Scientific Computing: Solving Linear Systems

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Outline

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Linear Spaces

 A vector space V is a set of elements called vectors x ∈ V that may be multiplied by a scalar c and added, e.g.,

$$\mathbf{z} = \alpha \mathbf{x} + \beta \mathbf{y}$$

- I will denote scalars with lowercase letters and vectors with lowercase bold letters.
- Prominent examples of vector spaces are ℝⁿ (or more generally ℂⁿ), but there are many others, for example, the set of polynomials in x.
- A subspace V' ⊆ V of a vector space is a subset such that sums and multiples of elements of V' remain in V' (i.e., it is closed).
- An example is the set of vectors in $x \in \mathbb{R}^3$ such that $x_3 = 0$.

Image Space

• Consider a set of *n* vectors $\mathbf{a}_1, \mathbf{a}_2, \cdots, \mathbf{a}_n \in \mathbb{R}^m$ and form a **matrix** by putting these vectors as columns

$$\mathbf{A} = [\mathbf{a}_1 \,|\, \mathbf{a}_2 \,|\, \cdots \,|\, \mathbf{a}_m] \in \mathbb{R}^{m,n}.$$

- I will denote matrices with bold capital letters, and sometimes write $\mathbf{A} = [m, n]$ to indicate dimensions.
- The matrix-vector product is defined as a linear combination of the columns:

$$\mathbf{b} = \mathbf{A}\mathbf{x} = x_1\mathbf{a}_1 + x_2\mathbf{a}_2 + \cdots + x_n\mathbf{a}_n \in \mathbb{R}^m.$$

The image im(A) or range range(A) of a matrix is the subspace of all linear combinations of its columns, i.e., the set of all b's.
 It is also sometimes called the column space of the matrix.

Dimension

- The set of vectors a₁, a₂, · · · , a_n are linearly independent or form a basis for ℝ^m if b = Ax = 0 implies that x = 0.
- The dimension r = dimV of a vector (sub)space V is the number of elements in a basis. This is a property of V itself and not of the basis, for example,

$$\dim \mathbb{R}^n = n$$

Given a basis A for a vector space V of dimension n, every vector of b ∈ V can be uniquely represented as the vector of coefficients x in that particular basis,

$$\mathbf{b} = x_1\mathbf{a}_1 + x_2\mathbf{a}_2 + \cdots + x_n\mathbf{a}_n.$$

A simple and common basis for ℝⁿ is {e₁,..., e_n}, where e_k has all components zero except for a single 1 in position k. With this choice of basis the coefficients are simply the entries in the vector, b ≡ x.

Kernel Space

• The dimension of the column space of a matrix is called the **rank** of the matrix $\mathbf{A} \in \mathbb{R}^{m,n}$,

$$r = \operatorname{rank} \mathbf{A} \leq \min(m, n).$$

- If $r = \min(m, n)$ then the matrix is of **full rank**.
- The **nullspace** null(**A**) or **kernel** ker(**A**) of a matrix **A** is the subspace of vectors **x** for which

$$Ax = 0.$$

- The dimension of the nullspace is called the nullity of the matrix.
- For a basis **A** the nullspace is $null(\mathbf{A}) = \{\mathbf{0}\}$ and the nullity is zero.

Orthogonal Spaces

- An inner-product space is a vector space together with an **inner or dot product**, which must satisfy some properties.
- The standard dot-product in \mathbb{R}^n is denoted with several different notations:

$$\mathbf{x} \cdot \mathbf{y} = (\mathbf{x}, \mathbf{y}) = \langle \mathbf{x}, \mathbf{y} \rangle = \mathbf{x}^T \mathbf{y} = \sum_{i=1}^n x_i y_i.$$

For Cⁿ we need to add complex conjugates (here ★ denotes a complex conjugate transpose, or adjoint),

$$\mathbf{x} \cdot \mathbf{y} = \mathbf{x}^* \mathbf{y} = \sum_{i=1}^n \bar{x}_i y_i.$$

• Two vectors \mathbf{x} and \mathbf{y} are **orthogonal** if $\mathbf{x} \cdot \mathbf{y} = \mathbf{0}$.

Part I of Fundamental Theorem

• One of the most important theorems in linear algebra is that the sum of rank and nullity is equal to the number of columns: For $\mathbf{A} \in \mathbb{R}^{m,n}$

rank
$$\mathbf{A}$$
 + nullity \mathbf{A} = n .

- In addition to the range and kernel spaces of a matrix, two more important vector subspaces for a given matrix **A** are the:
 - **Row space** or **coimage** of a matrix is the column (image) space of its transpose, im **A**^T.

Its dimension is also equal to the the rank.

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• Left nullspace or cokernel of a matrix is the nullspace or kernel of its transpose, ker A^T.

Part II of Fundamental Theorem

- The orthogonal complement V[⊥] or orthogonal subspace of a subspace V is the set of all vectors that are orthogonal to every vector in V.
- Let \mathcal{V} be the set of vectors in $x \in \mathbb{R}^3$ such that $x_3 = 0$. Then \mathcal{V}^{\perp} is the set of all vectors with $x_1 = x_2 = 0$.
- Second fundamental theorem in linear algebra:

im
$$\mathbf{A}^T = (\ker \mathbf{A})^{\perp}$$

ker $\mathbf{A}^T = (\operatorname{im} \mathbf{A})^{\perp}$

Linear Transformation

A function L : V → W mapping from a vector space V to a vector space W is a linear function or a linear transformation if

$$L(\alpha \mathbf{v}) = \alpha L(\mathbf{v})$$
 and $L(\mathbf{v}_1 + \mathbf{v}_2) = L(\mathbf{v}_1) + L(\mathbf{v}_2)$.

 Any linear transformation L can be represented as a multiplication by a matrix L

$$L(\mathbf{v}) = \mathbf{L}\mathbf{v}.$$

For the common bases of V = Rⁿ and W = R^m, the product w = Lv is simply the usual matix-vector product,

$$w_i = \sum_{k=1}^n L_{ik} v_k,$$

which is simply the dot-product between the *i*-th row of the matrix and the vector \mathbf{v} .

Matrix algebra

$$w_i = (\mathbf{L}\mathbf{v})_i = \sum_{k=1}^n L_{ik} v_k$$

• The composition of two linear transformations $\mathbf{A} = [m, p]$ and $\mathbf{B} = [p, n]$ is a matrix-matrix product $\mathbf{C} = \mathbf{AB} = [m, n]$:

$$z = A(Bx) = Ay = (AB)x$$

$$z_{i} = \sum_{k=1}^{n} A_{ik} y_{k} = \sum_{k=1}^{p} A_{ik} \sum_{j=1}^{n} B_{kj} x_{j} = \sum_{j=1}^{n} \left(\sum_{k=1}^{p} A_{ik} B_{kj} \right) x_{j} = \sum_{j=1}^{n} C_{ij} x_{j}$$
$$C_{ij} = \sum_{k=1}^{p} A_{lk} B_{kj}$$

Matrix-matrix multiplication is not commutative, AB \neq BA in general.

The Matrix Inverse

• A square matrix $\mathbf{A} = [n, n]$ is invertible or nonsingular if there exists a matrix inverse $\mathbf{A}^{-1} = \mathbf{B} = [n, n]$ such that:

$$AB = BA = I$$
,

where I is the identity matrix (ones along diagonal, all the rest zeros).

- The following statements are equivalent for $\mathbf{A} \in \mathbb{R}^{n,n}$:
 - A is invertible.
 - A is full-rank, rank A = n.
 - The columns and also the rows are linearly independent and form a basis for ℝⁿ.
 - The **determinant** is nonzero, det $\mathbf{A} \neq \mathbf{0}$.
 - Zero is not an eigenvalue of A.

Matrix Algebra

 Matrix-vector multiplication is just a special case of matrix-matrix multiplication. Note x^Ty is a scalar (dot product).

$$\mathbf{C}\left(\mathbf{A}+\mathbf{B}
ight)=\mathbf{C}\mathbf{A}+\mathbf{C}\mathbf{B}$$
 and $\mathbf{ABC}=\left(\mathbf{AB}
ight)\mathbf{C}=\mathbf{A}\left(\mathbf{BC}
ight)$

$$(\mathbf{A}^{T})^{T} = \mathbf{A} \text{ and } (\mathbf{A}\mathbf{B})^{T} = \mathbf{B}^{T}\mathbf{A}^{T}$$

$$\left(\mathbf{A}^{-1}\right)^{-1} = \mathbf{A} \text{ and } \left(\mathbf{A}\mathbf{B}\right)^{-1} = \mathbf{B}^{-1}\mathbf{A}^{-1} \text{ and } \left(\mathbf{A}^{\mathcal{T}}\right)^{-1} = \left(\mathbf{A}^{-1}\right)^{\mathcal{T}}$$

• Instead of matrix division, think of multiplication by an inverse:

$$\mathbf{A}\mathbf{B} = \mathbf{C} \quad \Rightarrow \quad \left(\mathbf{A}^{-1}\mathbf{A}\right)\mathbf{B} = \mathbf{A}^{-1}\mathbf{C} \quad \Rightarrow \quad \begin{cases} \mathbf{B} &= \mathbf{A}^{-1}\mathbf{C} \\ \mathbf{A} &= \mathbf{C}\mathbf{B}^{-1} \end{cases}$$

Vector norms

- Norms are the abstraction for the notion of a length or magnitude.
- For a vector $\mathbf{x} \in \mathbb{R}^n$, the *p*-norm is

$$\|\mathbf{x}\|_{p} = \left(\sum_{i=1}^{n} |x_{i}|^{p}\right)^{1/p}$$

and special cases of interest are:

- The 1-norm (L¹ norm or Manhattan distance), ||**x**||₁ = ∑_{i=1}ⁿ |x_i|
 The 2-norm (L² norm, Euclidian distance), ||**x**||₂ = √**x** · **x** = √∑_{i=1}ⁿ |x_i|²
 The ∞-norm (L[∞] or maximum norm), ||**x**||_∞ = max_{1≤i≤n} |x_i|
- Note that all of these norms are inter-related in a finite-dimensional setting.

Matrix norms

Matrix norm induced by a given vector norm:

$$\|\mathbf{A}\| = \sup_{\mathbf{x} \neq \mathbf{0}} \frac{\|\mathbf{A}\mathbf{x}\|}{\|\mathbf{x}\|} \quad \Rightarrow \|\mathbf{A}\mathbf{x}\| \le \|\mathbf{A}\| \|\mathbf{x}\|$$

- The last bound holds for matrices as well, $\|\mathbf{AB}\| \le \|\mathbf{A}\| \|\mathbf{B}\|$.
- Special cases of interest are:

• The 1-norm or column sum norm, $\|\mathbf{A}\|_1 = \max_j \sum_{i=1}^n |a_{ij}|$ • The ∞ -norm or row sum norm, $\|\mathbf{A}\|_{\infty} = \max_j \sum_{i=1}^n |a_{ij}|$

- **3** The 2-norm or **spectral norm**, $\|\mathbf{A}\|_2 = \sigma_1$ (largest singular value)
- The Euclidian or **Frobenius norm**, $\|\mathbf{A}\|_{F} = \sqrt{\sum_{i,j} |a_{ij}|^{2}}$ (note this is not an induced norm)

Matrices and linear systems

Conditioning of linear systems

• It is said that 70% or more of applied mathematics research involves solving systems of *m* linear equations for *n* unknowns:

$$\sum_{j=1}^n a_{ij}x_j = b_i, \quad i = 1, \cdots, m.$$

• Linear systems arise directly from **discrete models**, e.g., traffic flow in a city. Or, they may come through representing or more abstract **linear operators** in some finite basis (representation). Common abstraction:

$$Ax = b$$

• Special case: Square invertible matrices, m = n, det $\mathbf{A} \neq 0$:

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

• The goal: Calculate solution **x** given data **A**, **b** in the most numerically stable and also efficient way.

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Stability analysis

Perturbations on right hand side (rhs) only:

$$\mathbf{A} (\mathbf{x} + \delta \mathbf{x}) = \mathbf{b} + \delta \mathbf{b} \quad \Rightarrow \mathbf{b} + \mathbf{A} \delta \mathbf{x} = \mathbf{b} + \delta \mathbf{b}$$

$$\delta \mathbf{x} = \mathbf{A}^{-1} \delta \mathbf{b} \quad \Rightarrow \|\delta \mathbf{x}\| \le \|\mathbf{A}^{-1}\| \|\delta \mathbf{b}\|$$

Using the bounds

$$\left\| \mathbf{b} \right\| \leq \left\| \mathbf{A} \right\| \left\| \mathbf{x} \right\| \quad \Rightarrow \left\| \mathbf{x} \right\| \geq \left\| \mathbf{b} \right\| / \left\| \mathbf{A} \right\|$$

the relative error in the solution can be bounded by

$$\frac{\|\delta \mathbf{x}\|}{\|\mathbf{x}\|} \leq \frac{\|\mathbf{A}^{-1}\| \|\delta \mathbf{b}\|}{\|\mathbf{x}\|} \leq \frac{\|\mathbf{A}^{-1}\| \|\delta \mathbf{b}\|}{\|\mathbf{b}\| / \|\mathbf{A}\|} = \kappa(\mathbf{A}) \frac{\|\delta \mathbf{b}\|}{\|\mathbf{b}\|}$$

where the **conditioning number** $\kappa(\mathbf{A})$ depends on the matrix norm used:

$$\kappa(\mathbf{A}) = \|\mathbf{A}\| \|\mathbf{A}^{-1}\| \ge 1.$$

Conditioning Number

• The full derivation, not given here, estimates the uncertainty or perturbation in the solution:

$$\frac{\|\delta \mathbf{x}\|}{\|\mathbf{x}\|} \leq \frac{\kappa(\mathbf{A})}{1 - \kappa(\mathbf{A}) \frac{\|\delta \mathbf{A}\|}{\|\mathbf{A}\|}} \left(\frac{\|\delta \mathbf{b}\|}{\|\mathbf{b}\|} + \frac{\|\delta \mathbf{A}\|}{\|\mathbf{A}\|} \right).$$

The **worst-case conditioning** of the linear system is determined by $\kappa(\mathbf{A})$.

• Best possible error with rounding unit $u \approx 10^{-16}$:

Conditioning of linear systems

$$\frac{\left\|\delta\mathbf{x}\right\|_{\infty}}{\left\|\mathbf{x}\right\|_{\infty}} \lesssim 2u\kappa(\mathbf{A}),$$

- Solving an ill-conditioned system, $\kappa(\mathbf{A}) \gg 1$ (e.g., $\kappa = 10^{15}$!), should only be done if something special is known.
- The conditioning number can only be **estimated** in practice since **A**⁻¹ is not available (see MATLAB's *rcond* function).

GEM: Eliminating x_1

Step 1:

$$A = G$$

$$\begin{bmatrix} a_{AA}^{(A)} & a_{A2}^{(A)} & a_{A5}^{(A)} \\ \hline a_{2A}^{(A)} & a_{22}^{(A)} & a_{35}^{(A)} \\ \hline a_{3A}^{(A)} & a_{32}^{(A)} & a_{35}^{(A)} \\ \hline \end{bmatrix} \begin{bmatrix} x_{A} \\ \hline x_{L} \\ x_{L} \\ \hline x$$

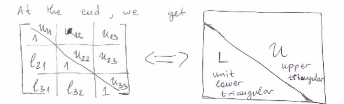
GEM: Eliminating x_2

Step 2: Q (1) **a**13⁽¹⁾ ×1 ×2 = (2)(2)(2)(2)(3)a(2) 22 Xz a⁽²⁾ 11 a(1) a(1) 12 Q (1) 13 $a_{23}^{(2)}$ a⁽²⁾

GEM: Backward substitution

 $\begin{array}{c} \text{Eliminate} \\ x_{3} \\ \text{entirely} \end{array} \begin{bmatrix} a_{11}^{(1)} & a_{12}^{(1)} \\ 0 & a_{22}^{(2)} \\ \text{solve for } x_{2} \\ \end{bmatrix} \begin{bmatrix} x_{1} \\ x_{2} \\ x_{3} \\ x_{2} \\ \end{bmatrix} = \begin{bmatrix} e_{1}^{(3)} - a_{13}^{(1)} \\ x_{3} \\ e_{2}^{(3)} - a_{23}^{(2)} \\ x_{3} \\ \end{bmatrix} = \begin{bmatrix} e_{1}^{(3)} \\ e_{2}^{(3)} - a_{23}^{(2)} \\ x_{3} \\ \end{bmatrix} = \begin{bmatrix} e_{1}^{(3)} \\ e_{2}^{(3)} \\$ IDEA: Store the multipliers in the lower triangle of A: Matrix U12 at Step k: UM U 13 $\begin{array}{c|c} l_{21} & a_{22}^{(2)} & a_{23}^{(2)} \\ \hline l_{11} & a_{23}^{(2)} & a_{33}^{(2)} \end{array}$ (k) l_{31} $a_{32}^{(2)}$ Example step 2 2

GEM as an LU factorization tool



• We have actually factorized A as

$$\mathbf{A} = \mathbf{L}\mathbf{U},$$

L is unit lower triangular ($l_{ii} = 1$ on diagonal), and **U** is upper triangular.

• GEM is thus essentially the same as the LU factorization method.

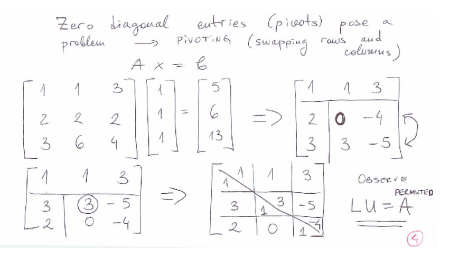
GEM in MATLAB

Sample MATLAB code (for learning purposes only, not real computing!): function A = MyLU(A)% LU factorization in-place (overwrite A) [n,m] = size(A);if (n = m); error ('Matrix not square'); end for k=1:(n-1) % For variable x(k)% Calculate multipliers in column k: A((k+1):n,k) = A((k+1):n,k) / A(k,k);% Note: Pivot element A(k,k) assumed nonzero! for i = (k+1):n% Eliminate variable x(k): $A((k+1):n,i) = A((k+1):n,i) - \dots$ A((k+1):n,k) * A(k,j);end

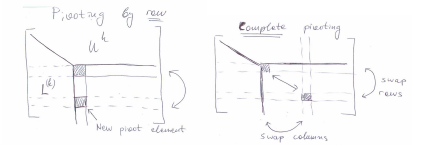
end

end

Pivoting



Pivoting during LU factorization



• **Partial (row) pivoting** permutes the rows (equations) of **A** in order to ensure sufficiently large pivots and thus numerical stability:

$\mathbf{PA} = \mathbf{LU}$

- Here **P** is a **permutation matrix**, meaning a matrix obtained by permuting rows and/or columns of the identity matrix.
- Complete pivoting also permutes columns, PAQ = LU.

Gauss Elimination Method (GEM)

- GEM is a general method for dense matrices and is commonly used.
- Implementing GEM efficiently and stably is difficult and we will not discuss it here, since others have done it for you!
- The **LAPACK** public-domain library is the main repository for excellent implementations of dense linear solvers.
- MATLAB uses a highly-optimized variant of GEM by default, mostly based on LAPACK.
- MATLAB does have **specialized solvers** for special cases of matrices, so always look at the help pages!

Solving linear systems

• Once an LU factorization is available, solving a linear system is simple:

$$Ax = LUx = L(Ux) = Ly = b$$

so solve for y using forward substitution.

This was implicitly done in the example above by overwriting ${\bf b}$ to become ${\bf y}$ during the factorization.

• Then, solve for x using backward substitution

$$\mathbf{U}\mathbf{x} = \mathbf{y}$$
.

• If row pivoting is necessary, the same applies but **L** or **U** may be permuted upper/lower triangular matrices,

$$\mathbf{A} = \widetilde{\mathbf{L}}\mathbf{U} = \left(\mathbf{P}^{\mathsf{T}}\mathbf{L}\right)\mathbf{U}.$$

In MATLAB

• In MATLAB, the **backslash operator** (see help on *mldivide*)

$$x = A \setminus b \approx A^{-1}b,$$

solves the linear system $\mathbf{A}\mathbf{x} = \mathbf{b}$ using the LAPACK library. Never use matrix inverse to do this, even if written as such on paper.

• Doing $x = A \setminus b$ is **equivalent** to performing an *LU* factorization and doing two **triangular solves** (backward and forward substitution):

$$\begin{split} [\tilde{L}, U] &= lu(A) \\ y &= \tilde{L} \backslash b \\ x &= U \backslash y \end{split}$$

• This is a carefully implemented **backward stable** pivoted LU factorization, meaning that the returned solution is as accurate as the conditioning number allows.

GEM Matlab example (1)

>> A = [1 2 3; 4 5 6; 7 8 0]; >> b=[2 1 -1]';

>> x=A^(-1)*b; x' % Don't do this! ans = -2.5556 2.1111 0.1111

>> x = A\b; x' % Do this instead ans = -2.5556 2.1111 0.1111

>> linsolve (A,b)' % Even more control ans = -2.5556 2.1111 0.1111

Gauss elimination and LU factorization

GEM Matlab example (2)

>> [L,U] = Iu(A) % Even better if resolving

L =	0.1429	1.0000	0
	0.5714	0.5000	1.0000
	1.0000	0	0
U =	7.0000	8.0000	0
	0	0.8571	3.0000
	0	0	4.5000

>>
$$y = L \setminus b$$
;
>> $x = U \setminus y$; x'
ans = -2.5556 2.1111 0.1111

Cost estimates for GEM

• For forward or backward substitution, at step k there are $\sim (n - k)$ multiplications and subtractions, plus a few divisions. The total over all n steps is

$$\sum_{k=1}^n (n-k) = \frac{n(n-1)}{2} \approx \frac{n^2}{2}$$

subtractions and multiplications, giving a total of $O(n^2)$ floating-point operations (FLOPs).

• The LU factorization itself costs a lot more, $O(n^3)$,

$$\mathsf{FLOPS} \approx \frac{2n^3}{3},$$

and the triangular solves are negligible for large systems.

• When many linear systems need to be solved with the same **A** the **factorization can be reused**.

Matrix Rescaling and Reordering

- Pivoting is not always sufficient to ensure lack of roundoff problems. In particular, **large variations** among the entries in **A should be avoided**.
- This can usually be remedied by changing the physical units for x and b to be the natural units x₀ and b₀.
- **Rescaling** the unknowns and the equations is generally a good idea even if not necessary:

$$\mathbf{x} = \mathbf{D}_x \tilde{\mathbf{x}} = \text{Diag} \{\mathbf{x}_0\} \, \tilde{\mathbf{x}} \text{ and } \mathbf{b} = \mathbf{D}_b \tilde{\mathbf{b}} = \text{Diag} \{\mathbf{b}_0\} \, \tilde{\mathbf{b}}.$$

$$\mathbf{A}\mathbf{x} = \mathbf{A}\mathbf{D}_{x}\tilde{\mathbf{x}} = \mathbf{D}_{b}\tilde{\mathbf{b}} \quad \Rightarrow \quad \left(\mathbf{D}_{b}^{-1}\mathbf{A}\mathbf{D}_{x}\right)\tilde{\mathbf{x}} = \tilde{\mathbf{b}}$$

- The rescaled matrix \$\tilde{A} = D_b^{-1}AD_x\$ should have a better conditioning.
- Also note that **reordering the variables** from most important to least important may also help.

Efficiency of Solution

 $\mathbf{A}\mathbf{x} = \mathbf{b}$

- The most appropriate algorithm really depends on the properties of the matrix **A**:
 - General **dense matrices**, where the entries in **A** are mostly non-zero and nothing special is known: Use *LU* factorization.
 - Symmetric $(a_{ij} = a_{ji})$ and also positive-definite matrices.
 - General **sparse matrices**, where only a small fraction of $a_{ij} \neq 0$.
 - Special **structured sparse matrices**, arising from specific physical properties of the underlying system.
- It is also important to consider **how many times** a linear system with the same or related matrix or right hand side needs to be solved.

Positive-Definite Matrices

- A real symmetric matrix **A** is positive definite iff (if and only if):
 - It is eigenvalues are real (follows from symmetry) and positive.
 - **2** $\forall x \neq \mathbf{0}, \mathbf{x}^T \mathbf{A} \mathbf{x} > 0$, i.e., the quadratic form defined by the matrix **A** is convex.
 - **③** There exists a *unique* lower triangular L, $L_{ii} > 0$,

$$\mathbf{A} = \mathbf{L}\mathbf{L}^{T},$$

termed the **Cholesky factorization** of **A** (symmetric *LU* factorization).

● For Hermitian complex matrices just replace transposes with adjoints (conjugate transpose), e.g., A^T → A^{*} (or A^H in the book).

Cholesky Factorization

• The MATLAB built in function

R = chol(A)

gives the Cholesky factorization and is a good way to **test for positive-definiteness**.

- The cost of a Cholesky factorization is about half the cost of LU factorization, $n^3/3$ FLOPS.
- Solving linear systems is as for LU factorization, replacing **U** with L^{T} .
- For Hermitian/symmetric matrices with positive diagonals MATLAB tries a Cholesky factorization first, *before* resorting to *LU* factorization with pivoting.

Special Matrices in MATLAB

- MATLAB recognizes (i.e., tests for) some special matrices automatically: banded, permuted lower/upper triangular, symmetric, Hessenberg, but **not** sparse.
- In MATLAB one may specify a matrix **B** instead of a single right-hand side vector **b**.
- The MATLAB function

$$X = linsolve(A, B, opts)$$

allows one to specify certain properties that speed up the solution (triangular, upper Hessenberg, symmetric, positive definite, none), and also estimates the condition number along the way.

• Use *linsolve* instead of backslash if you know (for sure!) something about your matrix.

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**:

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

where the choice of the L_2 norm leads to:

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

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} .

Normal Equations

• It can be shown that the least-squares solution satisfies:

$$oldsymbol{
abla} \Phi({f x}) = {f A}^{\mathcal{T}} \left[2 \left({f A} {f x} - {f b}
ight)
ight] = {f 0} \; ({\sf critical point})$$

• This gives the square linear system of normal equations

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

- If **A** is of full rank, rank (**A**) = *n*, it can be shown that **A**^T**A** is positive definite, and Cholesky factorization can be used to solve the normal equations.
- Multiplying A^T (n × m) and A (m × n) takes n² dot-products of length m, so O(mn²) operations

Problems with the normal equations

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

• The conditioning number of the normal equations is

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

- 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.

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).

Overdetermined Linear Systems

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:

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

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

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).

Computing the QR Factorization

- The QR factorization is closely-related to the orthogonalization of a set of n vectors (columns) {a₁, a₂,..., a_n} in ℝ^m, which is a common problem in numerical computing.
- Classical approach is the Gram-Schmidt method: To make a vector
 b orthogonal to a do:

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

• Repeat this in sequence: Start with $\tilde{a}_1 = a_1$, then make \tilde{a}_2 orthogonal to $\tilde{a}_1 = a_1$, then make \tilde{a}_3 orthogonal to span $(\tilde{a}_1, \tilde{a}_2) = \text{span}(a_1, a_2)$:

$$\begin{split} \tilde{\mathbf{a}}_1 &= \mathbf{a}_1 \\ \tilde{\mathbf{a}}_2 &= \mathbf{a}_2 - (\mathbf{a}_2 \cdot \mathbf{a}_1) \frac{\mathbf{a}_1}{(\mathbf{a}_1 \cdot \mathbf{a}_1)} \\ \tilde{\mathbf{a}}_3 &= \mathbf{a}_3 - (\mathbf{a}_3 \cdot \mathbf{a}_1) \frac{\mathbf{a}_1}{(\mathbf{a}_1 \cdot \mathbf{a}_1)} - (\mathbf{a}_3 \cdot \mathbf{a}_2) \frac{\mathbf{a}_2}{(\mathbf{a}_2 \cdot \mathbf{a}_2)} \end{split}$$

Overdetermined Linear Systems

Modified Gram-Schmidt Orthogonalization

• More efficient formula (standard Gram-Schmidt):

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

with cost $\approx 2mn^2$ FLOPS but is **not numerically stable** against roundoff errors (loss of orthogonality).

• A numerically stable alternative is the modified Gram-Schmidt, in which each orthogonalization is carried against each of the already-computed basis vectors.

As we saw in previous lecture, a small rearrangement of mathematically-equivalent approaches can produce a much more robust numerical method.

- A matrix where a substantial fraction of the entries are zero is called a **sparse matrix**. The difference with **dense matrices** is that only the nonzero entries are stored in computer memory.
- Exploiting sparsity is important for **large matrices** (what is large depends on the computer).
- The structure of a sparse matrix refers to the set of indices *i*, *j* such that $a_{ij} > 0$, and is visualized in MATLAB using *spy*.
- The structure of sparse matrices comes from the nature of the problem, e.g., in an inter-city road transportation problem it corresponds to the pairs of cities connected by a road.
- In fact, just counting the number of nonzero elements is not enough: the **sparsity structure** is the most important property that determines the best method.

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

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-zeros 15 UNDIRECTED GRAPH FOR SYMMETRIC MATRICES

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Sparse matrices in MATLAB

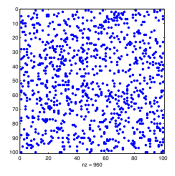
>> A = sparse(
$$[1 \ 2 \ 2 \ 4 \ 4]$$
, $[3 \ 1 \ 4 \ 2 \ 3]$, 1:5)
A =
(2,1) 2
(4,2) 4
(1,3) 1
(4,3) 5
(2,4) 3
>> nnz(A) % Number of non-zeros
ans = 5
>> whos A
A 4×4 120 double sparse
>> A = sparse([],[],[],4,4,5); % Pre-allocate memory
>> A(2,1)=2; A(4,2)=4; A(1,3)=1; A(4,3)=5; A(2,4)=3;

Sparse matrix factorization

an

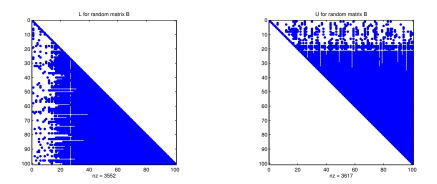
Random matrix \mathbf{B} and structured matrix \mathbf{X}

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



Fill-In

- There are general techniques for dealing with sparse matrices such as **sparse LU factorization**. How well they work depends on the structure of the matrix.
- When factorizing sparse matrices, the factors, e.g., L and U, can be much less sparse than A: fill-in.
- Pivoting (**reordering** of variables and equations) has a dual, sometimes conflicting goal:
 - **1** Reduce fill-in, i.e., **improve memory use**.
 - Reduce roundoff error, i.e., improve stability. Typically some threshold pivoting is used only when needed.
- For many sparse matrices there is a large fill-in and **iterative methods** are required.

Why iterative methods?

- Direct solvers are great for dense matrices and are implemented very well on modern machines.
- Fill-in is a major problem for certain sparse matrices and leads to extreme memory requirements.
- 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**.

Stationary Linear Iterative Methods

- 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** (many different choices/algorithms exist).

• 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 example, this fixed-point iteration is an iterative method:

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

Convergence of simple iterative methods

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

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

• 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,

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

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

• More precisely, $\lim_{k\to\infty} \mathbf{e}^{(k)} = \mathbf{0}$ for any $\mathbf{e}^{(0)}$ iff $\mathbf{B}^k \to \mathbf{0}$.

Spectral Radius

• Theorem: The simple iterative method converges iff the **spectral radius** of the iteration matrix is less than unity:

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

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

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

 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,

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

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 shown to be good if convergence is rapid.

• 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** $\mathbf{x}^{(0)}$.

Conclusions/Summary

• The conditioning of a linear system $\mathbf{A}\mathbf{x} = \mathbf{b}$ is determined by the condition number

$$\kappa(\mathbf{A}) = \|\mathbf{A}\| \|\mathbf{A}^{-1}\| \ge 1$$

- Gauss elimination can be used to solve general square linear systems and also produces a factorization **A** = **LU**.
- Partial pivoting is often necessary to ensure numerical stability during GEM and leads to PA = LU or $A = \tilde{L}U$.
- MATLAB has excellent linear solvers based on well-known public domain libraries like LAPACK. Use them!

Conclusions/Summary

- For symmetric positive definite matrices the Cholesky factorization
 A = LL^T is preferred and does not require pivoting.
- The *QR* factorization is a numerically-stable method for solving **full-rank non-square systems**.
- **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.