Numerical Matrix Analysis Notes #26 GMRES

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Image: A matrix

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26. GMRES

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GMRES

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- Moving Forward
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Setup and Notation Polynomial Approximation, and Convergence

Arnoldi Iteration $\rightsquigarrow A\vec{x} = \vec{b}$

Last time we looked at the Arnoldi Iteration as a procedure for finding eigenvalues. Next, we leverage it to solve $A\vec{x} = b$; introducing GMRES, the "Generalized Minimal RESiduals" strategy.



26. GMRES

Setup and Notation Moving Forward Polynomial Approximation, and Convergence

Structure, Notation, Idea

Problem Structure and Notation

We consider
$$A \in \mathbb{C}^{m \times m}$$
, with dim $(\operatorname{null}(A)) = 0$; $\vec{b} \in \mathbb{C}^m$;
 $K(A, \vec{b}, n) = \operatorname{span}\left(\vec{b}, A\vec{b}, \dots, A^{n-1}\vec{b}\right)$; and $\vec{x}_* = A^{-1}\vec{b}$ (exact solution).

GMRES Idea

At the n^{th} step, $\vec{x}_n \approx \vec{x}_*$ is the vector $\vec{x}_n \in K(A, \vec{b}, n)$ which minimizes $\|\vec{r_n}\|$, where $\vec{r_n} = (\vec{b} - A\vec{x_n})$; *i.e.* each $\vec{x_n}$ is the solution to a least squares problem over an *n*-dimensional (Krylov) subspace.

Many iterative optimization methods do something similar (at least in "spirit") — seeking approximately optimal approximations in carefully nested sequences of subspaces. (See [MATH 693A])



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GMRES GMRES: Matrix Polynomials Polynomial Approximation, and Convergence

GMRES: "Obvious" Strategy

With the Krylov matrix

$$K_n = \left[\vec{b} \ A\vec{b} \ \cdots \ A^{n-1}\vec{b} \right],$$

on hand, the "obvious" (ill-conditioned) way is to form

$$AK_n = \left[\begin{array}{cc} A\vec{b} & A^2\vec{b} & \cdots & A^n\vec{b} \end{array} \right],$$

which has the column space range(AK_n). We seek \vec{c}_n

$$ec{c}_n = \operatorname*{arg\,min}_{ec{c} \in \mathbb{C}^n} \| (AK_n) ec{c} - ec{b} \|, \quad ext{and} \ ec{x}_n = K_n ec{c}_n.$$

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Note: arg min "returns" the argument-that-minimizes the given function (objective).

The "Obvious" Strategy Fails (in Finite Precision)

A $Q_n R_n$ -factorization of AK_n would provide the necessary components of the pseudo-inverse necessary for identification of the solution to the least squares problem.

But, alas, this approach is numerically unstable, and wasteful (the R_n factor is not needed.)

Instead, we use the Arnoldi Iteration to construct Krylov Matrices Q_n , whose columns satisfy

span
$$(\vec{q}_1, \vec{q}_2, \ldots, \vec{q}_n) = K(A, \vec{b}, n),$$

thus we can represent $\vec{x_n} = Q_n \vec{y_n}$ rather than $\vec{x_n} = K_n \vec{c_n}$; the associated Least Squares Problem is

$$\vec{y}_n = \operatorname*{arg\,min}_{\vec{y} \in \mathbb{C}^n} \|AQ_n \vec{y} - \vec{b}\|.$$

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GMRES: Matrix Polynomials GMRES: Matrix Polynomials GMRES: Matrix Polynomials

"Shrinking" the Problem

As stated $\vec{y_n} = \arg \min_{\vec{y} \in \mathbb{C}^n} ||AQ_n \vec{y} - \vec{b}||$ is an $(m \times n)$ -dimensional Least Squares Problem, but using the structure of Krylov subspaces, its essential dimension is reduced to $((n + 1) \times n)$:

We use the "Arnoldi relation" $AQ_n = Q_{n+1}\tilde{H}_n$ to transform the problem into

$$ec{y}_n = rgmin_{ec{y} \in \mathbb{C}^n} \| Q_{n+1} \widetilde{H}_n ec{y} - ec{b} \|,$$

multiplication by Q_{n+1}^* preserves the norm, since both $(Q_{n+1}\tilde{H}_n\vec{y})$ and \vec{b} are — by construction — in the column space of Q_n ; we get

$$ec{y_n} = \mathop{\arg\min}_{ec{y} \in \mathbb{C}^n} \| ilde{H}_n ec{y} - Q_{n+1}^* ec{b} \|.$$

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Setup and Notation Moving Forward Polynomial Approximation, and Convergence

"Shrinking" the Problem

Finally, by construction of Q_n^{\ddagger} , we get $Q_{n+1}^*\vec{b} = \|\vec{b}\|\vec{e_1}$, so our problem is

$$ec{y_n} = rgmin_{ec{y} \in \mathbb{C}^n} \| ilde{H}_n ec{y} - eta ec{e_1} \|, \quad ext{where } eta = \| ec{b} \|;$$

and $\vec{x_n} = Q_n \vec{y_n}$.

 $\vec{e_1}$ is as usual the first standard basis vector in the appropriate space; it has a single "1" in the first component, and the remaining components are "0".



GMRES Algorithm

Algorithm (GMRES)

1: $\vec{b} \leftarrow \operatorname{random}(\mathbb{R}^{m \times 1}),$ 2: $\beta \leftarrow \|\vec{b}\|$ 3: $\vec{a}_1 \leftarrow \vec{b}/\beta$ 4: for $n \in \{1, 2, ...\}$ do 5: $\vec{v} \leftarrow A\vec{q}_n$ 6: **for** $j \in \{1, ..., n\}$ **do** 7: $h_{i,n} \leftarrow \vec{q}_i^* \vec{v}$ $\vec{v} \leftarrow \vec{v} - h_{i,n}\vec{q}_i$ 8: 9: end for $h_{n+1,n} \leftarrow \|\vec{v}\|$ 10: 11: $\vec{q}_{n+1} \leftarrow \vec{v}/h_{n+1,n}$ 12: $\vec{y}_n \leftarrow \arg\min_{\vec{v} \in \mathbb{C}^n} \|\tilde{H}_n \vec{v} - \beta \vec{e}_1\|$ 13: $\vec{x_n} \leftarrow Q_n \vec{v_n}$ 14: end for

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Comments

- In each step we solve an $((n + 1) \times n)$ Least Squares Problem with Hessenberg structure; the cost via QR-factorization is $\mathcal{O}(n^2)$ (exploiting the Hessenberg structure).
- It is possible to save work by identifying an updating strategy for the $Q_n R_n$ factorization of \tilde{H}_n from $Q_{n-1}R_{n-1} = \tilde{H}_{n-1}$. The cost is then one *Givens rotation*^{*} [T&B PROBLEMS 10.4 & 35.4] and $\mathcal{O}(n)$ work.
- * The Givens rotations are the building blocks for a slightly (50%) more expensive alternative to the Householder reflection method for computing the QR-factorization.

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Setup and Notation Moving Forward Polynomial Approximation, and Convergence

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Polynomial Class P_n

 $P_n = \{ \text{ Polynomials of degree} \leq n, \text{ with } p(0) = 1 \},$

i.e. the constant coefficient $c_0 = 1$.

Just as in the Arnoldi Iteration case, we can discuss the GMRES iteration in terms of polynomial approximations:

$$\vec{x}_n = q_n(A)\vec{b}$$

where $q_n(\cdot)$ is a polynomial of degree (n-1) with coefficients from the vector $\vec{c}_n = \arg \min_{\vec{c} \in \mathbb{C}^n} ||AK_n \vec{c} - \vec{b}||$.



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Polynomial Approximation

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With $p_n(z) = 1 - zq_n(z)$, we have

$$\vec{r}_n = \vec{b} - A\vec{x}_n = (I - Aq_n(A))\vec{b} = p_n(A)\vec{b},$$

for some $p_n \in P_n$.

GMRES solves the following problem

GMRES Approximation Problem

Find $p_n \in P_n$ such that

$$p_n = \underset{p \in P_n}{\arg\min} \| p(A) \vec{b} \|.$$

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Image: A matrix

Invariance Properties

Theorem

Let the GMRES iteration be applied to a matrix $A \in \mathbb{C}^{m \times m}$, then the following holds:

- [Scale-Invariance] If A is changed to σA for some $\sigma \in \mathbb{C}$, and \vec{b} is changed to $\sigma \vec{b}$, the residuals $\vec{r_n}$ change to $\sigma \vec{r_n}$.
- [INVARIANCE UNDER UNITARY TRANSFORMATIONS] If A is changed to UAU^* for some unitary matrix U, and \vec{b} is changed to $U\vec{b}$, the residuals $\vec{r_n}$ change to $U^*\vec{r_n}$.

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Convergence

Theorem (GMRES Convergence Property#1: Monotonic Convergence)

GMRES converges monotonically,

$$\|\vec{r}_{n+1}\| \leq \|\vec{r}_n\|.$$

This must be the case since we are minimizing over expanding subspaces, *i.e.* $K(A, \vec{b}, n) \subset K(A, \vec{b}, n+1)$.

Theorem (GMRES Converence Property#2: *m*-step Convergence)

In infinite precision, GMRES converges in at most m steps

 $\|\vec{r}_m\|=0.$

This must be the case since $K(A, \vec{b}, m) = \mathbb{C}^m$.



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The factor that gives us more useful convergence estimates is related to the polynomial p_n :

$$\frac{\|\vec{r}_n\|}{\|\vec{b}\|} \leq \inf_{p_n \in P_n} \|p_n(A)\|,$$

which brings us back to studying matrix polynomials related to Krylov subspaces.

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||*p_n*(*A*)|| Example: T&B-35.1 Example: T&B-35.2

How small can $||p_n(A)||$ be?

The standard way to get bounds on the behavior of $||p_n(A)||$ is to study polynomials on the spectrum $\lambda(A)$.

Definition

If p is a polynomial and $S \subset \mathbb{C}$, then

$$\|p\|_{\mathcal{S}} := \sup_{z \in \mathcal{S}} |p(z)|.$$

In the case where S is a finite set of points in the complex plane, the supremum (sup) is just the maximum (max).

When A is diagonalizable $A = V\Lambda V^{-1}$, then

 $\|p(A)\| \le \|V\| \|p(\Lambda)\| \|V^{-1}\| = \kappa(V) \|p\|_{\lambda(A)}.$

 $\kappa(V)$ is the conditioning of the Eigenbasis.



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||*p_n*(*A*)|| Example: T&B-35.1 Example: T&B-35.2

How small can $||p_n(A)||$ be?

Theorem

At step n of the GMRES iteration, the residual $\vec{r_n}$ satisfies

$$\frac{\|\vec{r}_n\|}{\|\vec{b}\|} \leq \inf_{p_n \in P_n} \|p_n(A)\| \leq \kappa(V) \inf_{p_n \in P_n} \|p_n\|_{\lambda(A)},$$

where $\lambda(A)$ is the set of eigenvalues of A, V is a non-singular matrix of eigenvectors (assuming A is diagonalizable), and $\|p_n\|_{\lambda(A)} = \sup_{z \in \lambda(A)} |p_n(z)|.$

As long as $\kappa(V)$ is not too large — *i.e.* the closer A is to being normal (unitarily diagonalizable) — and if polynomials p_n which decrease quickly on $\lambda(A)$ exist, then GMRES converges quickly.



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||*p*_n(*A*)|| **Example: T&B-35.1** Example: T&B-35.2



T&B-35.1

||*P*_n(A)|| Example: T&B-35.1 Example: T&B-35.2

- The eigenvalue spectrum of A is roughly contained in the disk of radius $\frac{1}{2}$, centered at z = 2.
- $\|p(A)\|$ is approximately minimized by $p(z) = (1 z/2)^n$;
- λ(I − A/2) is roughly contained in the disc of radius ¹/₄, centered at z = 0, so the convergence rate is ||p_n(A)|| = ||(I − A/2)ⁿ|| ~ ¹/_{4ⁿ}.
- A is quite well-conditioned: $\kappa(A) = 2.065$.
- A is "not too far" from normal: $\kappa(V) = 216.490$.



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||*p*_n(*A*)|| Example: T&B-35.1 Example: T&B-35.2

- m = 256; b = ones(m,1); th = (0:(m-1))*pi / (m-1);
- A = 2*eye(m) + 0.5 * randn(m)/sqrt(m) + diag(-2+2*sin(th)+i*cos(th));



• The eigenvalue spectrum of A now "surrounds" the origin.

Example: T&B-35.1

Example: T&B-35.2

- A is quite well-conditioned: $\kappa(A) = 3.802$.
- A is not too far from normal: $\kappa(V) = 150.711$.
- The convergence is quite slow in this case (observed $\sim~1.23^{-n}).$
- Note that the slowdown in convergence does not depend on conditioning, but on the location of the eigenvalues.
- Clearly, understanding the impact of the "structure" of the eigenvalue spectrum is a non-trivial topic...



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T&B-35.2⁺

- m = 256; b = ones(m,1); th = 1.5*(0:(m-1))*pi / (m-1);
 - A = 2*eye(m) + 0.5 * randn(m)/sqrt(m) + diag(-2+2*sin(th)+i*cos(th));



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T&B-35.2⁺⁺

m = 256; b = ones(m,1); th = 1.75*(0:(m-1))*pi / (m-1); A = 2*eye(m) + 0.5 * randn(m)/sqrt(m) + diag(-2+2*sin(th)+i*cos(th));



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T&B-35.2+++

||*P*_n(*A*)|| Example: T&B-35.1 Example: T&B-35.2

m = 1024; b = ones(m,1); th = 6.00*(0:(m-1))*pi / (m-1);

A = 2*eye(m) + 0.5 * randn(m)/sqrt(m) + diag(-2+(1+th/(6*pi)).*(2*sin(th)+i*cos(th)));



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