Numerical Methods I Solving Linear Systems: Sparse Matrices, Iterative Methods and Non-Square Systems

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- 2 Iterative Methods (briefly)
- 3 The QR Factorization



Banded Matrices

• Banded matrices are a very special but common type of sparse matrix, e.g., tridiagonal matrices

$$\begin{bmatrix} a_{1} & c_{1} & \mathbf{0} \\ b_{2} & a_{2} & \ddots & \\ & \ddots & \ddots & c_{n-1} \\ \mathbf{0} & b_{n} & a_{n} \end{bmatrix}$$

- There exist special techniques for banded matrices that are much faster than the general case, e.g, only 8*n* FLOPS and no additional memory for tridiagonal matrices.
- A general matrix should be considered sparse if it has sufficiently many zeros that exploiting that fact is advantageous: usually only the case for **large matrices** (what is large?)!

Sparse Matrices

Sparse Matrices

1 2 2 Sparse matrix 3 4 5 4 12 IN [3] 14) DIRECTED Graph representation: a21 -> No DES are variables (VERTICES) equations a42 (4) a_24 3 a13 -> ARCS (EDGES) are aug 14 3 the non-teros 15 UNDIRECTED GRAPH FOR SYMMETRIC MATRICES

Sparse matrices in MATLAB

>> A = sparse(
$$[1 \ 2 \ 2 \ 4 \ 4]$$
, $[3 \ 1 \ 4 \ 2 \ 3]$, 1:5)
A =
(2,1) 2
(4,2) 4
(1,3) 1
(4,3) 5
(2,4) 3
>> nnz(A)
ans = 5
>> whos A
A 4×4 120 double sparse
>> A = sparse([],[],[],4,4,5); % Pre-allocate memory
>> A(2,1)=2; A(4,2)=4; A(1,3)=1; A(4,3)=5; A(2,4)=3;

Sparse matrix factorization

>> B=sprand (4,4,0.25); % Density of 25% >> full(B) ans = 0.7655 0 0 0 0 0.7952 0 0 0 0.18690 0 0.48980 0 0 >> B=sprand (100,100,0.1); spy(B) >> X=gallery('poisson',10); spy(X) >> [L, U, P] = Iu(B); spy(L)>> p = symrcm(B); % Symmetric Reverse Cuthill-McKee of >> PBP=B(p,p); spy(PBP);>> [L,U,P] = Iu(PBP); spy(L);

Sparse Matrices

Random matrix \mathbf{B} and structured matrix \mathbf{X}

The MATLAB function spy shows where the nonzeros are as a plot



LU factors of random matrix B

Fill-in (generation of lots of nonzeros) is large for a random sparse matrix



LU factors of structured matrix X

Fill-in is much smaller for the sparse matrix but still non-negligible.



Matrix reordering

Matrix reordering cannot do much for the random matrix **B**, but it can help for structured ones!



Reducing fill-in by reordering \mathbf{X}

Fill-in was reduced by about 20% (from 1000 nonzeros to 800) by the reordering for the structured X, but does not help much for B. The actual numbers are different for different classes of matrices!

Sparse Matrices



Sparse Matrices

Importance of Sparse Matrix Structure

- Important to remember: While there are general techniques for dealing with sparse matrices that help greatly, it all depends on the structure (origin) of the matrix.
- Pivoting has a dual, sometimes conflicting goal:
 - Reduce fill-in, i.e., improve memory use: Still active subject of research!
 - Reduce roundoff error, i.e., improve stability. Typically some threshold pivoting is used only when needed.
- Pivoting for symmetric non-positive definite matrices is trickier: One can permute the diagonal entries only to **preserve symmetry**, but small diagonal entries require special treatment.
- For many sparse matrices **iterative methods** (briefly covered next lecture) are required to large fill-in.

Why iterative methods?

- Direct solvers are great for dense matrices and can be made to avoid roundoff errors to a large degree. They can also be implemented very well on modern machines.
- Fill-in is a major problem for certain sparse matrices and leads to extreme memory requirements (e.g., three-d.
- Some matrices appearing in practice are **too large** to even be represented explicitly (e.g., the Google matrix).
- Often linear systems only need to be **solved approximately**, for example, the linear system itself may be a linear approximation to a nonlinear problem.
- Direct solvers are much harder to implement and use on (massively) **parallel computers**.

Iterative Methods (briefly)

Stationary Linear Iterative Methods of First Order

- In iterative methods the core computation is iterative matrix-vector multiplication starting from an initial guess x⁽⁰⁾.
- Prototype is the linear recursion:

$$\mathbf{x}^{(k+1)} = \mathbf{B}\mathbf{x}^{(k)} + \mathbf{f},$$

where **B** is an **iteration matrix** somehow related to **A**.

• For this method to be **consistent**, we must have that the actual solution $\mathbf{x} = \mathbf{A}^{-1}\mathbf{b}$ is a **stationary point** of the iteration:

$$\mathbf{x} = \mathbf{B}\mathbf{x} + \mathbf{f} \quad \Rightarrow \quad \mathbf{A}^{-1}\mathbf{b} = \mathbf{B}\mathbf{A}^{-1}\mathbf{b} + \mathbf{f}$$

$$\mathbf{f} = \mathbf{A}^{-1}\mathbf{b} - \mathbf{B}\mathbf{A}^{-1}\mathbf{b} = (\mathbf{I} - \mathbf{B})\mathbf{x}$$

For this method to be stable, and thus convergent, the error
 e^(k) = x^(k) - x must decrease:

$$\mathbf{e}^{(k+1)} = \mathbf{x}^{(k+1)} - \mathbf{x} = \mathbf{B}\mathbf{x}^{(k)} + \mathbf{f} - \mathbf{x} = \mathbf{B}\left(\mathbf{x} + \mathbf{e}^{(k)}\right) + (\mathbf{I} - \mathbf{B})\mathbf{x} - \mathbf{x} = \mathbf{B}\mathbf{e}^{(k)}$$

Convergence of simple iterative methods

• We saw that the error propagates from iteration to iteration as

 $\mathbf{e}^{(k)} = \mathbf{B}^k \mathbf{e}^{(0)}.$

• When does this converge? Taking norms,

$$\left\|\mathbf{e}^{(k)}\right\| \leq \left\|\mathbf{B}\right\|^{k} \left\|\mathbf{e}^{(0)}\right\|$$

which means that $\|\boldsymbol{B}\| < 1$ is a sufficient condition for convergence.

- More precisely, $\lim_{k\to\infty} \mathbf{e}^{(k)} = \mathbf{0}$ for any $\mathbf{e}^{(0)}$ iff $\mathbf{B}^k \to \mathbf{0}$.
- Theorem: The method converges iff the **spectral radius** of the iteration matrix is less than unity:

$$\rho(\mathbf{B}) < 1.$$

Spectral Radius

 The spectral radius ρ(A) of a matrix A can be thought of as the smallest consistent matrix norm

$$ho(\mathbf{A}) = \max_{\lambda} |\lambda| \le \|\mathbf{A}\|$$

 The spectral radius often determines convergence of iterative schemes for linear systems and eigenvalues and even methods for solving PDEs because it estimates the asymptotic rate of error propagation:

$$ho(\mathbf{A}) = \lim_{k o \infty} \left\| \mathbf{A}^k \right\|^{1/k}$$

Termination

- The iterations of an iterative method can be terminated when:
 - The residual becomes small,

$$\left\| \mathbf{r}^{(k)} \right\| \le \varepsilon \left\| \mathbf{b} \right\|$$

This is good for well-conditioned systems.

2 The solution $\mathbf{x}^{(k)}$ stops changing, i.e., the **increment** becomes small,

$$\left[1-\rho(\mathbf{B})\right]\left\|\mathbf{e}^{(k)}\right\| \leq \left\|\mathbf{x}^{(k+1)}-\mathbf{x}^{(k)}\right\| \leq \varepsilon \left\|\mathbf{b}\right\|,$$

which can be seen to be good if convergence is rapid, $\rho(\mathbf{B}) \ll 1$.

• Usually a careful **combination** of the two strategies is employed along with some **safeguards**.

Fixed-Point Iteration

• A naive but often successful method for solving

x = f(x)

is the fixed-point iteration

$$x_{n+1}=f(x_n).$$

• In the case of a linear system, consider rewriting Ax = b as:

$$\mathbf{x} = (\mathbf{I} - \mathbf{A}) \, \mathbf{x} + \mathbf{b}$$

• Fixed-point iteration gives the consistent iterative method

$$\mathbf{x}^{(k+1)} = (\mathbf{I} - \mathbf{A}) \, \mathbf{x}^{(k)} + \mathbf{b}$$

Preconditioning

• The above method is consistent but it may not converge or may converge very slowly

$$\mathbf{x}^{(k+1)} = (\mathbf{I} - \mathbf{A}) \, \mathbf{x}^{(k)} + \mathbf{b}.$$

• As a way to speed it up, consider having a good approximate solver

$$\mathbf{P}^{-1}pprox \mathbf{A}^{-1}$$

called the $\ensuremath{\text{preconditioner}}$ (P is the preconditioning matrix), and transform

$$\mathbf{P}^{-1}\mathbf{A}\mathbf{x} = \mathbf{P}^{-1}\mathbf{b}$$

• Now apply fixed-point iteration to this modified system:

$$\mathbf{x}^{(k+1)} = \left(\mathbf{I} - \mathbf{P}^{-1}\mathbf{A}\right)\mathbf{x}^{(k)} + \mathbf{P}^{-1}\mathbf{b},$$

which now has an iteration matrix $\mathbf{I} - \mathbf{P}^{-1}\mathbf{A} \approx \mathbf{0}$, which means more rapid convergence.

Preconditioned Iteration

$$\mathbf{x}^{(k+1)} = \left(\mathbf{I} - \mathbf{P}^{-1}\mathbf{A}
ight)\mathbf{x}^{(k)} + \mathbf{P}^{-1}\mathbf{b}$$

 In practice, we solve linear systems with the matrix P instead of inverting it:

$$Px^{(k+1)} = (P - A)x^{(k)} + b = Px^{(k)} + r^{(k)},$$

where $\mathbf{r}^{(k)} = \mathbf{b} - \mathbf{A}\mathbf{x}^{(k)}$ is the **residual vector**.

• Finally, we obtain the usual form of a preconditioned stationary iterative solver

$$\mathbf{x}^{(k+1)} = \mathbf{x}^{(k)} + \mathbf{P}^{-1}\mathbf{r}^{(k)}.$$

• Note that convergence will be faster if we have a good initial guess $\mathbf{x}^{(0)}$.

Some Standard Examples

Splitting: $\mathbf{A} = \mathbf{L}_{A} + \mathbf{U}_{A} + \mathbf{D}$

• Since diagonal systems are trivial to solve, we can use the **Jacobi method**

$\mathbf{P} = \mathbf{D}.$

• Or since triangular systems are easy to solve by forward/backward substitution, we can use **Gauss-Seidel method**

$$\mathbf{P}=\mathbf{L}_{\mathcal{A}}+\mathbf{D}.$$

- Both of these converge for strictly diagonally-dominant matrices.
- Gauss-Seidel converges for **positive-definite matrices** (maybe slowly though!).

A Good Preconditioner

- Note that the matrix **A** is only used when calculating the residual through the **matrix-vector product** $Ax^{(k)}$.
- We must be able to do a **direct** linear solver for the preconditioner

$$\mathsf{P}\left(\Delta \mathsf{x}\right) = \mathsf{r}^{(k)},$$

so it must be in some sense simpler to deal with than **A**.

- Preconditioning is all about a **balance between fewer iterations to convergence and larger cost per iteration**.
- Making good preconditioners is in many ways an art and very **problem-specific**:

The goal is to make $P^{-1}A$ as close to being a normal (diagonalizable) matrix with **clustered eigenvalues** as possible.

In the Real World

- Some general preconditioning strategies have been designed, for example, **incomplete** *LU* **factorization** (MATLAB's *cholinc*).
- There are many **more-sophisticated iterative methods** (non-stationary, higher-order, etc) but most have the same **basic structure**:

At each iteration, solve a preconditioning linear system, do a matrix-vector calculation, and a convergence test.

- For positive-(semi)definite matrices the **Preconditioned Conjugate Gradient** method is good (MATLAB's *pcg*).
- For certain types of matrices specialized methods have been designed, such as **multigrid methods** for linear systems on large grids (PDE solvers in Numerical Methods II).

Non-Square Matrices

- In the case of **over-determined** (more equations than unknowns) or **under-determined** (more unknowns than equations), the solution to linear systems in general becomes **non-unique**.
- One must first define what is meant by a solution, and the common definition is to use a **least-squares formulation**:

$$\mathbf{x}^{\star} = \arg\min_{\mathbf{x}\in\mathbb{R}^n} \|\mathbf{A}\mathbf{x} - \mathbf{b}\|_2^2 = \arg\min_{\mathbf{x}\in\mathbb{R}^n} \Phi(\mathbf{x})$$

where the quadratic form is

$$\Phi(\mathbf{x}) = \left(\mathbf{A}\mathbf{x} - \mathbf{b}\right)^T \left(\mathbf{A}\mathbf{x} - \mathbf{b}\right).$$

- Sometimes the solution to the least-squares is still not unique:
 - Under-determined systems (not enough equations to fix all unknowns)
 - Singular systems, i.e., A that is not of full rank: Any solution to $A x_0 = 0$ can be added to x without changing the left hand side!
- Additional condition: Choose the **x*** that has **minimal Euclidean norm**.

Over-determined systems: Normal Equations

• Over-determined systems, m > n, can be thought of as fitting a linear model (linear regression):

The unknowns \mathbf{x} are the coefficients in the fit, the input data is in \mathbf{A} (one column per measurement), and the output data (observables) are in \mathbf{b} .

• Not worrying about technicalities, set the gradient to zero:

$$\Phi(\mathbf{x}) = (\mathbf{A}\mathbf{x} - \mathbf{b})^T (\mathbf{A}\mathbf{x} - \mathbf{b})$$
 and $\nabla \Phi(\mathbf{x}^{\star}) = \mathbf{0}$

 $abla \Phi(\mathbf{x}) = \mathbf{A}^{T} \left[2 \left(\mathbf{A} \mathbf{x} - \mathbf{b} \right) \right]$ (calculus with care for order and shapes)

• This gives the square linear system of normal equations

$$(\mathbf{A}^{\mathsf{T}}\mathbf{A})\mathbf{x}^{\star} = \mathbf{A}^{\mathsf{T}}\mathbf{b}.$$

If A is of full rank, rank (A) = n, it can be shown (do it!) that A^TA is positive definite, and Cholesky factorization can be used to solve the normal equations.

The QR Factorization

Problems with the normal equations

$$\left(\mathbf{A}^{\mathcal{T}} \mathbf{A}
ight) \mathbf{x}^{\star} = \mathbf{A}^{\mathcal{T}} \mathbf{b}.$$

• The conditioning number of the normal equations is

$$\kappa\left(\mathbf{A}^{T}\mathbf{A}\right) = \left[\kappa(\mathbf{A})\right]^{2}$$

- Furthermore, roundoff can cause **A**^T**A** to no longer appear as positive-definite and the Cholesky factorization will fail.
- If the normal equations are ill-conditioned, another approach is needed.
- Also note that multiplying \mathbf{A}^T $(n \times m)$ and \mathbf{A} $(m \times n)$ takes n^2 dot-products of length m, so $O(mn^2)$ operations, which can be much larger than $O(n^3)$ for the Cholesky factorization if $m \gg n$ (fitting lots of data with few parameters).

The QR factorization

 For nonsquare or ill-conditioned matrices of full-rank r = n ≤ m, the LU factorization can be replaced by the QR factorization:

 $\mathbf{A} = \mathbf{Q}\mathbf{R}$ $[m \times n] = [m \times n][n \times n]$

where **Q** has orthogonal columns, $\mathbf{Q}^T \mathbf{Q} = \mathbf{I}_n$, and **R** is a non-singular upper triangular matrix.

- Observe that orthogonal / unitary matrices are **well-conditioned** $(\kappa_2 = 1)$, so the *QR* factorization is numerically better (but also more expensive!) than the *LU* factorization.
- For matrices **not of full rank** there are modified *QR* factorizations but **the SVD decomposition is better** (next class).
- In MATLAB, the *QR* factorization can be computed using *qr* (with column pivoting).

The QR Factorization

Solving Linear Systems via QR factorization

$$\left(\mathbf{A}^{\mathcal{T}}\mathbf{A}
ight) \mathbf{x}^{\star} = \mathbf{A}^{\mathcal{T}}\mathbf{b}$$
 where $\mathbf{A} = \mathbf{Q}\mathbf{R}$

• Observe that **R** is the Cholesky factor of the matrix in the normal equations:

$$\mathbf{A}^{\mathsf{T}}\mathbf{A} = \mathbf{R}^{\mathsf{T}}\left(\mathbf{Q}^{\mathsf{T}}\mathbf{Q}\right)\mathbf{R} = \mathbf{R}^{\mathsf{T}}\mathbf{R}$$

$$\left(\mathbf{R}^{\mathcal{T}} \mathbf{R} \right) \mathbf{x}^{\star} = \left(\mathbf{R}^{\mathcal{T}} \mathbf{Q}^{\mathcal{T}} \right) \mathbf{b} \quad \Rightarrow \quad \mathbf{x}^{\star} = \mathbf{R}^{-1} \left(\mathbf{Q}^{\mathcal{T}} \mathbf{b} \right)$$

which amounts to solving a triangular system with matrix \mathbf{R} .

- This calculation turns out to be much **more numerically stable** against roundoff than forming the normal equations (and has similar cost).
- For under-determined full-rank systems, r = m ≤ n, one does a QR factorization of A^T = Q̃R̃ and the least-squares solution is

$$\mathbf{x}^{\star} = \tilde{\mathbf{Q}} \left(\tilde{\mathbf{R}}^{- \, \mathcal{T}} \mathbf{b} \right)$$

Practice: Derive the above formula and maybe prove least-squares. A. Doney (Courant Institute) Lecture III 9/23/2010 28 / 31

Computing the QR Factorization

Assume that

$$\exists x \text{ s.t. } b = Ax$$
, that is, $b \in range(A)$

$$\mathbf{b} = \mathbf{Q} \left(\mathbf{R} \mathbf{x}
ight) = \mathbf{Q} \mathbf{y} \quad \Rightarrow \quad \mathbf{x} = \mathbf{R}^{-1} \mathbf{y}$$

showing that the columns of ${\bf Q}$ form an **orthonormal basis** for the range of ${\bf A}$ (linear subspace spanned by the columns of ${\bf A}$).

- The *QR* factorization is thus closely-related to the **orthogonalization** of a set of *n* vectors (columns) {**a**₁, **a**₂,..., **a**_n} in \mathbb{R}^m .
- Classical approach is the Gram-Schmidt method: To make a vector
 b orthogonal to a do:

$$ilde{\mathbf{b}} = \mathbf{b} - (\mathbf{b} \cdot \mathbf{a}) \, rac{\mathbf{a}}{(\mathbf{a} \cdot \mathbf{a})}$$

Practice: Verify that $\tilde{\boldsymbol{b}}\cdot\boldsymbol{a}=\boldsymbol{0}$

• Repeat this in sequence: Start with $\tilde{a}_1 = a_1$, then make \tilde{a}_2 orthogonal to \tilde{a}_1 , then make \tilde{a}_3 orthogonal to \tilde{a}_2 and \tilde{a}_3 .

Modified Gram-Schmidt Orthogonalization

• More efficient formula (standard Gram-Schmidt):

The OR Eactorization

$$\tilde{\mathbf{a}}_{k+1} = \mathbf{a}_{k+1} - \sum_{j=1}^{k} \left(\mathbf{a}_{k+1} \cdot \mathbf{q}_{j} \right) \mathbf{q}_{j}, \quad \mathbf{q}_{k+1} = \frac{\tilde{\mathbf{a}}_{k+1}}{\|\tilde{\mathbf{a}}_{k+1}\|},$$

with cost $\sim mn^2$ FLOPS.

A mathematically-equivalent but numerically much superior against roundoff error is the modified Gram-Schmidt, in which each orthogonalization is carried in sequence and repeated against each of the already-computed basis vectors:
 Start with ã₁ = a₁, then make ã₂ orthogonal to ã₁, then make ã₃

orthogonal to \tilde{a}_2 and then make it orthogonal to \tilde{a}_3 .

- The modified procedure is twice more expensive, $\sim 2mn^2$ FLOPS, but usually worth it.
- **Pivoting** is strictly necessary for matrices not of full rank but it can also improve stability in general.

Conclusions

Conclusions/Summary

- **Sparse matrices** deserve special treatment but the details depend on the specific field of application.
- In particular, special sparse **matrix reordering** methods or iterative systems are often required.
- When **sparse direct methods** fail due to memory or other requirements, **iterative methods** are used instead.
- Convergence of iterative methods depends strongly on the matrix, and a good **preconditioner** is often required.
- There are **good libraries for iterative methods** as well (but you must supply your own preconditioner!).
- The *QR* factorization is a numerically-stable method for solving **full-rank non-square systems**.
- For rank-defficient matrices the singular value decomposition (SVD) is best, discussed in later lectures.