

## FAST POLAR DECOMPOSITION OF AN ARBITRARY MATRIX\*

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**Abstract.** The polar decomposition of an  $m \times n$  matrix  $A$  of full rank, where  $m \geq n$ , can be computed using a quadratically convergent algorithm of Higham [*SIAM J. Sci. Statist. Comput.*, 7 (1986), pp. 1160-1174]. The algorithm is based on a Newton iteration involving a matrix inverse. It is shown how, with the use of a preliminary complete orthogonal decomposition, the algorithm can be extended to arbitrary  $A$ . The use of the algorithm to compute the positive semidefinite square root of a Hermitian positive semidefinite matrix is also described. A hybrid algorithm that adaptively switches from the matrix inversion based iteration to a matrix multiplication based iteration due to Kovarik, and to Björck and Bowie, is formulated. The decision when to switch is made using a condition estimator. This "matrix multiplication rich" algorithm is shown to be more efficient on machines for which matrix multiplication can be executed 1.5 times faster than matrix inversion.

**Key words.** polar decomposition, complete orthogonal decomposition, matrix square root, matrix multiplication, Schulz iteration, condition estimator

**AMS(MOS) subject classification.** 65F05

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**1. Introduction.** A polar decomposition of a matrix  $A \in \mathbb{C}^{m \times n}$  is a factorization  $A = UH$ , where  $H \in \mathbb{C}^{n \times n}$  is Hermitian positive semidefinite and  $U \in \mathbb{C}^{m \times n}$  is unitary; here we define unitary to mean that  $U$  has orthonormal rows or columns according as  $m \leq n$  or  $m \geq n$ . The decomposition always exists,  $H$  is the unique Hermitian positive semidefinite square root of  $A^*A$  (i.e.,  $H = (A^*A)^{1/2}$ ), and  $U$  is unique if and only if  $A$  has full rank (these properties are proved in § 2).

The polar decomposition is well known in the case  $m \geq n$  (see [8] and [11], for example). We have followed Horn and Johnson [14] in extending the definition to  $m \leq n$ . The consistency of the definition can be seen in the result that for any  $m$  and  $n$  the unitary polar factor  $U$  is a nearest unitary matrix to  $A$  in the Frobenius norm (this is a straightforward extension of a result from [6]). Because of the role it plays in solving this and other nearness problems, computation of the polar decomposition is required in several applications [13]. A recent application, which motivated the work here, is the computation of block reflectors (generalizations of Householder matrices) [19]. Here, the polar decomposition of an arbitrary matrix must be computed, and it is desirable to do this efficiently on vector and parallel computers.

The polar decomposition can be obtained directly from the singular value decomposition (SVD). Higham [11] describes an alternative approach based on a Newton iteration involving a matrix inverse. The iteration is defined for square, nonsingular matrices only, but in [11] it is pointed out how a preliminary QR decomposition enables the treatment of  $A \in \mathbb{C}^{m \times n}$  with  $m \geq n$  and  $\text{rank}(A) = n$ . It is also shown in [11] how the iteration can be used to compute the square root of a Hermitian positive definite matrix. According to the traditional model of computational cost based on operation counts, the iterative algorithm is generally of similar expense to

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the SVD approach, but is much more efficient when the matrix is nearly unitary. In an attempt to improve the performance of the iterative algorithm on machines that execute matrix multiplication at high efficiency, Schreiber and Parlett [19] propose the use of an inner Schulz iteration to compute most of the matrix inverses; they show that this leads to an increase in efficiency if matrix multiplication can be done at a rate 6.8 times faster than matrix inversion. Gander [7] develops a family of iteration methods for computing the polar decomposition of a rectangular matrix of full rank; his family includes a variant of the Newton iteration of [11].

The purpose of this work is twofold. First, we extend the algorithm of [11] so that it is applicable to arbitrary  $A$ . Our technique is to use an initial complete orthogonal decomposition so as to extract an appropriate square, nonsingular matrix. We might say that the complete orthogonal decomposition is to the polar decomposition what Chan's preliminary QR factorization is to the SVD! We also show how to use the algorithm of [11] to compute the square root of a (singular) Hermitian positive semidefinite matrix. Second, we introduce a modification of the Schulz inner iteration idea of [19] that reduces the cutoff ratio of multiplication speed to inversion speed from 6.8 to 2, or to 1.5 if advantage is taken of a symmetric matrix product.

**2. Iterative polar decomposition of an arbitrary matrix.** The basic algorithm of [11] is as follows. It converges quadratically for any square, nonsingular  $A$ . We use a MATLAB-like algorithmic notation, and denote by  $A^{-*}$  the conjugate transpose of  $A^{-1}$ .

ALGORITHM 2.1.  $[U, H] = \text{polar} \cdot \text{square}(A, \delta)$ .

% Input arguments: square, nonsingular  $A$ ; convergence tolerance  $\delta$ .

% Output arguments:  $U, H$ .

$X_0 = A$ ;  $k = -1$

repeat

$k = k + 1$

$\gamma_k = (\|X_k^{-1}\|_1 \|X_k^{-1}\|_\infty / (\|X_k\|_1 \|X_k\|_\infty))^{1/4}$

$X_{k+1} = \frac{1}{2}(\gamma_k X_k + X_k^{-*} / \gamma_k)$

until  $\|X_{k+1} - X_k\|_1 \leq \delta \|X_{k+1}\|_1$

$U = X_{k+1}$

$H = \frac{1}{2}(U^* A + A^* U)$

To adapt the algorithm to arbitrary  $A \in \mathbb{C}^{m \times n}$  we begin by computing a complete orthogonal decomposition (COD)

$$A = P \begin{bmatrix} R & 0 \\ 0 & 0 \end{bmatrix} Q^*,$$

where  $P \in \mathbb{C}^{m \times m}$  and  $Q \in \mathbb{C}^{n \times n}$  are unitary, and  $R \in \mathbb{C}^{r \times r}$  is nonsingular and upper triangular (we exclude the trivial case  $A = 0$ , for which  $R$  is empty). This decomposition may be computed using a QR factorization with column pivoting followed by a further Householder reduction step (see [9, p. 169] for the details). Now we apply Algorithm 2.1 to  $R$ , obtaining  $R = U_R H_R$ , and we "piece together" the polar factors of  $A$ . We have

$$\begin{aligned} A &= P \begin{bmatrix} U_R H_R & 0 \\ 0 & 0 \end{bmatrix} Q^* \\ &= P \begin{bmatrix} U_R & 0 \\ 0 & I_{m-r, n-r} \end{bmatrix} \begin{bmatrix} H_R & 0 \\ 0 & 0 \end{bmatrix} Q^* \\ &= P \begin{bmatrix} U_R & 0 \\ 0 & I_{m-r, n-r} \end{bmatrix} Q^* \cdot Q \begin{bmatrix} H_R & 0 \\ 0 & 0 \end{bmatrix} Q^* \\ &\equiv UH, \end{aligned}$$

where  $I_{m-r,n-r}$  denotes the  $(m-r) \times (n-r)$  identity matrix. Note that  $I_{m-r,n-r}$  could be replaced by any unitary matrix of the same dimensions; this shows the nonuniqueness of the unitary polar factor when  $r = \text{rank}(A) < \min(m, n)$ . Note also that even though  $U^*U \neq I$  when  $m < n$ ,  $H = U^*UH$  for all  $m$  and  $n$ ; thus  $A^*A = HU^*UH = H^2$ , so that  $H = (A^*A)^{1/2}$ .

In evaluating  $U$  and  $H$  advantage can be taken of the zero blocks in the products. Denoting by  $Q_1$  the first  $r$  columns of  $Q$ , we have

$$(2.1) \quad H = Q_1 H_R Q_1^*.$$

For  $U$  we partition

$$P = \begin{pmatrix} r & m-r \\ P_1 & P_2 \end{pmatrix}$$

and we distinguish the two cases:

$$(2.2a) \quad m \geq n \Rightarrow U = [P_1, P_2] \begin{bmatrix} U_R & 0 \\ 0 & \begin{bmatrix} I_{n-r} \\ 0 \end{bmatrix} \end{bmatrix} Q^* = \begin{bmatrix} P_1 U_R & P_2 \begin{bmatrix} I_{n-r} \\ 0 \end{bmatrix} \end{bmatrix} Q^*,$$

$$(2.2b) \quad m \leq n \Rightarrow U = [P_1, P_2] \begin{bmatrix} U_R & 0 \\ 0 & [I_{m-r}, 0] \end{bmatrix} Q^* = [P_1 U_R, P_2 [I_{m-r}, 0]] Q^*,$$

in which, respectively, the last  $m - n$  columns of  $P_2$  and the last  $n - m$  rows of  $Q^*$  need not participate in the multiplication.

To summarise, we have the following algorithm.

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ALGORITHM 2.2. [U, H] = polar(A, ε, δ).
% A ≠ 0 is arbitrary.
[P, R, Q] = COD(A, ε)
[UR, HR] = polar · square(R, δ)
Form U, H according to (2.1) and (2.2).
```

As the notation indicates, in floating-point arithmetic a tolerance  $\epsilon$  is required for the complete orthogonal decomposition to determine a numerical rank (i.e., the dimension of  $R$ ). The natural approach is to set to zero all rows of the trapezoidal QR factor of  $A$  that are negligible (in some measure) relative to  $\epsilon$  (see [9, p. 166]). The choice of  $\epsilon$  is important, since a small change in  $\epsilon$  can produce a large change in the computed  $U$  when  $A$  is rank-deficient. However, a redeeming feature is that whatever the choice of  $\epsilon$ , and irrespective of how well the QR factorization reveals rank, Algorithm 2.2 is stable, that is, the computed polar factors  $\hat{U}, \hat{H}$  satisfy

$$\hat{U}\hat{H} = A + E,$$

where  $\|E\|_F \approx \max\{\epsilon, \delta\} \|A\|_F$ ; this follows from the empirical stability of Algorithm 2.1 (see [11]) together with the stability of the additional orthogonal transformations in Algorithm 2.2.

The operation count of Algorithm 2.2 breaks down as follows, using the ‘‘flop’’ notation [9, p. 32]. The complete orthogonal decomposition requires  $2mnr - r^2(m + n) + 2r^3/3 + r^2(n - r)$  flops [9, pp. 165, 170]. Algorithm 2.1 requires, typically, eight iterations (assuming  $\delta \approx 10^{-16}$ ), and hence  $(7 + \frac{2}{3})r^3$  flops (taking into account the triangularity

of  $R$ ). And formation of  $H$  and  $U$  requires at most  $nr^2 + n^2r/2$  and  $mr^2 + \max\{nm^2, mn^2\}$  flops, respectively. By comparison, computing a polar decomposition via the Golub-Reinsch SVD algorithm requires approximately  $8mn^2 + 25n^3/6$  flops when  $m \geq n$ . The Golub-Reinsch SVD algorithm does not take advantage of rank-deficiency, although it could be modified to do so by using an initial complete orthogonal decomposition as above.

Of course, operation counts are not always a reliable guide to the actual computational cost on modern vector and parallel computers. An alternative performance indicator is the amount of matrix multiplication in an algorithm, since matrix multiplication can be performed very efficiently on many modern computers [1], [3], [18], [20]. As we will see in the next section, Algorithm 2.1 can be modified so that it is rich in matrix multiplication. In the complete orthogonal decomposition in Algorithm 2.2 the second Householder reduction step can be accomplished using the matrix multiplication rich WY representation of [3], [20]. In the initial QR factorization effective use of the WY representation is precluded by the column pivoting. One alternative is to use Bischof's local pivoting and incremental condition estimation technique [2], which does not hinder exploitation of the WY form. Another alternative is to compute a QR factorization *without* pivoting, and then to apply Chan's post-processing algorithm [5] for obtaining a rank-revealing QR factorization.

Finally, we show how to use Algorithm 2.1 to compute the Hermitian positive semidefinite square root of a Hermitian positive semidefinite  $A \in \mathbb{C}^{n \times n}$ . First, we compute a Cholesky decomposition with pivoting,

$$\Pi^T A \Pi = R^* R, \quad R = \begin{bmatrix} R_{11} & R_{12} \\ 0 & 0 \end{bmatrix},$$

where  $R_{11} \in \mathbb{C}^{r \times r}$  is nonsingular and upper triangular. Then Householder transformations are used to zero  $R_{12}$  (as in the complete orthogonal decomposition):

$$U^* \Pi^T A \Pi U = \begin{bmatrix} T_{11}^* & \\ 0 & \end{bmatrix} [T_{11} \quad 0], \quad T_{11} \in \mathbb{C}^{r \times r} \text{ upper triangular.}$$

Next, Algorithm 2.1 applied to  $T_{11}$  yields  $T_{11} = U_T H_T$ , whence, with  $Q = \Pi U$ ,

$$A = Q \begin{bmatrix} H_T^2 & 0 \\ 0 & 0 \end{bmatrix} Q^* = \left( Q \begin{bmatrix} H_T & 0 \\ 0 & 0 \end{bmatrix} Q^* \right)^2 \equiv X^2.$$

Square roots of semidefinite matrices are required in some statistical applications [10]. An alternative to this polar decomposition approach is to make use of an eigendecomposition; the relative merits are similar to those discussed above for the SVD.

**3. A hybrid iteration.** To make Algorithm 2.1 rich in matrix multiplication rather than matrix inversion, Schreiber and Parlett [19] use an inner Schulz iteration,

$$(3.1) \quad Z_{j+1} = Z_j + (I - Z_j X_k) Z_j, \quad Z_0 = X_k^{-1},$$

to compute  $X_k^{-1}$  on all iterations after the first. This approach takes advantage of the fact that since the  $X_k$  are converging quadratically,  $X_{k-1}^{-1}$  is an increasingly good approximation to  $X_k^{-1}$ . The Schulz iteration (3.1) is a Newton iteration and so also converges quadratically. Schreiber and Parlett observe that for the matrices in their application (which are often well conditioned) the typical number of inner iterations required for convergence is 6, 5, 3, 2, 1, leading to 17 iterations in total, or 34 matrix

multiplications. If the matrix inverses were computed directly, five inverses would be needed. This suggests that the modified algorithm will be faster than Algorithm 2.1 if matrix multiplication can be done at a rate  $34/5$  times faster than matrix inversion.

Further experimentation with the inner Schulz iteration led us to feel that it is unnecessary to run the inner iteration to convergence, and we considered employing just *one* Schulz iteration, with the starting matrix  $Z_0 = X_k^*$  ( $\approx X_k^{-1}$  since  $X_k$  converges to a unitary matrix). Thus the basic iteration

$$(3.2) \quad X_{k+1} = \frac{1}{2} \left( \gamma_k X_k + \frac{1}{\gamma_k} X_k^{-*} \right)$$

is replaced by (setting  $\gamma_k = 1$ )

$$(3.3) \quad \begin{aligned} X_{k+1} &= \frac{1}{2} (X_k + (Z_0^* + Z_0^* (I - Z_0 X_k)^*)) \\ &= X_k (I + \frac{1}{2} (I - X_k^* X_k)). \end{aligned}$$

This is precisely the quadratically convergent iteration of Kovarik [15] and Björck and Bowie [4] for computing the unitary polar factor! Hence, just a single inner Schulz iteration is enough to maintain quadratic convergence.

The convergence of (3.3) is described by the following relation, noted in [17]:

$$R_{k+1} = \frac{3}{4} R_k^2 + \frac{1}{4} R_k^3,$$

where  $R_k = I - X_k^* X_k$ . (Using this relation, we can show that the asymptotic error constant is  $\frac{3}{2}$  for (3.3) compared with  $\frac{1}{2}$  for (3.2) [11].) If  $\|R_k\| < 1$ , then

$$\|R_{k+1}\| < \frac{3}{4} \|R_k\|^2 + \frac{1}{4} \|R_k\|^2 = \|R_k\|^2 < \|R_k\|.$$

To maximise the number of matrix multiplications we therefore need to switch from iteration (3.2) to iteration (3.3) as soon as the convergence condition

$$(3.4) \quad \|X_k^* X_k - I\| \leq \theta < 1$$

is satisfied; to ensure fast convergence  $\theta$  should not be too close to 1. As explained below, typically (3.4) is satisfied for  $k=3$  with  $\theta=0.6$  (and obviously for  $k=0$  if  $X_0 = A$  happens to be nearly unitary). Rather than expend a matrix multiplication testing (3.4) we can use the matrix norm estimator CONEST from [12]. This computes a lower bound for  $\|C\|_1$  by sampling several matrix-vector products  $Cx$  and  $C^*x$ ; thus we can estimate  $\|X_k^* X_k - I\|_1$ , without forming  $X_k^* X_k$ , in  $O(r^2)$  flops (for  $r \times r$   $X_k$ ). A suitable way to use the estimate is to test whether it is less than  $\lambda\theta$ , where  $\lambda < 1$ . If so,  $X_k^* X_k - I$  is formed, in preparation for (3.3), and its norm is taken. If (3.4) is satisfied then (3.3) is used—otherwise we revert to iteration (3.2). The optimum choice of  $\lambda$  depends on the desired bias between wasting a matrix multiplication in an abortive switch of iteration, and not switching soon enough. The estimate from CONEST is almost always correct to within a factor 3, so  $\lambda \cong \frac{1}{3}$  is appropriate. In practice we have found that the performance of the algorithm is fairly insensitive to the choices of  $\theta$  and  $\lambda$ .

To summarise, our hybrid inversion/multiplication algorithm is as follows.

```
ALGORITHM 3.1. [U, H] = polar · mult (A, δ, λ, θ).
% A must be a square, nonsingular matrix.
X0 = A; k = -1; μ = 1; switched = false
repeat
```

```

k = k + 1
if switched
  R = I - X_k^* X_k; μ = ||R||_1
  evaluate (3.3)
else
  μ = CONEST (I - X_k^* X_k)
  if μ > λθ
    evaluate (3.2)
  else
    R = I - X_k^* X_k; μ = ||R||_1
    if μ > θ, evaluate (3.2), else evaluate (3.3), switched = true; end
  end
end
until μ ≤ δ
U = X_{k+1}
H = ½(U^* A + A^* U)

```

Since iteration (3.3) requires two matrix multiplications, and iteration (3.2) requires one inversion, Algorithm 3.1 will be more efficient than Algorithm 2.1 if matrix multiplication can be done at twice the rate of matrix inversion; thus, compared with using the full inner Schulz iteration, the “cutoff ratio” is 2 instead of 6.8. Moreover, if advantage is taken of the symmetry of the second matrix product in (3.3) the cutoff ratio is reduced to 1.5. The overall speedup depends on the ratio of inversions to multiplications, which in turn depends on the conditioning of the matrix, as discussed below.

All the algorithms mentioned here have been coded and tested in PC-MATLAB [16], running on an IBM PC-AT. For this machine the unit roundoff  $u \approx 2.22 \times 10^{-16}$ . We used  $\theta = .6$ ,  $\lambda = .75$ ,  $\delta = \sqrt{r} u$ , where  $r$  is the dimension of the matrix  $A$  in Algorithms 2.1 and 3.1, and  $\varepsilon = \max(m, n)|t_{11}|u$  in the complete orthogonal decomposition, where  $T$  is the triangular factor from the QR factorization with complete pivoting.

The following comments summarise our numerical experience, based on a wide variety of test matrices.

- Algorithms 2.1 and 3.1 usually require the same number of iterations. Occasionally Algorithm 3.1 requires one more iteration due to the larger error constant for iteration (3.3).

- In general, the typical number of iterations for Algorithm 3.1 is seven to nine, within the switch of iteration on iteration three or four.

TABLE 3.1

$k$	$\kappa_F(X_{k+1})$	$\ X_k^* X_k - I\ _1^{-1}$	Iteration	$\gamma_k$
0	2.6265E7	1.1380E10	(3.2)	3.1546E-3
1	5.3197E2	2.6233E4	(3.2)	8.0931E-3
2	4.0886E0	8.0962E-2 <sup>2</sup>	(3.3)	
3	3.9959E0	4.4915E-3	(3.3)	
4	4.0000E0	1.3686E-5	(3.3)	
5	4.0000E0	1.2607E-10	(3.3)	
6	4.0000E0	1.5765E-17	(3.3)	

<sup>1</sup> Norm estimate when (3.2) is used; exact quantity when (3.3) is used.

<sup>2</sup> Norm estimate exact to five digits.

• For well-conditioned matrices ( $\kappa_2(A) \leq 10$ , say), as are common in certain applications (see [13]), Algorithm 3.1 tends to require at most seven iterations and the switch is on iteration one, two, or three.

We present the results for one representative matrix in detail: MATLAB's "gallery(5)," which is the  $5 \times 5$  nilpotent matrix

$$A = \begin{bmatrix} -9 & 11 & -21 & 63 & -252 \\ 70 & -69 & 141 & -421 & 1684 \\ -575 & 575 & -1149 & 3451 & -13801 \\ 3891 & -3891 & 7782 & -23345 & 93365 \\ 1024 & -1024 & 2048 & -6144 & 24572 \end{bmatrix}.$$

Using Algorithm 3.1 within Algorithm 2.2, the numerical rank is diagnosed as 4, and Algorithm 3.1 is presented with a triangular matrix having one singular value of order  $10^5$  and three of order 1. Table 3.1 summarises the iteration. The backward error  $\|A - \hat{U}\hat{H}\|_1 = 4.7u\|A\|_1$ .

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