1 Householder transformations

The Gram-Schmidt orthogonalization procedure is not generally recommended for numerical use. Suppose we write \( A = [a_1 \ldots a_m] \) and \( Q = [q_1 \ldots q_m] \). The essential problem is that if \( r_{jj} \ll \|a_j\|_2 \), then cancellation can destroy the accuracy of the computed \( q_j \); and in particular, the computed \( q_j \) may not be particularly orthogonal to the previous \( q_j \). Actually, loss of orthogonality can build up even if the diagonal elements of \( R \) are not exceptionally small. This is Not Good, and while we have some tricks to mitigate the problem, we need a different approach if we want the problem to go away.

Recall that one way of expressing the Gaussian elimination algorithm is in terms of Gauss transformations that serve to introduce zeros into the lower triangle of a matrix. Householder transformations are orthogonal transformations (reflections) that can be used to similar effect. Reflection across the plane orthogonal to a unit normal vector \( v \) can be expressed in matrix form as

\[
H = I - 2vv^T.
\]

Now suppose we are given a vector \( x \) and we want to find a reflection that transforms \( x \) into a direction parallel to some unit vector \( y \). The right reflection is through a hyperplane that bisects the angle between \( x \) and \( y \) (see Figure 1), which we can construct by taking the hyperplane normal to \( x - \|x\|y \). That is, letting \( u = x - \|x\|y \) and \( v = u/\|u\| \), we have

\[
(I - 2vv^T)x = x - 2\left( x + \|x\|y \right) \left( x^T x + \|x\| x^T y \right) \frac{1}{\|x\|^2 + 2x^T y \|x\| + \|x\|^2 \|y\|^2}
\]

\[
= x - (x - \|x\|y)
\]

\[
= \|x\|y.
\]

If we use \( y = \pm e_1 \), we can get a reflection that zeros out all but the first element of the vector \( x \). So with appropriate choices of reflections, we can take a matrix \( A \) and zero out all of the subdiagonal elements of the first column.

Now think about applying a sequence of Householder transformations to introduce subdiagonal zeros into \( A \), just as we used a sequence of Gauss
transformations to introduce subdiagonal zeros in Gaussian elimination. This leads us to the following algorithm to compute the QR decomposition:

```matlab
function [Q,R] = hqr1(A)
    % Compute the QR decomposition of an m-by-n matrix A using
    % Householder transformations.

    [m,n] = size(A);
    Q = eye(m);                 % Orthogonal transform so far
    R = A;                      % Transformed matrix so far

    for j = 1:n
        % -- Find H = I-tau*w*w' to put zeros below R(j,j)
        normx = norm(R(j:end,j));
        s = sign(R(j,j));
        u1 = R(j,j) - s*normx;
        w = R(j:end,j)/u1;
        w(1) = 1;
        tau = -s*u1/normx;

        % -- R := HR, Q := QH
        R(j:end,:) = R(j:end,:)-(tau*w)*(w'*R(j:end,:));
        Q(:,j:end) = Q(:,j:end)-(Q(:,j:end)*w)*(tau*w)';
    end
```

Note that there are two valid choices of $u_1$ at each step; we make the choice that avoids cancellation in the obvious version of the formula.

As with $LU$ factorization, we can re-use the storage of $A$ by recognizing that the number of nontrivial parameters in the vector $w$ at each step is the
same as the number of zeros produced by that transformation. This gives us
the following:

```matlab
function [A,tau] = hqr2(A)
% Compute the QR decomposition of an m-by-n matrix A using
% Householder transformations, re-using the storage of A
% for the Q and R factors.

[m,n] = size(A);
tau = zeros(n,1);

for j = 1:n
    % -- Find H = I-tau*w*w' to put zeros below A(j,j)
    normx = norm(A(j:end,j));
    s = -sign(A(j,j));
    u1 = A(j,j) - s*normx;
    w = A(j:end,j)/u1;
    w(1) = 1;
    A(j+1:end,j) = w(2:end);
    A(j,j) = s*normx;
    tau(j) = -s*u1/normx;

    % -- R := HR
    A(j:end,j+1:end) = A(j:end,j+1:end) -
        (tau(j)*w)*(w'*A(j:end,j+1:end));
end

If we ever need Q or QT explicitly, we can always form it from the com-
pressed representation. We can also multiply by Q and QT implicitly:

```
Givens rotations

Householder reflections are one of the standard orthogonal transformations used in numerical linear algebra. The other standard orthogonal transformation is a Givens rotation:

\[
G = \begin{bmatrix}
c & -s \\
s & c
\end{bmatrix}
\]

where \(c^2 + s^2 = 1\). Note that

\[
G = \begin{bmatrix}
c & -s \\
s & c
\end{bmatrix}
\begin{bmatrix}
x \\
y
\end{bmatrix}
= \begin{bmatrix}
cx - sy \\
rx + cy
\end{bmatrix}
\]

so if we choose

\[
s = \frac{-y}{\sqrt{x^2 + y^2}}, \quad c = \frac{x}{\sqrt{x^2 + y^2}}
\]

then the Givens rotation introduces a zero in the second column. More generally, we can transform a vector in \(\mathbb{R}^m\) into a vector parallel to \(e_1\) by a sequence of \(m - 1\) Givens rotations, where the first rotation moves the last element to zero, the second rotation moves the second-to-last element to zero, and so forth.

For some applications, introducing zeros one by one is very attractive. In some places, you may see this phrased as a contrast between algorithms based on Householder reflections and those based on Givens rotations, but this is not quite right. Small Householder reflections can be used to introduce one zero at a time, too. Still, in the general usage, Givens rotations seem to be the more popular choice for this sort of local introduction of zeros.

Stability of QR

It is not too difficult to show that applying a Givens rotations or Householder reflector to a matrix is backward-stable: if \(P\) is the desired transformation, the floating point result of \(PA\) is

\[
\tilde{P}A = (P + E)A, \quad \|E\| \leq O(\epsilon_{\text{mach}})\|A\|.
\]
Moreover, orthogonal matrices are perfectly conditioned! Taking a product of \( j \) matrices is also fine; the result has backward error bounded by \( jO(\epsilon_{\text{mach}})||A|| \). As a consequence, QR decomposition by Givens rotations or Householder transformations is ultimately backward stable.

The stability of orthogonal matrices in general makes them a marvelous building block for numerical linear algebra algorithms, and we will take advantage of this again when we discuss eigenvalue solvers.

\section{Sparse QR}

Just as was the case with LU, the QR decomposition admits a sparse variant. And, as with LU, sparsity of the matrix \( A \in \mathbb{R}^{m \times n} \) alone is not enough to guarantee sparsity of the factorization! Hence, as with solving linear systems, our recommendation for solving sparse least squares problems varies depending on the actual sparse structure.

Recall that the \( R \) matrix in QR factorization is also the Cholesky factor of the Gram matrix: \( G = A^T A = R^T R \). Hence, the sparsity of the \( R \) factor can be inferred from the sparsity of \( G \) using the ideas we talked about when discussing sparse Cholesky. If the rows of \( A \) correspond to experiments and columns correspond to factors, the nonzero structure of \( G \) is determined by which experiments share common factors: in general \( g_{ij} \neq 0 \) if any experiment involves both factors \( i \) and factor \( j \). So a very sparse \( A \) matrix may nonetheless yield a completely dense \( G \) matrix. Of course, if \( R \) is dense, that is not the end of the world! Factoring a dense \( n \times n \) matrix is pretty cheap for \( n \) in the hundreds or even up to a couple thousand, and solves with the resulting triangular factors are quite inexpensive.

If one forms \( Q \) at all, it is often better to work with \( Q \) as a product of (sparse) Householder reflectors rather than forming the elements of \( Q \). One may also choose to use a “\( Q \)-less QR decomposition” in which the matrix \( Q \) is not kept in any explicit form; to form \( Q^T b \) in this case, we would use the formulation \( Q^T b = R^{-T} A^T b \).

As with linear solves, least squares solves can be “cleaned up” using iterative refinement. This is a good idea in particular when using \( Q \)-less QR. If \( \tilde{A} \) is an approximate least squares solve (e.g. via the slightly-unstable
normal equations approach), iterative refinement looks like

\[ r^k = b - Ax^k \]
\[ x^{k+1} = x^k - \tilde{R}^{-1}(\tilde{R}^T(A^T r_k)) \]

This approach can be useful even when \( A \) is moderately large and dense; for example, \( \tilde{R} \) might be computed from a (scaled) QR decomposition of a carefully selected subset of the rows of \( A \).

## 5 Constrained case

Consider the weighted least squares problem

\[
\text{minimize } \sum_{i=1}^{m} w_i r_i^2
\]

where \( w_1 \) is much larger than the others. If we let \( w_1 \to \infty \) while the others are fixed, what happens? We essentially say that we care about enforcing the first equation above all others, and in the limit we are solving the constrained least squares problem

\[
\text{minimize } \sum_{i=2}^{m} w_i r_i^2 \text{ s.t. } r_1 = 0.
\]

Unfortunately, if we actually try to compute this way, we are dancing on dangerous ground; as \( w_1 \) goes to infinity, so does the condition number of the least squares problem. But this is only an issue with the weighted formulation; we can formulate the constrained problem in other ways that are perfectly well-behaved.

In the remainder of this section, we address two ways of handling the linearly constrained least squares problem

\[
\text{minimize } \|Ax - b\|^2 \text{ s.t. } C^T x = d,
\]

by either eliminating variables (the null-space method) or adding variables (the method of Lagrange multipliers).
5.1 Null space method

In the null space method, we write an explicit expression for the solutions to \( C^T x = d \) in the form \( x_p + W z \) where \( x_p \) is a particular solution to \( C^T x_p = d \) and \( W \) is a basis for the null space of \( C^T \). Perhaps the simplest particular solution is \( x_p = (C^T)^+d \), the solution with minimal norm; we can compute both this particular solution and an orthonormal null space basis quickly using a full QR decomposition of \( C \):

\[
C = \begin{bmatrix} Q_1 & Q_2 \end{bmatrix} \begin{bmatrix} R_1 \\ 0 \end{bmatrix}, \quad x_p = Q_1 R_1^{-T} d, \quad W = Q_2.
\]

Note that

\[
C^T x_p = (R_1^T Q_1^T) x_p = d,
\]

so this is indeed a particular solution. Having written an explicit parameterization for all solutions of the constraint equations, we can minimize the least squares objective with respect to the reduced set of variables

\[
\text{minimize } \| A(x_p + W z) - b \|^2 = \|(AW) z - (b - Ax_p)\|^2.
\]

This new least squares problem involves a smaller set of variables (which is good); but in general, even if \( A \) is sparse, \( AW \) will not be. So it is appropriate to have a few more methods in our arsenal.

5.2 Lagrange multipliers

An alternate method is the method of Lagrange multipliers. This is an algebraic technique for adding equations to enforce constraints.

One way to approach the Lagrange multiplier method is to look at the equations for a constrained minimum. In order not to have a downhill direction, we require that the directional derivatives be zero in any direction consistent with the constraint; that is, we require \( C x = d \) and

\[
\delta x^T A^T r = 0 \quad \text{when} \quad C^T \delta x = 0.
\]

The constraint says that admissible \( \delta x \) are orthogonal to the columns of \( C \); the objective tells us the admissible \( \delta x \) should be orthogonal to the residual. So we need that \( A^T r \) should lie in the column span of \( C \); that is,

\[
A^T r = -C\lambda
\]
for some $\lambda$, and $Cx = d$. Putting this together, we have the KKT equations

$$
\begin{bmatrix}
A^T A & C \\
C^T & 0
\end{bmatrix}
\begin{bmatrix}
x \\
\lambda
\end{bmatrix}
=
\begin{bmatrix}
A^T b \\
d
\end{bmatrix}.
$$

These bordered normal equations are not the end point for constrained least squares with Lagrange multipliers, any more than the normal equations are the end point for unconstrained least squares. Rather, we can use this as a starting point for clever manipulations involving our favorite factorizations (QR and SVD) that reduce the bordered system to a more computationally convenient form.