

The Francis Algorithm

Recall that we have a shifted QR iteration that converges quickly to a *reduced* Hessenberg matrix if the s_i are close to an eigenvalue of H (in fact, only one iteration does it if s_i is an eigenvalue of H (this is called an *ultimate shift*)). If H is reduced, we can decouple (or deflate) and continue with a strictly smaller problem than before. Here is the iteration:

$$\begin{aligned} Q_i R_i &= H_i - s_i I \\ H_{i+1} &= R_i Q_i + s_i I = Q_i^T H_i Q_i \end{aligned}$$

Nonsymmetric real matrices may have complex eigenvalues, which must occur in conjugate pairs: $u + iv$ and $u - iv$. If we apply a complex shift $s_i = u + iv$ to the iteration, then Q_i , R_i and H_{i+1} will be complex. This requires more storage, and more computation (one complex multiplication requires 4 real multiplications and 2 real additions).

Now if we immediately follow the $s_i = u + iv$ shift iteration with an iteration with shift $s_{i+1} = u - iv$, then everything becomes again real. We can understand this by noting that two iterations above gives the same H_{i+2} as the single iteration

$$\begin{aligned} QR &= (H_i - s_i I)(H_i - s_{i+1} I) \\ H_{i+2} &= Q^T H_i Q. \end{aligned}$$

Even more: $R = R_{i+1} R_i$ and $Q = Q_i Q_{i+1}$. So here is a way to apply two complex conjugate shifts in succession using only real arithmetic! While it appears, that we have bought this efficiency at the cost of forming $G = (H_i - s_i I)(H_i - s_{i+1} I)$, a wonderful uniqueness result comes to our aid:

The *Implicit Q Theorem* says that if $H_{i+2} = Q^T H_i Q$ is unreduced, then it is essentially uniquely determined by H_i and the first column of Q .

Thus we need only compute enough of G to determine the first column of Q , the remainder of Q is discovered through a Hessenberg reduction. Here are the details:

$$Q R e_1 = G e_1 \implies Q e_1 = \pm G e_1 / \|G\|_2.$$

So define a Householder reflector, P_0 , so that $P_0 G e_1 = \alpha e_1$ (the first step of the Householder QR factorization of G), and apply this *not* to G , but to H_i as a similarity transform: $B = P_0 H_i P_0$. B is no longer Hessenberg, it has a bulge at $b_{41} \neq 0$, so we need reflectors P_1, P_2, \dots, P_{n-2} to “chase the bulge” from b_{41} to b_{52} to ... to $b_{n,n-2}$, resp. to “re-Hessenbergize” H_i :

$$H_{i+2} = P_{n-2} \cdots P_1 P_0 H_i P_0 P_1 \cdots P_{n-2} = Q^T H_i Q.$$

All of this was developed in J. Francis’ 1961 paper, along with a scheme for choosing the shifts s_i and s_{i+1} as the eigenvalues of the lower right 2×2 submatrix of H_i . While fallable, this method – augmented with schemes for detecting subdiagonal elements “small enough” to allow decoupling and deflation – is the state of the art general purpose eigenvalue method.