Lecture 13

Symmetric Factorizations & Iterative Methods

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Goals for today...

- Review LDL^T factorization
- Review Cholesky factorization.
- Discussion of Matrix solve costs.
- Iterative methods

Reminder: Goal of Factorization?

- Different solves have different costs:
 - $Ax = b \cos \theta (n^3)$ via Gaussian elimination.
 - Ly = b or Uy = b costs $O(n^2)$ via forward/backward substitution.
 - \rightarrow A = LU costs $\Im(n^3)$ to compute L and U (basically just Gaussian Elimination), but each successive solve is just 2 traingular solves.
- What if we know that $A = A^T$, can we speed up computation of L and U?

Use SYMMETRY! YRTEMMYS esU

Suppose

$$A = A^T$$
.

$$\Rightarrow LU = A = LDL^T$$
.

LDL[™] Factorization

If A is symmetric, we can factor A as

$$A = LDL^T$$

where L is the same lower triangular matrix made during LU factorization (don't need to do any work to create L^T .

How do we make *D*?

If you multiply out $A = LDL^T$ diagonal by diagonal and take advantage of all the zeros, you get

$$d_{11}=a_{11},$$
 $d_{22}=a_{22}-d_{11}\ell_{21}^2,$ $d_{33}=a_{33}-d_{11}\ell_{31}^2-d_2\ell_{32}^2,$.

Listing 1: LDL^T Factorization

```
given A
       output L, D
       for j = 1 \dots n
          \ell_{ii} = 1
           d_i = a_{ii} - \sum_{\nu=1}^{j-1} d_{\nu} \ell_{i\nu}^2. % COULD BE NEGATIVE
7
           for i = j + 1 ... n
                   \ell_{ii} = 0
                    \ell_{ij} = \left(a_{ij} - \sum_{\nu=1}^{j-1} \ell_{i\nu} d_{\nu} \ell_{j\nu}\right) / d_{j}
            end
       end
```

Special form of the LU factorization (for symmetric A).

$\tilde{L}\tilde{L}^T$: Cholesky Factorization

- A must be symmetric and positive definite (SPD)
- A is Positive Definite (PD) if for all $x \neq 0$ the following holds

$$x^T A x > 0$$

- Positive definite gives us an all positive D in $A = LDL^T$
- Define $D^{1/2}$ via $\sqrt{d_{ii}}$.
- $A = LDL^T = LD^{1/2}D^{1/2}L^T$.
- Define $\tilde{L} = LD^{1/2}$.
- $A = \tilde{L}\tilde{L}^T$, i.e. $\tilde{L} = \tilde{U}^T$ (note, L had ones on the diagonal, \tilde{L} does not).
 - Half as many flops as LU!
 - Only calculate L not U

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Cholesky Factorization

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Listing 2: Cholesky

Why SPD?

In general, SPD gives us

- non singular
 - If $x^T A x > 0$, for all nonzero x
 - Then $\lambda > 0$ for all eigenvalues λ
- No pivoting
 - From algorithm, can derive that $|I_{ki}| \leqslant \sqrt{a_{kk}}$
 - Elements of \tilde{L} do not grow with respect to A
 - For short proof see book
- Cholesky faster than LU
 - No pivoting
 - Only calculate L
 , not U

Why SPD?

A matrix is Positive Definite (PD) if for all $x \neq 0$ the following holds

$$x^T A x > 0$$

- For SPD matrices, use the Cholesky factorization, $A = \tilde{L}\tilde{L}^T$
- Cholesky Factorization
 - Requires no pivoting
 - Requires one half as many flops as LU factorization, that is only calculate L
 not L and U.
 - Cholesky will be more than twice as fast as LU because no pivoting means no data movement
- Use MATLAB's built-in cho1 function for routine work

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Motivation Revisted

Multiple right hand sides

- Solve Ax = b for k different b vectors
- Using LU factorization, the cost is $O(n^3) + O(kn^2)$
- Using Gaussian Elimination, the cost is $O(kn^3)$

If A is symmetric

- Save 50% of the flops with LDL^T factorization
- Save 50% of the flops and many memory operations with Cholesky $(\tilde{L}\tilde{L}^T)$ factorization

See time_LU_vs_Cholesky.m

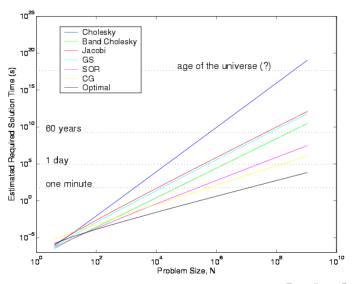
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Summary: Complexity of Linear Solves

- Ax = b
- diagonal system: O(n)
- tridiagonal system: O(n) (bigger constant)
- m-band system: $\mathcal{O}(m^2n)$
- upper or lower triangular system: $O(n^2)$
- full system with GE: O(n³)
- scaled partial pivoting adds O(n²)
- full system with LU: O(n³)
- LU back solve: O(n²)
- m different right-hand sides (when using LU): $O(n^3 + mn^2)$

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Summary: Complexity



Approximate solutions

So far, we are seeking "exact" (up to floating point error) solutions x^* to

$$Ax = b$$
.

What if we only need an approximation \hat{x} to x^* ?

We would like some \hat{x} so that $\|\hat{x} - x^*\| \le \epsilon$, where ϵ is some tolerance.

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The Residual

We can't actually evaluate

$$e = x^* - \hat{x}$$

But...

For
$$x = x^*$$

$$b - Ax \equiv 0$$

For
$$x = \hat{x}$$

$$b - Ax \not\equiv 0$$

We call $\hat{r} = b - A\hat{x}$ the *residual* (Recall from Lecture 7). It is a different way to measure the "error". In fact

$$\hat{r} = b - A\hat{x}$$

$$= Ax^* - A\hat{x}$$

$$= A\hat{e}$$

How big is the residual?

For a given approximation, \hat{x} to x, how "big" is the residual $\hat{r} = b - A\hat{x}$?

- ||r|| gives a magnitude
- $||r||_1 = \sum_{j=1}^n |r_i|$
- $||r||_2 = \left(\sum_{j=1}^n r_i^2\right)^{1/2}$
- $||r||_{\infty} = \max_{1 \leqslant j \leqslant n} |r_i|$

Notes



Approximating x...

Suppose we made a wild guess to the solution x of Ax = b:

$$x^{(0)} \approx x$$

How do I improve $x^{(0)}$?

Ideally:

$$x^{(1)} = x^{(0)} + e^{(0)}$$

but to obtain $e^{(0)}$, we must know x^* . Not a viable method.

Ideally (another way):

$$x^{(1)} = x^{(0)} + e^{(0)}$$

$$= x^{(0)} + (x^* - x^{(0)})$$

$$= x^{(0)} + (A^{-1}b - x^{(0)})$$

$$= x^{(0)} + A^{-1}(b - Ax^{(0)})$$

$$= x^{(0)} + A^{-1}r^{(0)}$$

An iteration

Again, the method

$$x^{(1)} = x^{(0)} + A^{-1}r^{(0)}$$

is nonsense since A^{-1} is needed.

What if we approximate A^{-1} ? Suppose $Q^{-1} \approx A^{-1}$ and is cheap to evaluate, then

$$x^{(1)} = x^{(0)} + Q^{-1}r^{(0)}$$

is a good step.

continuing...

$$x^{(k)} = x^{(k-1)} + Q^{-1}r^{(k-1)}$$
 (1)

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What kind of Q^{-1} do I need?

Rewrite:

$$x^{(k)} = x^{(k-1)} + Q^{-1}(b - Ax^{(k-1)})$$

This becomes

$$Qx^{(k)} = Qx^{(k-1)} + (b - Ax^{(k-1)})$$

= $(Q - A)x^{(k-1)} + b$

Or

$$x^{(k)} = Q^{-1}(Q - A)x^{(k-1)} + Q^{-1}b$$
 (2)

Often (1) is more practical for programming, while (2) is useful to do math. There are exceptions however.

Choices, choices....

We want Q^{-1} to be simple/cheap to apply. Ideally it will also be a "good" approximate of A^{-1} .

Let $A = D - C_L - C_U$ where D is diagonal, C_L and C_U are strictly lower and upper triangular.

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Choices, choices....

We want Q^{-1} to be simple/cheap to apply. Ideally it will also be a "good" approximate of A^{-1} .

Let $A = D - C_L - C_U$ where D is diagonal, C_L and C_U are strictly lower and upper triangular.

What if Q=D? Super cheap to invert, maybe a bad approximation... What if $Q=D-C_L$? Still kind of cheap, probably a better approximation...

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Two Popular Choices

Example

Jacobi iteration approximates A with Q = diag(A).

```
1 X = X^{(0)}

2 3 Q = D

3 for k = 1 to for k = 1 to for k = 1

6 for k = 1 to for k = 1

7 for k = 1 to for k = 1

8 for k = 1 to for k = 1

9 for k = 1 to for k = 1

10 for k = 1 to for k = 1

10 for k = 1 to for k = 1

11 for k = 1 to for k = 1

12 for k = 1 to for k = 1

13 for k = 1 to for k = 1

14 for k = 1 to for k = 1

15 for k = 1 to for k = 1

16 for k = 1 to for k = 1

17 for k = 1

18 for k = 1 to for k = 1

19 for k = 1

10 for k = 1

10
```

Two Popular Choices

Let $A = D - C_U - C_L$, where C_U and C_L are the negative of the strictly upper and lower triangular, respectively, of A.

Example

Gauss-Seidel iteration approximates A with Q = lowertri(A).

```
1 X = X^{(0)}

2 Q = D - C_L

3 Q = D - C_L

5 for k = 1 to k_{max}

6 r = b - Ax

7 if ||r|| \le tol, stop

8 X = X + Q^{-1}r

10 end
```

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Why D and $D - C_L$?

Look again at the iteration

$$x^{(k)} = x^{(k-1)} + Q^{-1}r^{(k-1)}$$

Looking at the error:

$$x^* - x^{(k)} = x^* - x^{(k-1)} - Q^{-1}r^{(k-1)}$$

Gives

$$e^{(k)} = e^{(k-1)} - Q^{-1}Ae^{(k-1)}$$

or

$$e^{(k)} = (I - Q^{-1}A)e^{(k-1)}$$

or

$$e^{(k)} = (I - Q^{-1}A)^k e^{(0)}$$

Why D and $D - C_L$?

We want

$$e^{(k)} = (I - Q^{-1}A)^k e^{(0)}$$

to converge.

Our iteration converges when $||I - Q^{-1}A|| < 1$, because

$$\|e^{(k)}\| = \|(I - Q^{-1}A)^k e^{(0)}\|$$

 $\leq \|I - Q^{-1}A\|^k \|e^{(0)}\|$

goes to 0, as $k \to \infty$.

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Notes



Matrix Norms (Recall Lecture 7)

What is $||I - Q^{-1}A||$?

•
$$||A||_1 = \max_{1 \leq j \leq n} \sum_{i=1}^n |a_{ij}|$$

•
$$\|A\|_{\infty} = \max_{1 \leqslant i \leqslant n} \sum_{j=1}^{n} |a_{ij}|$$

Other norms (you may be familiar with):

- $\|A\|_2 = \sqrt{\rho(A^TA)}$
- $\rho(A) = \max_{1 \leqslant j \leqslant n} |\lambda_i|$
- $||A||_2 = \rho(A)$ for symmetric A

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Again, why do Jacobi and Gauss-Seidel work?

Jacobi, Gauss-Seidel (sufficient) Convergence Theorem

If A is diagonally dominant, then the Jacobi and Gauss-Seidel methods converge for any initial guess $x^{(0)}$.

Definition: Diagonal Dominance

A matrix is diagonally dominant if

$$|a_{ii}| > \sum_{j=1, j\neq i}^{n} |a_{ij}|$$

for all i.

Notes



Smart Jacobi Algorithm

The Jacobi algorithm uses the matrix representation:

$$x^{(k)} = D^{-1}(C_L + C_U)x^{(k-1)} + D^{-1}b$$

Or componentwise,

$$x_i^{(k)} = -\sum_{j=1, j \neq i}^n \left(\frac{a_{ij}}{a_{ii}}\right) x_j^{(k-1)} + \frac{b_i}{a_{ii}}$$

So each sweep (from k-1 to k) uses O(n) operations per vector element (potentially $O(n^2)$ total).

If, for each row i, $a_{ij} \neq 0$ for at most m values, each sweep uses O(mn) operations.

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Smart Gauss-Seidel Algorithm

The Gauss-Seidel algorithm uses the matrix representation:

$$x^{(k)} = (D - C_L)^{-1}C_Ux^{(k-1)} + (D - C_L)^{-1}b$$

Component-wise:

$$x_i^{(k)} = -\sum_{j=1}^{i-1} \left(\frac{a_{ij}}{a_{ii}}\right) x_j^{(k)} - \sum_{j=i+1}^{n} \left(\frac{a_{ij}}{a_{ii}}\right) x_j^{(k-1)} + \frac{b_i}{a_{ii}}$$

So again each sweep (from k-1 to k) uses O(n) operations per vector element.

If, for each row i, $a_{ij} \neq 0$ for at most m values, each sweep uses O(mn) operations.

The difference is that in the Jacobi method, updates are saved (and not used) in a new vector. With Gauss-Seidel, an update to an element $x_i^{(k)}$ is used immediately.

Example:

Lets do Jacobi and Gauss-Seidel:

Example

Consider

$$A = \begin{bmatrix} 2 & -1 & 0 \\ -1 & 3 & -1 \\ 0 & -1 & 2 \end{bmatrix}, \qquad b = \begin{bmatrix} 1 \\ 8 \\ -5 \end{bmatrix}, \qquad x^{(0)} = \begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix}$$

See iterative_methods.m

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