1 The slippery inverse

The concept of the inverse of a matrix is generally more useful in theory than in numerical practice. We work with the inverse implicitly all the time through solving linear systems via LU; but we rarely form it explicitly, unless the inverse has some special structure we want to study or to use.

If we did want to form $A^{-1}$ explicitly, the usual approach is to compute $PA = LU$, then use that factorization to solve the systems $Ax_k = e_k$, where $e_k$ is the $k$th column of the identity matrix and $x_k$ is thus the $k$th column of the identity matrix. As discussed last time, forming the $LU$ factorization takes $n^3/3$ multiply-adds ($2n^3/3$ flops), and a pair of triangular solves takes $n^2$ multiply-add operations. Therefore, computing the inverse explicitly via an LU factorization takes about $n^3$ multiply-add operations, or roughly three times as much arithmetic as the original LU factorization. Furthermore, multiplying by an explicit inverse is almost exactly the same amount of arithmetic work as a pair of triangular solves. So computing and using an explicit inverse is, on balance, more expensive than simply solving linear systems using the LU factorization.

To make matters worse, multiplying by the explicit inverse of a matrix is not a backward stable algorithm. Even if we could compute $A^{-1}$ essentially exactly, only committing rounding errors when storing the entries and when performing matrix-vector multiplication, we would find $\hat{f}(A^{-1}b) = (A^{-1} + F)b$, where $|F| \leq n\epsilon_{\text{mach}}|A^{-1}|$. But this corresponds to a backward error of roughly $-AF\hat{A}$, which is potentially much larger than $\|A\|$.

In summary: you should get used to the idea that any time you see an inverse in the description of a numerical method, it is probably shorthand for “solve a linear system here.” Except in special circumstances, forming and multiplying by an explicit inverse is both slower and less numerically stable than solving a linear system by Gaussian elimination.

2 Iterative refinement revisited

At the end of last lecture, we discussed iterative refinement:

$$x_{k+1} = x_k + \hat{A}^{-1}(b - Ax_k).$$
The fixed point for this iteration is $x = A^{-1}b$, and we can write a simple recurrence for the error $e_{k+1} = x_k - x$:

$$e_{k+1} = \hat{A}^{-1}E e_k$$

where $E = \hat{A} - A$. Therefore, if $\|\hat{A}^{-1}E\| < 1$, then iterative refinement converges — in exact arithmetic.

In floating point arithmetic, we actually compute something like

$$x_{k+1} = x_k + \hat{A}_k^{-1}(b - Ax_k + \delta_k) + \mu_k,$$

where $\hat{A}_k = A + E_k$ accounts for the backward error $E_k$ in the approximate solve, $\delta_k$ is an error associated with computing the residual, and $\mu_k$ is an error associated with the update. This gives us the error recurrence

$$e_{k+1} = \hat{A}_k^{-1}E e_k + \hat{A}_k^{-1}\delta_k + \mu_k$$

If $\|\delta_k\| < \alpha$, $\|\mu_k\| < \beta$, and $\|A^{-1}E_k\| < \gamma < 1$ for all $k$, then we can show that

$$\|x_k - x\| \leq \gamma^k\|x_0 - x\| + \frac{\alpha\|A^{-1}\| + \beta}{1 - \gamma}.$$

If we evaluate the residual in the obvious way, we typically have

$$\alpha \leq c_1\epsilon_{\text{mach}}\|A\|\|x\|,$$
$$\beta \leq c_2\epsilon_{\text{mach}}\|x\|,$$

for some modest $c_1$ and $c_2$; and for large enough $k$, we end up with

$$\frac{\|x_k - x\|}{\|x\|} \leq C_1\epsilon_{\text{mach}}\kappa(A) + C_2\epsilon_{\text{mach}}.$$

That is, iterative refinement leads to a relative error not too much greater than we would expect due to a small relative perturbation to $A$; and we can show that in this case the result is backward stable. And if we use mixed precision to evaluate the residual accurately enough relative to $\kappa(A)$ (i.e. $\alpha\kappa(A) \lesssim \beta$) we can actually achieve a small forward error.
3 Condition estimation

Suppose now that we want to compute \( \kappa_1(A) \) (or \( \kappa_\infty(A) = \kappa_1(A^T) \)). The most obvious approach would be to compute \( A^{-1} \), and then to evaluate \( \|A^{-1}\|_1 \) and \( \|A\|_1 \). But the computation of \( A^{-1} \) involves solving \( n \) linear systems for a total cost of \( O(n^3) \) — the same order of magnitude as the initial factorization. Error estimates that cost too much typically don’t get used, so we want a different approach to estimating \( \kappa_1(A) \), one that does not cost so much. The only piece that is expensive is the evaluation of \( \|A^{-1}\|_1 \), so we will focus on this.

Note that \( \|A^{-1}x\|_1 \) is a convex function of \( x \), and that \( \|x\|_1 \leq 1 \) is a convex set. So finding

\[
\|A^{-1}\|_1 = \max_{\|x\|_1 \leq 1} \|A^{-1}x\|_1
\]

is a convex optimization problem. Also, note that \( \| \cdot \|_1 \) is differentiable almost everywhere: if all the components of \( y \) are nonzero, then

\[
\xi^T y = \|y\|_1, \quad \text{for } \xi = \text{sign}(y);
\]

and if \( \delta y \) is small enough so that all the components of \( y + \delta y \) have the same sign as the corresponding components of \( y \), then

\[
\xi^T (y + \delta y) = \|y + \delta y\|_1
\]

More generally, we have

\[
\xi^T u \leq \|\xi\|_\infty \|u\|_1 = \|u\|_1,
\]

i.e. even when \( \delta y \) is big enough so that the linear approximation to \( \|y + \delta y\|_1 \) no longer holds, we at least have a lower bound.

Since \( y = A^{-1}x \), we actually have that

\[
|\xi^T A^{-1}(x + \delta x)| \leq \|A^{-1}(x + \delta x)\|,
\]

with equality when \( \delta x \) is sufficiently small (assuming \( y \) has no zero components). This suggests that we move from an initial guess \( x \) to a new guess \( x_{\text{new}} \) by maximizing

\[
|\xi^T A^{-1} x_{\text{new}}| \quad \text{over } \|x_{\text{new}}\| \leq 1.
\]

This actually yields \( x_{\text{new}} = e_j \), where \( j \) is chosen so that the \( j \)th component of \( z^T = \xi^T A^{-1} \) has the greatest magnitude.

Putting everything together, we have the following algorithm
% Hager's algorithm to estimate \( \|A^{-1}\|_1 \)
% We assume solveA and solveAT are \( O(n^2) \) solution algorithms
% for linear systems involving \( A \) or \( A' \) (e.g. via LU)

\( x = \text{ones}(n,1)/n; \)  \% Initial guess
while true
    \( y = \text{solveA}(x); \)  \% Evaluate \( y = A^{-1} x \)
    \( x_i = \text{sign}(y); \)  \% and \( z = A^{-T} \text{sign}(y) \), the
    \( z = \text{solveAT}(x_i); \)  \% (sub)gradient of \( x \rightarrow \|A^{-1} x\|_1. \)
    \% Find the largest magnitude component of \( z \)
    \[ z_{norn}, j \] = \text{max}(\text{abs}(z));
    \% znorn = \( |z_j| \) is our lower bound on \( |A^{-1} e_j| \).
    \% If this lower bound is no better than where we are now, quit
    if znorn <= \( \text{norm}(y,1) \)
        \( \text{invA}_nortest = \text{norm}(y,1); \)
        break;
    end
    \% Update \( x \) to \( e_j \) and repeat
    \( x = \text{zeros}(n,1); x(j) = 1; \)
end

This method is not infallible, but it usually gives estimates that are the right order of magnitude. There are various alternatives, refinements, and extensions to Hager's method, but they generally have the same flavor of probing \( A^{-1} \) through repeated solves with \( A \) and \( A^T \).

4 Scaling

Suppose we wish to solve \( Ax = b \) where \( A \) is ill-conditioned. Sometimes, the ill-conditioning is artificial because we made a poor choice of units, and it appears to be better conditioned if we write

\[ D_1AD_2y = D_1b, \]

where \( D_1 \) and \( D_2 \) are diagonal scaling matrices. If the original problem was poorly scaled, we will likely find \( \kappa(D_1AD_2) \ll \kappa(A) \), which may be great for Gaussian elimination. But by scaling the matrix, we are really changing the
norms that we use to measure errors — and that may not be the right thing to do.

For physical problems, a good rule of thumb is to non-dimensionalize before computing. The non-dimensionalization will usually reveal a good scaling that (one hopes) simultaneously is appropriate for measuring errors and does not lead to artificially inflated condition numbers.

5 Symmetric matrices

5.1 Quadratic forms

A matrix $A$ is symmetric if $A = A^T$. For each symmetric matrix $A$, there is an associated quadratic form $x^T A x$. Even if you forgot them from our lightning review of linear algebra, you are likely familiar with quadratic forms from a multivariate calculus class, where they appear in the context of the second derivative test. One expands

$$F(x + u) = F(x) + F'(x)u + \frac{1}{2} u^T H(x) u + O(\|u\|^3),$$

and notes that at a stationary point where $F'(x) = 0$, the dominant term is the quadratic term. When $H$ is positive definite or negative definite, $x$ is a strong local minimum or maximum, respectively. When $H$ is indefinite, with both negative and positive eigenvalues, $x$ is a saddle point. When $H$ is semi-definite, one has to take more terms in the Taylor series to determine whether the point is a local extremum.

If $B$ is a nonsingular matrix, then we can write $x = By$ and $x^T A x = y^T (B^T A B) y$. So an “uphill” direction for $A$ corresponds to an “uphill” direction for $B^T A B$; and similarly with downhill directions. More generally, $A$ and $B^T A B$ have the same inertia, where the inertia of a symmetric $A$ is the triple

(# pos eigenvalues, # zero eigenvalues, # neg eigenvalues).

Now suppose that $A = LU$, where $L$ is unit lower triangular and $U$ is upper triangular. If we let $D$ be the diagonal part of $U$, we can write $A = LDM^T$, where $L$ and $M$ are both unit lower triangular matrices. Noting that $A^T = (LDM^T)^T = MDL^T = M(LD)^T$ and that the $LU$ factorization
of a matrix is unique, we find $M = L$ and $LD = DM^T = U$. Note that $D$ has the same inertia as $A$.

The advantage of the $LDL^T$ factorization over the $LU$ factorization is that we need only compute and store one triangular factor, and so $LDL^T$ factorization costs about half the flops and storage of $LU$ factorization. We have the same stability issues for $LDL^T$ factorization that we have for ordinary $LU$ factorization, so in general we might compute

$$PAP^T = LDL^T,$$

where the details of various pivoting schemes are described in the book.

### 5.2 Positive definite matrices

A symmetric matrix is positive definite if $x^T Ax > 0$ for all nonzero $x$. If $A$ is symmetric and positive definite, then $A = LDL^T$ where $D$ has all positive elements (because $A$ and $D$ have the same inertia). Thus, we can write $A = (LD^{1/2})(LD^{1/2})^T = \hat{L}\hat{L}^T$. The matrix $\hat{L}$ is a Cholesky factor of $A$.

There are several useful properties of SPD matrices that we will use from time to time:

1. The inverse of an SPD matrix is SPD.

   **Proof:** If $x^T Ax > 0$ for all $x \neq 0$, then we cannot have $Ax = 0$ for nonzero $x$. So $A$ is necessarily nonsingular. Moreover,

   $$x^T A^{-1} x = (A^{-1} x)^T A (A^{-1} x)$$

   must be positive for nonzero $x$ by positive-definiteness of $A$. Therefore, $A^{-1}$ is SPD.

2. Any minor of an SPD matrix is SPD.

   **Proof:** Without loss of generality, let $M = A_{11}$. Then for any appropriately sized $x$,

   $$x^T M x = \begin{bmatrix} x^T \\ 0 \end{bmatrix} A \begin{bmatrix} x \\ 0 \end{bmatrix} > 0$$

   for $x \neq 0$. Therefore, $M$ is positive definite.
3. Any Schur complement of an SPD matrix is SPD

**Proof:** A Schur complement in $A$ is the inverse of a minor of an inverse of $A$. By the two arguments above, this implies that any Schur complement of an SPD matrix is SPD.

4. If $M$ is a minor of $A$, $\kappa_2(M) \leq \kappa_2(A)$.

**Proof:** The largest and smallest singular values of an SPD matrix are the same as the largest and smallest eigenvalues; they can be written as

$$\sigma_1(A) = \max_{\|x\|_2=1} x^T Ax, \quad \sigma_{\text{min}}(A) = \min_{\|x\|_2=1} x^T Ax.$$ 

Without loss of generality, let $M = A_{11}$. Then

$$\sigma_1(M) = \max_{\|x\|_2=1} x^T M x = \max_{\|x\|_2=1} \left[ x \atop 0 \right]^T A \left[ x \atop 0 \right] \leq \max_{\|z\|_2=1} z^T Az = \sigma_1(A)$$

and similarly $\sigma_{\text{min}}(M) \geq \sigma_{\text{min}}(A)$. The condition numbers are therefore

$$\kappa_2(M) = \frac{\sigma_1(M)}{\sigma_{\text{min}}(M)} \leq \frac{\sigma_1(A)}{\sigma_{\text{min}}(A)} = \kappa_2(A).$$

5. If $S$ is a Schur complement in $A$, $\kappa_2(S) \leq \kappa_2(A)$.

**Proof:** This is left as an exercise.

6 **Cholesky**

The algorithm to compute the Cholesky factor of an SPD matrix is close to the Gaussian elimination algorithm. In the first step, we would write

$$\begin{bmatrix} a_{11} & a_{21}^T \\ a_{21} & A_{22} \end{bmatrix} = \begin{bmatrix} l_{11} & 0 \\ l_{21} & L_{22} \end{bmatrix} \begin{bmatrix} l_{11} & l_{21}^T \\ 0 & L_{22}^T \end{bmatrix},$$

or

$$a_{11} = l_{11}^2$$
$$a_{21} = l_{21} l_{11}$$
$$A_{22} = L_{22} L_{22}^T + l_{21} l_{21}^T.$$
The first two equations allow us to compute the first column of $L$; the last equation tells us that the rest of $L$ is the Cholesky factor of a Schur complement, $L_{22}L_{22}^T = A_{22} - l_{21}l_{21}^T$. Continuing in this fashion, we have the algorithm

```
for j = 1:n

    % Compute the jth column of L
    A(j,j) = sqrt(A(j,j));
    A(j+1:end,j) = A(j+1:end,j)/A(j,j);

    % Update trailing submatrix
    A(j+1:end,j+1:end) = A(j+1:end,j+1:end) - A(j+1:end,j)*A(j+1:end,j)';

end

% Lower triangle was overwritten by L
L = tril(A);
```

Like the nearly-identical Gaussian elimination algorithm, we can rewrite the Cholesky algorithm in block form for better cache use. Unlike Gaussian elimination, we do just fine using Cholesky without pivoting\(^1\).

\(^1\)Pivoting can still be useful for near-singular matrices, but unpivoted Cholesky is backward stable