I)$,

$$\mathbb{E}[\boldsymbol{Y}] := \mathbb{E}\left[\frac{1}{m}\sum_{k=1}^{m} \boldsymbol{y}_k \boldsymbol{a}_k \boldsymbol{a}_k^{\top}\right] = \underbrace{\|\boldsymbol{x}^{\star}\|_2^2 \boldsymbol{I} + 2\boldsymbol{x}^{\star} \boldsymbol{x}^{\star^{\top}}}_{\text{leading eigenvector: } \pm \boldsymbol{x}^{\star}}$$



1. initialize within

local basin sufficiently close to x^\star

(restricted) strongly convex; no saddles / spurious local mins



2. iterative refinement

A highly incomplete list of two-stage methods

phase retrieval:

- Netrapalli, Jain, Sanghavi '13
- Candès, Li, Soltanolkotabi '14
- Chen, Candès '15
- Cai, Li, Ma'15
- Wang, Giannakis, Eldar '16
- Zhang, Zhou, Liang, Chi'16
- Kolte, Ozgur '16
- Zhang, Chi, Liang '16
- Soltanolkotabi '17
- Vaswani, Nayer, Eldar'16
- Chi, Lu'16
- Wang, Zhang, Giannakis, Akcakaya, Chen'16
- Tan, Vershynin '17
- Ma, Wang, Chi, Chen '17
- Duchi, Ruan'17
- Jeong, Gunturk '17
- Yang, Yang, Fang, Zhao, Wang, Neykov '17
- Qu, Zhang, Wright '17
- Goldstein, Studer '16
- Bahmani, Romberg '16
- Hand, Voroninski '16
- Wang, Giannakis, Saad, Chen'17
- Barmherzig, Sun '17
- ...

other problems:

- Keshavan, Montanari, Oh'09
- Sun, Luo'14
- Chen, Wainwright '15
- Tu, Boczar, Simchowitz, Soltanolkotabi, Recht '15
- Zheng, Lafferty '15
- Balakrishnan, Wainwright, Yu'14
- Chen, Suh '15
- Chen, Candès '16
- Li, Ling, Strohmer, Wei '16
- Yi, Park, Chen, Caramanis'16
- Jin, Kakade, Netrapalli '16
- Huang, Kakade, Kong, Valiant '16
- Ling, Strohmer'17
- Li, Ma, Chen, Chi'18
- Aghasi, Ahmed, Hand '17
- Lee, Tian, Romberg '17
- Li, Chi, Zhang, Liang '17
- Cai, Wang, Wei'17
- Abbe, Bandeira, Hall '14
- Chen, Kamath, Suh, Tse '16
- Zhang, Zhou'17
- Boumal '16
- Zhong, Boumal '17
- .

Theorem 1 (Candès, Li, Soltanolkotabi'14)

Under i.i.d. Gaussian design, WF with spectral initialization achieves

$$\mathsf{dist}(oldsymbol{x}^t,oldsymbol{x}^\star)\lesssim \left(1-rac{\eta}{4}
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with high prob., provided that step size $\eta \lesssim 1/n$ and sample size: $m \gtrsim n \log n$.

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- Iteration complexity: $O(n \log \frac{1}{\epsilon})$
- Sample complexity: $O(n \log n)$

 $\operatorname{dist}(\boldsymbol{x}^t, \boldsymbol{x}^\star) := \min\{\|\boldsymbol{x}^t \pm \boldsymbol{x}^\star\|_2\}$

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with high prob., provided that step size and sample size: .

- Iteration complexity: $O(n \log \frac{1}{\epsilon})$
- Sample complexity: $O(n \log n)$
- Derived based on (worst-case) local geometry

Theorem 2 (Ma, Wang, Chi, Chen '17)

Under i.i.d. Gaussian design, WF with spectral initialization achieves

$$\mathsf{dist}(\boldsymbol{x}^t, \boldsymbol{x}^\star) \lesssim \left(1 - \frac{\eta}{2}\right)^t \|\boldsymbol{x}^\star\|_2$$

with high prob., provided that step size $\eta \simeq 1/\log n$ and sample size $m \gtrsim n \log n$.

- Iteration complexity: $O(n \log \frac{1}{\epsilon}) \searrow O(\log n \log \frac{1}{\epsilon})$
- Sample complexity: $O(n \log n)$
- Derived based on finer analysis of GD trajectory

What does optimization theory say about WF?

Gaussian designs: $a_k \stackrel{\text{i.i.d.}}{\sim} \mathcal{N}(\mathbf{0}, I_n), \quad 1 \leq k \leq m$

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Finite-sample level $(m \asymp n \log n)$

 $\nabla^2 f(\boldsymbol{x}) \succ \boldsymbol{0}$

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$$\nabla^2 f(\boldsymbol{x}) \succ \boldsymbol{0}$$
 but ill-conditioned (even locally)

condition number $\asymp n$

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 (even locally) (even locally)

condition number $\asymp n$

Consequence (Candès et al '14): WF attains ε -accuracy within $O(n \log \frac{1}{\varepsilon})$ iterations if $m \asymp n \log n$

Generic optimization theory gives pessimistic bounds

WF converges in ${\cal O}(n)$ iterations

 $\mathsf{WF}\xspace$ converges in O(n) iterations

Step size taken to be $\eta = O(1/n)$

WF converges in O(n) iterations $\label{eq:stepsize} \square$ Step size taken to be $\eta = O(1/n)$

This choice is suggested by worst-case optimization theory

WF converges in O(n) iterations $\begin{array}{c} & & & \\ & & \\ & & \\ \end{array}$ Step size taken to be $\eta = O(1/n)$ $\begin{array}{c} & \\ & \\ & \\ \end{array}$ This choice is suggested by worst-case optimization theory $\begin{array}{c} & \\ & \\ & \\ \end{array}$ Does it capture what really happens?

Numerical efficiency with $\eta_t = 0.1$



Vanilla GD (WF) converges fast for a constant step size!

A second look at gradient descent theory

Which local region enjoys both strong convexity and smoothness?

$$\nabla^2 f(\boldsymbol{x}) = \frac{1}{m} \sum_{k=1}^m \left[3 (\boldsymbol{a}_k^\top \boldsymbol{x})^2 - (\boldsymbol{a}_k^\top \boldsymbol{x}^\star)^2 \right] \boldsymbol{a}_k \boldsymbol{a}_k^\top$$
Which local region enjoys both strong convexity and smoothness?

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• Not sufficiently smooth if x and a_k are too close (coherent)

Which local region enjoys both strong convexity and smoothness?



• x is incoherent w.r.t. sampling vectors $\{a_k\}$ (incoherence region)

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• x is incoherent w.r.t. sampling vectors $\{a_k\}$ (incoherence region)

Prior works suggest enforcing regularization (e.g. truncation, projection, regularized loss) to promote incoherence











GD implicitly forces iterates to remain incoherent with $\{a_k\}$ $\max_k |a_k^\top (x^t - x^*)| \lesssim \sqrt{\log n} ||x^*||_2, \quad \forall t$

 cannot be derived from generic optimization theory; relies on finer statistical analysis for entire trajectory of GD

Theorem 3 (Ma, Wang, Chi, Chen '17)

Under i.i.d. Gaussian design, WF with spectral initialization achieves

• $\max_k |\boldsymbol{a}_k^\top \boldsymbol{x}^t| \lesssim \sqrt{\log n} \, \|\boldsymbol{x}^\star\|_2$ (incoherence)

Theorem 3 (Ma, Wang, Chi, Chen '17)

Under i.i.d. Gaussian design, WF with spectral initialization achieves

- $\max_k |\boldsymbol{a}_k^\top \boldsymbol{x}^t| \lesssim \sqrt{\log n} \, \|\boldsymbol{x}^\star\|_2$ (incoherence)
- dist $(\boldsymbol{x}^t, \boldsymbol{x}^\star) \lesssim (1 \frac{\eta}{2})^t \| \boldsymbol{x}^\star \|_2$ (linear convergence)

provided that step size $\eta \simeq 1/\log n$ and sample size $m \gtrsim n \log n$.

• Attains ε accuracy within $O(\log n \log \frac{1}{\varepsilon})$ iterations

For each $1 \leq l \leq m$, introduce leave-one-out iterates $x^{t,(l)}$ by dropping lth measurement





• Leave-one-out iterate $x^{t,(l)}$ is independent of a_l



- Leave-one-out iterate $oldsymbol{x}^{t,(l)}$ is independent of $oldsymbol{a}_l$
- Leave-one-out iterate $oldsymbol{x}^{t,(l)}pprox$ true iterate $oldsymbol{x}^t$



- Leave-one-out iterate $x^{t,(l)}$ is independent of a_l
- Leave-one-out iterate $oldsymbol{x}^{t,(l)}pprox$ true iterate $oldsymbol{x}^t$

$$\implies x^t$$
 is nearly independent of a_l

• Several prior works use sample-splitting: require fresh samples at each iteration; not practical but helps analysis



• Several prior works use sample-splitting: require fresh samples at each iteration; not practical but helps analysis



• This tutorial: reuses all samples in all iterations



Can we further improve sample complexity (to O(n))?

Truncated spectral initialization



problem: unless $m \gg n$, dangerous to use empirical average because large observations $y_k = (\boldsymbol{a}_k^\top \boldsymbol{x}^\star)^2$ bear too much influence

Truncated spectral initialization



- **problem:** unless $m \gg n$, dangerous to use empirical average because large observations $y_k = (\boldsymbol{a}_k^\top \boldsymbol{x}^\star)^2$ bear too much influence
- **solution:** discard high leverage samples and compute leading eigenvector of truncated sum

$$\frac{1}{m}\sum_{k=1}^{m} y_k \boldsymbol{a}_k \boldsymbol{a}_k^\top \cdot \mathbf{1}_{\{|\boldsymbol{y}_k| \le \alpha^2 \operatorname{Avg}(|\boldsymbol{y}_j|)\}}$$





Original image



Spectral initialization



Spectral initialization



Truncated spectral initialization

Precise asymptotic characterization (Lu, Li'17)

(1,1)

- $m/n \asymp 1$
- i.i.d. Gaussian design

Fig. credit: Lu, Li '17

Theorem 4 (Lu, Li'17, Mondelli, Montanari'17)

There exist analytical formulas $\rho(\cdot)$ and constants α_{\min} and α_{\max} s.t.

$$\underbrace{\frac{(\boldsymbol{x}^{\star\top}\boldsymbol{x}^{0})^{2}}{\|\boldsymbol{x}^{\star}\|_{2}^{2}\|\boldsymbol{x}^{0}\|_{2}^{2}}}_{\text{min}} \longrightarrow \begin{cases} 0, & \text{if } m/n < \alpha_{\min} \\ \rho(m/n), & \text{if } m/n > \alpha_{\max} \end{cases}$$

Theoretical prediction vs. simulations

image size: 64×64



Fig. credit: Lu, Li'17

WF (GD):
$$\boldsymbol{x}^{t+1} = \boldsymbol{x}^t - \frac{\eta}{m} \sum_k \nabla f_k(\boldsymbol{x}^t)$$





Problem: descent direction might have large variability

Solution: variance reduction via trimming

More adaptive rule: $\boldsymbol{x}^{t+1} = \boldsymbol{x}^t - \frac{\eta}{m} \sum_{k \in \mathcal{T}_t} \nabla f_k(\boldsymbol{x}^t)$



Solution: variance reduction via trimming

More adaptive rule: $\boldsymbol{x}^{t+1} = \boldsymbol{x}^t - \frac{\eta}{m} \sum_{k \in \mathcal{T}_t} \nabla f_k(\boldsymbol{x}^t)$



• T_t trims away excessively large grad components

$$\mathcal{T}_{t} := \left\{ k: \quad \left\| \nabla f_{k}(\boldsymbol{x}^{t}) \right\|_{2} \; \lesssim \; \text{typical-size} \Big\{ \left\| \nabla f_{l}(\boldsymbol{x}^{t}) \right\|_{2} \Big\}_{1 \leq l \leq m} \right\}$$

Slight bias + much reduced variance

(1) Regularized spectral initialization: $x^0 \leftarrow$ principal component of

$$\frac{1}{m} \sum_{k \in \mathcal{T}_0} y_k \, \boldsymbol{a}_k \boldsymbol{a}_k^\top$$

(2) Follow adaptive gradient descent

$$\boldsymbol{x}^{t} = \boldsymbol{x}^{t} - \frac{\eta_{t}}{m} \sum_{k \in \mathcal{T}_{t}} \nabla f_{k}(\boldsymbol{x}^{t})$$

Adaptive and iteration-varying rules: discard high-leverage data $\{y_k: k \notin \mathcal{T}_t\}$

Theoretical guarantees (noiseless data)



Theorem 5 (Chen, Candès '15)

Suppose $a_k \stackrel{i.i.d.}{\sim} \mathcal{N}(\mathbf{0}, I_n)$ and sample size $m \gtrsim n$. With high prob., $\operatorname{dist}(\boldsymbol{x}^t, \boldsymbol{x}^\star) := \min \|\boldsymbol{x}^t \pm \boldsymbol{x}^\star\|_2 \leq \nu (1-\rho)^t \|\boldsymbol{x}^\star\|_2$

where $0 < \nu, \rho < 1$ are universal constants

Empirical success rate (noiseless data)



Empirical success rate vs. sample size

Stability vis a vis noise?

- Noisy data: $y_k = (\boldsymbol{a}_k^{ op} \boldsymbol{x}^\star)^2 + \eta_k$
- Signal-to-noise ratio:

$$\mathsf{SNR} := rac{\sum_k (oldsymbol{a}_k^ opoldsymbol{x}^\star)^4}{\sum_k \eta_k^2} pprox rac{3m \|oldsymbol{x}^\star\|_2^4}{\|oldsymbol{\eta}\|_2^2}$$

• i.i.d. Gaussian design $\boldsymbol{a}_k \overset{\text{i.i.d.}}{\sim} \mathcal{N}(\boldsymbol{0}, \boldsymbol{I}_n)$
- Noisy data: $y_k = (\boldsymbol{a}_k^{ op} \boldsymbol{x}^\star)^2 + \eta_k$
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• i.i.d. Gaussian design $\boldsymbol{a}_k \overset{\text{i.i.d.}}{\sim} \mathcal{N}(\boldsymbol{0}, \boldsymbol{I}_n)$

Theorem 6 (Chen, Candès '15)

Relative error of TWF converges to $O(\frac{1}{\sqrt{\text{SNR}}})$

Relative MSE vs. SNR (Poisson data)



Empirical evidence: relative MSE scales inversely with SNR

This accuracy is nearly un-improvable (empirically)

Comparison with ideal MLE (with phase info. revealed)

ideal knowledge: $y_k \sim \text{Poisson}(|\boldsymbol{a}_k^\top \boldsymbol{x}^\star|^2)$ and $\varepsilon_k = \text{sign}(\boldsymbol{a}_k^\top \boldsymbol{x}^\star)$



Little loss due to missing phases!

This accuracy is nearly un-improvable (theoretically)

- Poisson data: $y_k \stackrel{\text{ind.}}{\sim} \text{Poisson}(|\boldsymbol{a}_k^\top \boldsymbol{x}^\star|^2)$
- Signal-to-noise ratio:

$$\mathsf{SNR} \; \approx \; \frac{\sum_k |\boldsymbol{a}_k^\top \boldsymbol{x}^\star|^4}{\sum_k \mathsf{Var}(y_k)} \; \approx \; 3 \|\boldsymbol{x}^\star\|_2^2$$

This accuracy is nearly un-improvable (theoretically)

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- Signal-to-noise ratio:

$$\mathsf{SNR} ~pprox ~ rac{\sum_k |oldsymbol{a}_k^ op oldsymbol{x}^\star|^4}{\sum_k \mathsf{Var}(y_k)} ~pprox ~ 3 \|oldsymbol{x}^\star\|_2^2$$

Theorem 7 (Chen, Candès '15)

Under i.i.d. Gaussian design, for any estimator \widehat{x} ,

$$\inf_{\widehat{\boldsymbol{x}}} \sup_{\boldsymbol{x}^{\star}: \|\boldsymbol{x}^{\star}\|_{2} \geq \log^{1.5} m} \frac{\mathbb{E}\left[\operatorname{dist}\left(\widehat{\boldsymbol{x}}, \boldsymbol{x}^{\star}\right) \mid \{\boldsymbol{a}_{k}\}\right]}{\|\boldsymbol{x}^{\star}\|_{2}} \gtrsim \frac{1}{\sqrt{\mathsf{SNR}}},$$

provided that sample size $m \asymp n$

Other examples: low-rank matrix estimation

Problem: complete a rank-r matrix \boldsymbol{M} from partial entries: $M_{i,j}$, $(i,j)\in \Omega$

• random sampling: (i,j) is included in Ω independently with prob. p

find low-rank \widehat{M} s.t. $\mathcal{P}_{\Omega}(\widehat{M}) = \mathcal{P}_{\Omega}(M)$

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find low-rank
$$\widehat{oldsymbol{M}}$$
 s.t. $\mathcal{P}_\Omega(\widehat{oldsymbol{M}})=\mathcal{P}_\Omega(oldsymbol{M})$

Strong convexity and smoothness do not hold in general

 $\rightarrow~$ need to regularize loss function by promoting $\ensuremath{\text{incoherent}}$ solutions

Definition 8 (Incoherence for matrix completion)

A rank-r matrix M with eigendecomposition $M = U \Sigma U^\top$ is said to be $\mu\text{-incoherent}$ if

$$ig\|oldsymbol{U}ig\|_{2,\infty} \leq \sqrt{rac{\mu}{n}}ig\|oldsymbol{U}ig\|_{ ext{F}} = \sqrt{rac{\mu r}{n}}$$



Let $M = X^{\star}X^{\star op}$. Observe

$$Y_{i,j} = M_{i,j} + E_{i,j}, \quad (i,j) \in \Omega$$

where $(i, j) \in \Omega$ independently with prob. p, and $E_{i,j} \sim \mathcal{N}(0, \sigma^2)^1$

$$\mathsf{minimize} \left\| \mathcal{P}_\Omega \big(\widehat{\boldsymbol{M}} - \boldsymbol{Y} \big) \right\|_\mathrm{F}^2 \quad \mathsf{s.t.} \quad \mathsf{rank}(\widehat{\boldsymbol{M}}) \leq r$$

¹can be relaxed to sub-Gaussian noise and the asymmetric case

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where $(i, j) \in \Omega$ independently with prob. p, and $E_{i,j} \sim \mathcal{N}(0, \sigma^2)^1$

$$\begin{split} \mininimize \left\| \mathcal{P}_{\Omega}(\widehat{\boldsymbol{M}} - \boldsymbol{Y}) \right\|_{\mathrm{F}}^{2} \quad \text{s.t.} \quad \operatorname{rank}(\widehat{\boldsymbol{M}}) \leq r \\ \mininimize_{\boldsymbol{X} \in \mathbb{R}^{n \times r}} \quad \underbrace{f(\boldsymbol{X}) = \sum_{(j,k) \in \Omega} \left(\boldsymbol{e}_{j}^{\top} \boldsymbol{X} \boldsymbol{X}^{\top} \boldsymbol{e}_{k} - Y_{j,k} \right)^{2}}_{\text{unregularized least-squares loss}} \end{split}$$

¹can be relaxed to sub-Gaussian noise and the asymmetric case

1. spectral initialization: let $U^0 \Sigma^0 U^{0\top}$ be rank-r eigendecomposition of

$$\frac{1}{p}\mathcal{P}_{\Omega}(\boldsymbol{Y}).$$

and set $oldsymbol{X}^0 = oldsymbol{U}^0 \left(oldsymbol{\Sigma}^0
ight)^{1/2}$

2. gradient descent updates:

$$\boldsymbol{X}^{t+1} = \boldsymbol{X}^t - \eta_t \nabla f(\boldsymbol{X}^t), \qquad t = 0, 1, \cdots$$

Define the optimal rotation from the tth iterate $oldsymbol{X}^{t}$ to $oldsymbol{X}^{\star}$ as

$$oldsymbol{Q}^t := \mathsf{argmin}_{oldsymbol{R} \in \mathcal{O}^{r imes r}} ig\| oldsymbol{X}^t oldsymbol{R} - oldsymbol{X}^\star ig\|_{\mathrm{F}}$$

where $\mathcal{O}^{r \times r}$ is the set of $r \times r$ orthonormal matrices

• orthogonal Procrustes problem

Theorem 9 (Noiseless MC, Ma, Wang, Chi, Chen '17)

Suppose $M = X^* X^{*\top}$ is rank-*r*, incoherent and well-conditioned. Vanilla GD (with spectral initialization) achieves

•
$$\| oldsymbol{X}^t oldsymbol{Q}^t - oldsymbol{X}^\star \|_{ ext{F}} \lesssim rac{
ho^t \mu r rac{1}{\sqrt{np}} \| oldsymbol{X}^\star \|_{ ext{F}}$$
 ,

•
$$\| \mathbf{X}^t \mathbf{Q}^t - \mathbf{X}^\star \| \lesssim \rho^t \mu r \frac{1}{\sqrt{np}} \| \mathbf{X}^\star \|$$
, (spectral)

•
$$\| oldsymbol{X}^t oldsymbol{Q}^t - oldsymbol{X}^\star \|_{2,\infty} \lesssim oldsymbol{
ho}^t \mu r \sqrt{rac{\log n}{np}} \| oldsymbol{X}^\star \|_{2,\infty}$$
, (incoherence)

where $0<\rho<1,$ if the step size $\eta\asymp 1/\sigma_{max}$ and the sample complexity $n^2p\gtrsim \mu^3nr^3\log^3n$

Theorem 9 (Noiseless MC, Ma, Wang, Chi, Chen '17)

Suppose $M = X^* X^{*\top}$ is rank-*r*, incoherent and well-conditioned. Vanilla GD (with spectral initialization) achieves

•
$$\| oldsymbol{X}^t oldsymbol{Q}^t - oldsymbol{X}^\star \|_{ ext{F}} \lesssim rac{
ho^t \mu r rac{1}{\sqrt{np}} \| oldsymbol{X}^\star \|_{ ext{F}}$$
 ,

•
$$\| \mathbf{X}^t \mathbf{Q}^t - \mathbf{X}^\star \| \lesssim \rho^t \mu r \frac{1}{\sqrt{np}} \| \mathbf{X}^\star \|$$
, (spectral)

•
$$\| \mathbf{X}^t \mathbf{Q}^t - \mathbf{X}^\star \|_{2,\infty} \lesssim \rho^t \mu r \sqrt{rac{\log n}{np}} \| \mathbf{X}^\star \|_{2,\infty}$$
, (incoherence)

where $0<\rho<1,$ if the step size $\eta\asymp 1/\sigma_{max}$ and the sample complexity $n^2p\gtrsim \mu^3nr^3\log^3n$

• vanilla GD converges linearly for matrix completion!

Numerical evidence for noiseless data



Relative error of $X^t X^{t\top}$ (measured by $\|\cdot\|_{\rm F}$, $\|\cdot\|$, $\|\cdot\|_{\infty}$) vs. iteration count for matrix completion, where n = 1000, r = 10, p = 0.1, and $\eta_t = 0.2$

$$\mathsf{minimize}_{\boldsymbol{X} \in \mathbb{R}^{n \times r}} \quad f(\boldsymbol{X}) = \sum_{(j,k) \in \Omega} \left(\boldsymbol{e}_j^\top \boldsymbol{X} \boldsymbol{X}^\top \boldsymbol{e}_k - Y_{j,k} \right)^2$$

Related theory promotes incoherence explicitly:

$$\mathsf{minimize}_{\boldsymbol{X} \in \mathbb{R}^{n \times r}} \quad f(\boldsymbol{X}) = \sum_{(j,k) \in \Omega} \left(\boldsymbol{e}_j^\top \boldsymbol{X} \boldsymbol{X}^\top \boldsymbol{e}_k - Y_{j,k} \right)^2$$

Related theory promotes incoherence explicitly:

- regularized loss (solve $\min_{\mathbf{X}} f(\mathbf{X}) + Q(\mathbf{X})$ instead)
 - $\circ~$ e.g. Keshavan, Montanari, Oh '10, Sun, Luo '14, Ge, Lee, Ma '16

$$\mathsf{minimize}_{\boldsymbol{X} \in \mathbb{R}^{n \times r}} \quad f(\boldsymbol{X}) = \sum_{(j,k) \in \Omega} \left(\boldsymbol{e}_j^\top \boldsymbol{X} \boldsymbol{X}^\top \boldsymbol{e}_k - Y_{j,k} \right)^2$$

Related theory promotes incoherence explicitly:

- regularized loss (solve min_X f(X) + Q(X) instead)
 e.g. Keshavan, Montanari, Oh '10, Sun, Luo '14, Ge, Lee, Ma '16
- projection onto set of incoherent matrices

 $\circ~$ e.g. Chen, Wainwright '15, Zheng, Lafferty '16

$$\boldsymbol{X}^{t+1} = \mathcal{P}_{\mathcal{C}}\left(\boldsymbol{X}^{t} - \eta_{t}\nabla f\left(\boldsymbol{X}^{t}\right)\right), \qquad t = 0, 1, \cdots$$

Are carefully-designed initialization or saddle-point escaping schemes necessary for fast convergence?

Initialization



• Spectral initialization gets us reasonably close to truth

Initialization



- Spectral initialization gets us reasonably close to truth
- Cannot initialize GD from anywhere, e.g. it might get stucked at local stationary points (e.g. saddle points)

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Can we initialize GD randomly, which is simpler and model-agnostic?

Numerical efficiency of randomly initialized GD

$$\eta_t = 0.1, \ \boldsymbol{a}_i \sim \mathcal{N}(\boldsymbol{0}, \boldsymbol{I}_n), \ m = 10n, \ \boldsymbol{x}^0 \sim \mathcal{N}(\boldsymbol{0}, n^{-1} \boldsymbol{I}_n)$$



Numerical efficiency of randomly initialized GD

$$\eta_t = 0.1$$
, $\boldsymbol{a}_i \sim \mathcal{N}(\boldsymbol{0}, \boldsymbol{I}_n)$, $m = 10n$, $\boldsymbol{x}^0 \sim \mathcal{N}(\boldsymbol{0}, n^{-1}\boldsymbol{I}_n)$



Randomly initialized GD enters local basin within a few iterations

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Randomly initialized GD enters local basin within a few iterations

A geometric analysis



- if $m \gtrsim n \log^3 n$, then (Sun et al. '16)
 - $\circ\;$ there is no spurious local mins
 - all saddle points are strict (i.e. associated Hessian matrices have at least one sufficiently negative eigenvalue)

A geometric analysis



• With such benign landscape, GD with random initialization converges to global min almost surely (Lee et al. '16)

No convergence rate guarantees for vanilla GD!

Exponential growth of signal strength in Stage 1



Exponential growth of signal strength in Stage 1



Numerically, $O(\log n)$ iterations are enough to enter local region

Linear / geometric convergence in Stage 2



Linear / geometric convergence in Stage 2



Numerically, GD converges linearly within local region

These numerical findings can be formalized when $a_i \overset{\text{i.i.d.}}{\sim} \mathcal{N}(\mathbf{0}, I_n)$:

Theorem 10 (Chen, Chi, Fan, Ma'18)

Under i.i.d. Gaussian design, GD with $m{x}^0 \sim \mathcal{N}(m{0}, n^{-1} m{I}_n)$ achieves

$$\operatorname{dist}(\boldsymbol{x}^{t}, \boldsymbol{x}^{\star}) \leq \gamma (1-\rho)^{t-T_{\gamma}} \| \boldsymbol{x}^{\star} \|_{2}, \qquad t \geq T_{\gamma}$$

for $T_{\gamma} \lesssim \log n$ and some constants $\gamma, \rho > 0$, provided that step size $\eta \asymp 1$ and sample size $m \gtrsim n \operatorname{poly} \log m$

Theoretical guarantees for randomly initialized GD

 $\operatorname{dist}(\boldsymbol{x}^{t}, \boldsymbol{x}^{\star}) \leq \gamma (1-\rho)^{t-T_{\gamma}} \|\boldsymbol{x}^{\star}\|_{2}, \quad t \geq T_{\gamma} \asymp \log n$



Theoretical guarantees for randomly initialized GD





• Stage 1: takes $O(\log n)$ iterations to reach $dist(\boldsymbol{x}^t, \boldsymbol{x}^\star) \leq \gamma$

Theoretical guarantees for randomly initialized GD





- Stage 1: takes $O(\log n)$ iterations to reach $dist(\boldsymbol{x}^t, \boldsymbol{x}^\star) \leq \gamma$
- Stage 2: linear convergence
Theoretical guarantees for randomly initialized GD





• near-optimal compututational cost:

 $-O(\log n + \log \frac{1}{\varepsilon})$ iterations to yield ε accuracy

Theoretical guarantees for randomly initialized GD





• near-optimal compututational cost:

 $-O(\log n + \log \frac{1}{\varepsilon})$ iterations to yield ε accuracy

• near-optimal sample size: $m\gtrsim n\mathsf{poly}\log m$

Saddle-escaping schemes?



Randomly initialized GD never hits saddle points in phase retrieval!

Other saddle-escaping schemes

	iteration complexity	num of iterations needed to escape saddles	local iteration complexity
Trust-region	$n^7 + \log \log \frac{1}{\varepsilon}$	n^7	$\log \log \frac{1}{\varepsilon}$
Perturbed GD (Jin et al. '17)	$n^3 + n \log \frac{1}{\varepsilon}$	n^3	$n\log \frac{1}{\varepsilon}$
Perturbed accelerated GD (Jin et al. '17)	$n^{2.5} + \sqrt{n} \log \frac{1}{\varepsilon}$	n ^{2.5}	$\sqrt{n}\log\frac{1}{\varepsilon}$
GD (Chen et al. '18)	$\log n + \log \frac{1}{\varepsilon}$	$\log n$	$\log \frac{1}{\varepsilon}$

Generic optimization theory yields highly suboptimal convergence guarantees

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