ALGORITHMS FOR HESSENNBERG-TRIANGULAR REDUCTION OF FIEDLER LINEARIZATION OF MATRIX POLYNOMIALS

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Abstract. Small- to medium-sized polynomial eigenvalue problems can be solved by linearizing the matrix polynomial and solving the resulting generalized eigenvalue problem using the QZ algorithm. The QZ algorithm, in turn, requires an initial reduction of a matrix pair to Hessenberg-triangular (HT) form. In this paper, we discuss the design and evaluation of high-performance parallel algorithms and software for HT reduction of a specific linearization of matrix polynomials of arbitrary degree. The proposed algorithm exploits the sparsity structure of the linearization to reduce the number of operations and improve the cache reuse compared to existing algorithms for unstructured inputs. Experiments on both a workstation and a high-performance computing system demonstrate that our structure-exploiting parallel implementation can outperform both the general LAPACK routine DGGEHR and the prototype implementation DGGEHR3 of a general blocked algorithm.

Key words. Hessenberg-triangular reduction, polynomial eigenvalue problem, linearization, blocked algorithm, parallelization

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1. Introduction. A complex matrix polynomial of degree $d$ takes the form

$$P(\lambda) = \lambda^d P_d + \lambda^{d-1} P_{d-1} + \cdots + \lambda P_1 + P_0,$$

where $P_k \in \mathbb{C}^{n \times n}$ and $P_d \neq 0$. Throughout this paper we assume that $P(\lambda)$ is regular, that is, that $\det P(\lambda)$ is not identically zero. The polynomial eigenvalue problem consists of finding scalars $\lambda \in \mathbb{C}$ and nonzero vectors $x, y \in \mathbb{C}^n$ satisfying the equations $P(\lambda)x = 0$ and $y^* P(\lambda) = 0$. The vectors $x$ and $y$ are right and left eigenvectors, respectively, associated with the eigenvalue $\lambda$. The finite eigenvalues of $P$ are roots of the scalar equation $\det P(\lambda) = 0$. When $\det P(\lambda) = 0$ has degree $r < dn$, we add $dn - r$ eigenvalues at infinity, which correspond to the zero eigenvalues of the reversal polynomial $\lambda^d P(1/\lambda)$.

A standard approach to solving a polynomial eigenvalue problem is to construct from the polynomial $P$ a linear polynomial

$$L(\lambda) = A + \lambda B, \quad A, B \in \mathbb{C}^{dn \times dn},$$

with the same spectrum as $P$. The associated generalized eigenvalue problem

$$(1.1) \quad L(\lambda) z = 0, \quad w^* L(\lambda) = 0$$

can then be solved using standard techniques and readily available software. Unless the dimension $dn$ of $L$ is very large, dense transformation-based methods can be
used to solve (1.1). This amounts to reducing the matrix pair $(A, B)$ to Hessenberg-triangular (HT) form by a unitary equivalence transformation $(A, B) \mapsto (Q^* AZ, Q^* BZ)$, where $Q, Z \in \mathbb{C}^{d \times d}$ are unitary matrices, followed by the QZ algorithm to obtain a generalized Schur form, again with a unitary equivalence transformation.

There are infinitely many linearizations to choose from, and they tend to be structured and relatively sparse [1, 4, 14]. In this paper, we exploit the structure of a particular Fiedler linearization to devise an efficient algorithm for HT reduction that has lower arithmetic and communication costs than general HT reduction algorithms.

The Fiedler linearization of interest is illustrated for the case $d = 5$ as

\begin{equation}
L(\lambda) = A + \lambda B = \begin{bmatrix}
P_4 & P_3 & P_2 & P_1 & -I \\
-I & 0 & 0 & 0 & 0 \\
0 & -I & 0 & 0 & 0 \\
0 & 0 & -I & 0 & 0 \\
0 & 0 & 0 & P_0 & 0
\end{bmatrix} + \lambda \begin{bmatrix}
P_5 & 0 & 0 & 0 & 0 \\
0 & I & 0 & 0 & 0 \\
0 & 0 & I & 0 & 0 \\
0 & 0 & 0 & I & 0 \\
0 & 0 & 0 & 0 & I
\end{bmatrix},
\end{equation}

with a straightforward generalization to arbitrary degrees (see also Appendix A). The sparsity structure is illustrated in pictorial form in Figure 1. This particular linearization is used by the eigensolver *quadeig* for dense quadratic ($d = 2$) eigenvalue problems [5]. It has several desirable properties. First, $L(\lambda)$ is always a linearization of $P(\lambda)$ in the sense that $E(\lambda)L(\lambda)F(\lambda) = \text{diag}(P(\lambda), I_{(d-1)n})$ for some matrix polynomials $E(\lambda)$ and $F(\lambda)$ with constant nonzero determinant. As a result, $\det P(z) = \gamma \cdot \det L(\lambda)$ for some nonzero constant $\gamma$, so that $P$ and $L$ have the same finite eigenvalues. In fact $L(\lambda)$ is a strong linearization: it has the same finite and infinite elementary divisors as $P$ [13, Thm. 3]. The pencil $L(\lambda)$ in (1.2) is easy to construct from the coefficients $F_i$ of $P(\lambda)$, without performing any arithmetic operation. The right and left eigenvectors $x$ and $y$ of $P(\lambda)$ are easily recovered from those of $L(\lambda)$. Indeed, if $z$ and $w$ are right and left eigenvectors of $L(\lambda)$, then

\[
x = \begin{cases}
(e_j^T \otimes I)z & \text{if } \lambda = \infty, \\
(e_j^T \otimes I)z & 1 \leq j \leq d - 1 \text{ if } \lambda \in \mathbb{C},
\end{cases}
\quad y = \begin{cases}
(e_j^T \otimes I)w & \text{if } \lambda = \infty, \\
(e_j^T \otimes I)w & j = 1, d \text{ if } \lambda \in \mathbb{C},
\end{cases}
\]

where $e_j$ denotes the $j$th column of the $n \times n$ identity matrix and $\otimes$ denotes the Kronecker product. Also, when $\|P_i\|_2 \approx 1$ for $i = 0: d$, then $L(\lambda)$ has good conditioning and backward stability properties [3, 6, 7]. Moreover, the pair $(A, B)$ is almost already in HT form since $A$ is in upper block Hessenberg form with $2n - 1$ subdiagonals, and $B$ is in block upper triangular form. It is straightforward to transform $(A, B)$ using unitary equivalence such that $A$ becomes upper block Hessenberg with only $n$ subdiagonals, and $B$ becomes upper triangular. Furthermore, we show in Appendix A that prior to the HT reduction, deflation of zero and infinite eigenvalues contributed by
singular trailing and leading coefficient matrices can be performed without altering the block structure of \((A, B)\) in (1.2). This is in part due to the position of \(P_0\) and \(P_d\) in \(A\) and \(B\).

The rest of the paper is organized as follows. The preprocessing step is described in section 2. A known algorithm for reduction from block HT form to proper HT form is recalled in section 3, and its effect on the sparsity structure when applied to inputs of the form (1.2) is analyzed. This is combined with recent work on cache-efficient algorithms for the unstructured case in section 4, and the construction of cache-efficient algorithms tailored for the structured case is described thereafter. In section 5 we discuss a parallelization scheme suitable for multicore-based systems. Numerical results are presented in section 6, and section 7 concludes.

2. Preprocessing. If \(P_0\) is singular, then \(P(\lambda)\) has zero eigenvalues since \(P(0)x = P_0x = 0\) for \(x \in \text{null}(P_0)\). Similarly, if \(P_d\) is singular, then \(P(\lambda)\) has eigenvalues at infinity: these are the zero eigenvalues of the reversal polynomial \(\lambda^d P(1/\lambda)\). Although the QZ algorithm handles eigenvalues at infinity well [15], numerical experiments by Kågström and Kressner [8] show that if infinite eigenvalues are not extracted before starting the QZ steps, then they may never be detected due to the effect of rounding errors in floating point arithmetic. It is then desirable to deflate these eigenvalues before the HT reduction step. Starting from the linearization (1.2), we show in Appendix A how to deflate zero and infinite eigenvalues of \(P(\lambda)\) contributed by singular leading and trailing coefficients, resulting in a pencil \(A + \lambda B\) of dimension \((d-2)n + r_0 + r_d\) with the same block structure as in (1.2) but with blocks of different dimensions. Here \(r_i = \text{rank}(P_i)\) for \(i = 0, d\).

To avoid clutter in the notation and for simplicity we assume in what follows that no deflation occurred so that all the blocks in (1.2) are \(n \times n\). The block upper triangular matrix \(B\) can immediately be reduced to upper triangular form by a QR factorization of its top left block (i.e., \(P_d\)). The lower bandwidth of the block upper Hessenberg matrix \(A\) can be reduced from \(2n - 1\) to \(n\) by a QR factorization of its bottom right subdiagonal block (i.e., \(P_0\)). More specifically, let \(P_d = Q_d R_d\) and \(P_0 = Q_0 R_0\) be QR factorizations of \(P_d\) and \(P_0\), respectively. By defining the transformation matrices

\[
Q = \begin{bmatrix} Q_d & 0 & 0 \\ 0 & I_{(d-2)n} & 0 \\ 0 & 0 & Q_0 \end{bmatrix}, \quad Z = \begin{bmatrix} I_n & 0 & 0 \\ 0 & I_{(d-2)n} & 0 \\ 0 & 0 & Q_0 \end{bmatrix}
\]

(see Figure 2 for their pictorial form), the unitary equivalence transformation \((A, B) \leftrightarrow (Q^*AZ, Q^*BZ)\) accomplishes the desired preprocessing step. Following this transformation, the matrix pair \((A, B)\) is in block HT form with \(n\) subdiagonals in \(A\). An example for the case \(d = 5\) takes the form

\[
A + \lambda B = \begin{bmatrix} Q_5P_4 & Q_5^*P_3 & Q_5^*P_2 & Q_5^*P_1 & -Q_5^*Q_0 \\ -I & 0 & 0 & 0 & 0 \\ 0 & -I & 0 & 0 & 0 \\ 0 & 0 & -I & 0 & 0 \\ 0 & 0 & 0 & R_0 & 0 \end{bmatrix} + \lambda \begin{bmatrix} R_5 & 0 & 0 & 0 & 0 \\ 0 & I & 0 & 0 & 0 \\ 0 & 0 & I & 0 & 0 \\ 0 & 0 & 0 & I & 0 \\ 0 & 0 & 0 & 0 & I \end{bmatrix}
\]

(2.2)

The pictorial form of (2.2) is illustrated in Figure 3.

3. Exploiting structure. This section introduces the basic bulge-chasing algorithm that forms the foundation of the proposed algorithms.
3.1. The basic algorithm. Assume that the pair \((A, B)\) has already been preprocessed as explained in section 2 and that \(Q\) and \(Z\) are the resulting transformation matrices. The aim of this section is to recall a known algorithm that takes a matrix pair in block HT form and reduces it to proper HT form via a bulge-chasing procedure. This algorithm appears as the second stage of a two-stage HT reduction algorithm for general dense matrix pairs [2, 9].

Algorithm 1 is a reformulation of the bulge-chasing algorithm using our notation. The inputs to Algorithm 1 include the degree, \(d\), and size, \(n\), of the polynomial matrix as well as the preprocessed pair \((A, B)\) and the resulting transformation matrices \(Q\) and \(Z\). The algorithm reduces \((A, B)\) from block HT form with \(n\) nonzero subdiagonals to proper HT form by systematically applying a carefully chosen set of plane rotations. The product of the rotations corresponds to a unitary equivalence transformation of \((A, B)\). The algorithm returns the reduced matrix pair \((A, B)\) in proper HT form along with the updated transformation matrices \(Q\) and \(Z\).

In section 3.1.1 we define the computational kernels that are introduced in Algorithm 1. The algorithm itself and its effect on unstructured inputs are described in section 3.2. In section 3.3 we present a block partitioning of \(A\) and \(B\) that comes naturally from the algorithm. The definition of the block partitioning depends on the iteration index \(c\) in the sense that the block boundaries shift one step southeast from one iteration to the next. To avoid ambiguities we introduce notation for referring to a specific block in a specific iteration. The effect of the algorithm on the sparsity structure of a preprocessed pair \((A, B)\) from section 2 is analyzed in section 3.4. The result of this analysis forms the basis for our further developments. How structures in the transformation matrices are exploited is described in sections 3.5 and 3.6, and a structure-exploiting algorithm is presented in section 3.7.
Algorithm 1: $A, B, Q, Z \leftarrow \text{HTBasic}(n, d, A, B, Q, Z)$. Given a matrix pair $(A, B)$ with matrices of size $dn \times dn$ in upper block HT form with $n$ nonzero subdiagonals in $A$, this algorithm reduces $(A, B)$ to proper HT form and accumulates the associated transformations into the given transformation matrices $Q$ and $Z$.

1. for $c \leftarrow 1$ to $dn - 2$ do
2.   for $r \leftarrow \min\{dn, c + n\}$ down to $c + 2$ do
3.     $\tilde{c} \leftarrow c$;
4.     $\tilde{r} \leftarrow r$;
5.     while $\tilde{r} \leq dn$ do
6.        /* Zero out $A(\tilde{r}, \tilde{c})$ with a row rotation. */
7.            $G \leftarrow \text{Zero}(A, \tilde{r}, \tilde{c})$;
8.            $\text{LeftRot}(G, A, \tilde{r} - 1, \tilde{r}, \tilde{c}, dn)$;
9.            $\text{LeftRot}(G, B, \tilde{r} - 1, \tilde{r}, \tilde{r} - 1, dn)$;
10.           $\text{RightRot}(G, Q, 1, dn, \tilde{r} - 1, \tilde{r})$;
11.          /* Move from $A$ to $B$. */
12.             $\tilde{c} \leftarrow \tilde{r} - 1$;
13.           /* Zero out $B(\tilde{r}, \tilde{c})$ using a column rotation. */
14.             $G \leftarrow \text{RightZero}(B, \tilde{r}, \tilde{c})$;
15.             $\text{RightRot}(G, A, 1, \min\{dn, \tilde{c} + n + 1\}, \tilde{c}, \tilde{c} + 1)$;
16.             $\text{RightRot}(G, B, 1, \tilde{r}, \tilde{c}, \tilde{c} + 1)$;
17.           /* Move from $B$ to $A$. */
18.             $\tilde{r} \leftarrow \tilde{c} + n + 1$;

3.1.1. Computational kernels introduced in HTBasic. The four computational kernels that are introduced in Algorithm 1 are described below.

- $G \leftarrow \text{Zero}(A, r, c)$.
  Constructs a row rotation $G$ that zeroes out $A(r, c)$ using $A(r - 1, c)$ as a pivot.

- $G \leftarrow \text{RightZero}(A, r, c)$.
  Constructs a column rotation $G$ that zeroes out $A(r, c)$ using $A(r, c + 1)$ as a pivot.

- $\text{LeftRot}(G, A, r_1, r_2, c_1, c_2)$.
  Applies a row rotation $G$ to rows $r_1$ and $r_2$ of $A$. The effect is limited to the column range $c_1 : c_2$.

- $\text{RightRot}(G, A, r_1, r_2, c_1, c_2)$.
  Applies a column rotation $G$ to columns $c_1$ and $c_2$ of $A$. The effect is limited to the row range $r_1 : r_2$.

3.2. The effect of HTBasic on unstructured inputs. Algorithm 1 systematically zeroes out the undesired nonzero entries below the first subdiagonal of $A$. The entries are zeroed one column at a time from left to right (loop over $c$ on line 1) and from the bottom up within each column (loop over $r$ on line 2). The entry $A(r, c)$ is zeroed out by a row rotation involving rows $r - 1$ and $r$ constructed on line 6.

A row rotation on rows $r - 1$ and $r$ introduces an undesired nonzero entry (a fill or bulge) in the subdiagonal entry $B(r, r - 1)$. Unless the bulge is removed, the lower
triangular part of $B$ will rapidly fill in, and its sparsity structure will be destroyed. Therefore, the bulge is removed using a column rotation that acts on columns $r - 1$ and $r$. The rotation is constructed on line 11 and applied to $A$, $B$, and $Z$ on the following lines. This creates, however, a second bulge at $A(r + n, r - 1)$ unless $r + n > dn$, in which case no new bulge appears. At most one bulge is present at any given time either in $A$ or in $B$. It helps to interpret the algorithm as if it moves a single bulge back and forth between $A$ and $B$ until the bulge finally disappears near the bottom right corner. The precise movement of the bulge is illustrated in Figure 4.

Algorithm 1 uses the two variables $\tilde{r}$ and $\tilde{c}$ to keep track of the current position of the bulge. In each iteration of the while-loop on line 5, an entry is zeroed out in $A$, then a bulge is generated and zeroed out in $B$, and finally the bulge reappears in $A$. The loop continues to chase the bulge until it disappears.

3.3. Block partitioning induced by HTBasic. The computations associated with iteration $c$ of the outer loop in Algorithm 1 induce a natural block partitioning of $A$ and $B$, as illustrated by the solid lines in Figures 4 and 5 for the first iteration ($c = 1$). The block partitioning depends on the loop index $c$, and for iteration $c \in \{1, 2, \ldots, dn - 2\}$, the $(1, 1)$ block has size $c \times c$. The other blocks on the diagonal all have size $n \times n$, except possibly the last block, which may be smaller. It follows that the blocks on the subdiagonal generally have size $n \times n$, except possibly the first and the last blocks, which may be rectangular. As the partitioning is symmetric, the number of blocks in either dimension is

$$N_c = \left\lceil \frac{dn - c}{n} \right\rceil + 1.$$  

The dependence on $c$ manifests itself as a shift by one down the diagonal from one iteration to the next, as illustrated by the dashed lines (red online) in Figure 5 for the second iteration ($c = 2$).

To unambiguously refer to a specific block relative to a given partitioning/iteration, we introduce the following notation. The block $(i, j)^X_c$ refers to block $(i, j)$ of matrix $X$ (either $X = A$ or $X = B$) with respect to the block partitioning in iteration $c$. 

![Figure 4. Illustration of the bulge movement in Algorithm 1 for the case $d = 5$, $n = 4$, $c = 1$, and $r = 4$ when the algorithm is applied to unstructured inputs. The matrix $A$ is displayed on the left and $B$ on the right.](image)

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3.4. The effect of HTBasic on structured inputs. We now study the effect of Algorithm 1 on the sparsity structure when the input is a preprocessed matrix pair as defined in section 2. Figure 6 illustrates the generic sparsity structure of $A$ and $B$ for the case $d = 5$ and $n = 4$. Even though Algorithm 1 eventually destroys the structure of all four of the involved matrices ($A$, $B$, $Q$, and $Z$), the process turns out to be gradual, and the evolving sparsity structure can be exploited, as we will soon explain, to reduce both the arithmetic and communication costs.

Informally, the initial sparsity structure shown in Figure 6 evolves during the execution of Algorithm 1 as follows. The dense upper triangular block of $B$ grows slowly by one column to the right for each iteration of the outer for-loop. The dense first block row of $A$ also grows slowly by one row for each iteration of the outer for-loop. The upper triangular block of $A$, however, grows rapidly by one row for each iteration of the inner for-loop. Moreover, this block also grows slowly by one column for each iteration of the outer for-loop.

We refer to one iteration of the inner for-loop on line 2 of Algorithm 1 as a sweep, and Figure 7 illustrates the sparsity structure obtained after the first sweep. Note
the appearance of a new column on the triangular block of $B$, a new row on the
dense first block row of $A$, and a new row and column on the upper triangular block
of $A$. The two remaining sweeps in the current iteration of the outer for-loop each
add one row to the triangular block of $A$, but they have no other effect on the sparsity
structures of either $A$ or $B$. After completing the first iteration of the outer for-loop,
the sparsity structure takes on the form illustrated in Figure 8. The effect on the
sparsity structure is similar for all iterations. It follows that after $d-2$ iterations the
two dense blocks of $A$ will fuse together, and after $n$ iterations the triangular block
of $A$ will reach the east boundary of the matrix.

These observations suggest that some of the arithmetic operations required to
update the matrix $B$ can be eliminated since they operate on zeros and thus have
no effect. A fraction of the arithmetic operations required to update the matrix $A$
can be eliminated as well, but in general this cost reduction is more modest since
the structure of $A$ is rapidly destroyed. We exploit in our algorithms only the block
**3.5. Exploiting structure in the transformation matrices.** The transformation matrices $Q$ and $Z$ are thickly banded matrices with some additional structure within the band at all times during the whole accumulation process. More precisely, they each take the form of a skyline matrix where the top and bottom skylines are nondecreasing. Formally, let $t_j$ and $b_j$ denote the row index of the first, respectively, last (structural) nonzero in column $j$. Then $t_{j+1} \geq t_j$ and $b_{j+1} \geq b_j$. By explicitly keeping track of the $t_j$ and $b_j$ during the accumulation procedure, it is straightforward and inexpensive to determine the minimal range of rows to which a given rotation must be applied and thereby optimally reduce the number of arithmetic operations.

**3.6. Exploiting redundancy in the transformation matrices.** The transformation matrices $Q$ and $Z$ obtained from the preprocessing step (section 2) are actually identical except for their top left $n \times n$ block. Due to the identities on the main block diagonal of $B$, many of the row and column rotations are identical, too (see section 3.7). As it turns out, while $Q$ and $Z$ gradually diverge, they maintain a shrinking number of identical columns. Specifically, the number of columns shared by $Q$ and $Z$ shrinks by one column for every column reduced. Immediately before reducing column $c$, the matrices $Q$ and $Z$ can be partitioned into column blocks

$$Q = \begin{bmatrix} Q_1 & Q_2 \end{bmatrix} \quad \text{and} \quad Z = \begin{bmatrix} Z_1 & Z_2 \end{bmatrix}$$

such that both $Q_1$ and $Z_1$ have $\min\{dn, n + c - 1\}$ columns and $Q_1 \neq Z_1$ (in general) and $Q_2 = Z_2$ (always). During the reduction of column $c$ in Algorithm 1, a sequence of rotations will engage the rightmost parts of $Q_1$ as well as the leftmost column of $Q_2$. Similarly, another sequence of rotations will engage the rightmost parts of $Z_1$ and the leftmost column of $Z_2$. The subsequent rotations will affect only $Q_2$ and $Z_2$ and, crucially, will not touch the leftmost columns of either $Q_2$ or $Z_2$. Since the lower part of $B$ contains only identities, the remaining rotations accumulated into $Q$ and $Z$ are identical, and hence the last columns of these matrices will remain identical. In summary, at the end of iteration $c$, the leftmost columns of $Q_2$ and $Z_2$ will differ, but the other columns will remain identical. This allows us to avoid the redundant accumulation of rotations into both $Q_2$ and $Z_2$ by accumulating only into $Q_2$ and prior to iteration $c$ copying the leftmost column of $Q_2$ into the corresponding column of $Z_2$. See Figure 9 for a pictorial representation.

![Diagram](image_url)
3.7. The structure-exploiting algorithm. This section presents the algorithm HTStructured (Algorithm 2), which exploits most of the aforementioned structures. In addition, the algorithm pipelines the bulge-chasing procedure to improve the cache reuse (which is the topic of section 4 below).

The steps of HTBasic (Algorithm 1) can be reordered such that all bulges stemming from iteration \( c \) are chased together in a pipelined fashion. This is made possible by the associativity of matrix multiplication. In floating point arithmetic the different ordering leads to differences in the computed results, but both HTBasic and HTStructured are backward stable. We refer the reader to section 3.7.1 for descriptions of the computational kernels that are introduced in the structure-exploiting algorithm.

Algorithm 2: \( A, B, Q, Z \leftarrow \text{HTStructured}(n, d, A, B, Q, Z) \).

```plaintext
/* For each column left to right. */
for \( c \leftarrow 1 \) to \( dn - 2 \) do
  /* Copy one column from \( Q \) to \( Z \) to avoid the redundancy. */
  if \( c + n \leq dn \) then \( Z(:,c+n) \leftarrow Q(:,c+n) \);
  /* Locate column \( c \) of \( A \). */
  \( r_1, r_2 \leftarrow c + 1, \min\{dn,c+n\} \);
  /* Zero out column \( c \) of \( A \) using row rotations. */
  \( \mathcal{G} \leftarrow \text{LeftZero}(A, r_1, r_2, c) \);
  LeftSkyline(\( \mathcal{G}, A, r_1, r_2, c+1, dn \));
  /* Move from \( A \) to \( B \). */
  \( c_1, c_2 \leftarrow r_1, r_2 \);
  /* Generate and zero out bulges in \( B \). */
  \( \mathcal{G} \leftarrow \text{LeftRightZero}(\mathcal{G}, B, r_1, r_2, c_1, c_2) \);
  Right(\( \mathcal{G}, B, 1, r_1 - 1, c_1, c_2 \));
  RightSkyline(\( \mathcal{G}, Z, c_1, c_2 \));
  /* Chase bulges until they disappear. */
  loop
    /* Move from \( B \) to \( A \). */
    \( r_1, r_2 \leftarrow c_1 + n, \min\{dn,c_2+n\} \);
    /* Update \( A \) using column rotations. */
    Right(\( \mathcal{G}, A, 1, \min\{dn,r_1 - 1\}, c_1, c_2 \));
    /* Exit if all bulges have disappeared. */
    if \( r_1 > dn \) then exit loop;
    /* Generate and zero out bulges in \( A \). */
    \( \mathcal{G} \leftarrow \text{RightLeftZero}(\mathcal{G}, A, r_1, r_2, c_1, c_2) \);
    Left(\( \mathcal{G}, A, r_1, r_2, c_2 + 1, \min\{dn,(d-1)n+c\} \));
    RightSkyline(\( \mathcal{G}, Q, r_1, r_2 \));
    /* Move from \( A \) to \( B \). */
    \( c_1, c_2 \leftarrow r_1, r_2 \); martyr
```

Algorithm 2 uses the concept of a sequence of rotations, that is, a sequence \( \mathcal{G} = \langle G_1, G_2, \ldots, G_m \rangle \) consisting of \( m \) rotations acting on adjacent pairs of rows or columns.

More specifically, for an increasing sequence \( \mathcal{G} \), the rotation \( G_k \) acts on rows/columns \( i + k - 1 \) and \( i + k \) for some constant base index \( i \). In other words, the rotation
sequence $G$ affects the $m+1$ rows or columns $i: i+m$. For a decreasing sequence $G$, rotation $G_k$ acts on rows or columns $i+m-k$ and $i+m-k+1$. Again, the sequence affects the rows or columns $i: i+m$. We encounter only decreasing sequences in what follows.

The structure-exploiting algorithm exploits the structure of $B$ and the structure of the last block column of $A$ (but ignores the structure of its interior). The structures of the transformation matrices and the redundancy present in their accumulation are also exploited. The variables $r_1$, $r_2$, $c_1$, and $c_2$ keep track of the submatrix (of $A$ or $B$, depending on the context) that currently contains (or is about to contain) the set of bulges. The outer loop iterates over the columns from left to right. Column $c$ of $A$ is zeroed out on line 4, and the resulting sequence of rotations is immediately applied to $A$ and accumulated into $Q$. The resulting bulges in $B$ are generated and immediately zeroed out on line 8 followed by further updates of $B$ and accumulation into $Z$. Entering the loop on line 11, the column rotations are applied to $A$ on line 13. A sequence of row rotations is constructed on line 15 to zero out the resulting bulges in $A$ followed by updates of $A$ and accumulation into $Q$. The affected portion of $B$ is an identity matrix, and therefore any row rotation generates a bulge that is zeroed out by the same rotation applied to the columns. Hence, the net effect on $B$ is nil, and there is no need to explicitly generate the bulges or even reference $B$.

3.7.1. Computational kernels introduced in HTStructured. Algorithm 2 introduces the following computational kernels.

- $G \leftarrow \text{LeftZero}(A, r_1, r_2, c)$.
  Constructs a decreasing sequence $G$ consisting of $r_2 - r_1$ row rotations that zeroes out all entries except the first one in the column vector $A(r_1: r_2, c)$. The vector is updated in the process.

- $G \leftarrow \text{Left}(G, A, r_1, r_2, c_1, c_2)$.
  Applies a decreasing sequence consisting of $r_2 - r_1$ row rotations $G$ to the submatrix $A(r_1: r_2, c_1: c_2)$.

- $G \leftarrow \text{Right}(G, A, r_1, r_2, c_1, c_2)$.
  Applies a decreasing sequence consisting of $c_2 - c_1$ column rotations $G$ to the submatrix $A(r_1: r_2, c_1: c_2)$.

- $G \leftarrow \text{RightSkyline}(G, A, c_1, c_2)$.
  Identical to Right except that the skyline sparsity pattern of the matrix determines the optimal range of rows to update with each rotation.

- $G \leftarrow \text{LeftRightZero}(G, B, r_1, r_2, c_1, c_2)$.
  Applies a decreasing sequence consisting of $r_2 - r_1$ row rotations $G$ to the upper triangular and square submatrix $B(r_1: r_2, c_1: c_2)$ while at the same time constructing and applying a decreasing sequence consisting of $c_2 - c_1$ column rotations $G$ that zeroes out the bulges created by $G$. The updated submatrix is again upper triangular.

- $G \leftarrow \text{RightLeftZero}(G, A, r_1, r_2, c_1, c_2)$.
  Applies a decreasing sequence consisting of $c_2 - c_1$ column rotations $G$ to the upper trapezoidal submatrix $A(r_1: r_2, c_1: c_2)$ while at the same time constructing and applying a decreasing sequence consisting of $r_2 - r_1$ row rotations $G$ that zeroes out the bulges created by $G$. The updated submatrix is again upper trapezoidal.

3.8. Cost analysis. In this section, we analyze the arithmetic cost of three different algorithms to understand how large the impact of exploiting structure is. The following are the algorithms included in the analysis.

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**Algorithm I.** Basic algorithm (DGGHRD in LAPACK), where $A$, $Q$, and $Z$ are dense and $B$ is triangular.

**Algorithm II.** The HTBasic algorithm on unstructured inputs, where $A$ has $n$ subdiagonals, $Q$ and $Z$ are dense, and $B$ is triangular.

**Algorithm III.** The HTStructured algorithm, where $A$ has $n$ subdiagonals and a structured final block column, $Q$ and $Z$ are structured, $B$ is structured, and the redundancy in the accumulation of $Q$ and $Z$ is exploited.

For a fixed degree $d$, the arithmetic complexity of all three algorithms is $O(n^3)$. The coefficient of the leading $n^3$ term is for each algorithm a third degree polynomial in $d$. This polynomial can be written as the sum of six polynomials, one each for left/right updates of $A$, left/right updates of $B$, and right updates of $Q$ and $Z$. The analysis does not include the overhead of accumulating rotations and applying them as matrix-matrix multiplications. The application of a rotation to a pair of scalars is counted as six arithmetic operations.

The coefficients of the polynomials reported below were obtained through the simulation of the algorithms using a binary matrix to keep track of structural nonzeros. The exact arithmetic operation counts were obtained for simulation of the algorithms using a binary matrix to keep track of structural nonzeros. The exact arithmetic operation counts were obtained for $n = 100$ and $d = 2, 3, \ldots, 10$. After dividing the counts by $n^3$, a third degree polynomial in $d$ was fitted in the least squares sense, and the resulting coefficients were rounded to the nearest integer. The results were checked by obtaining counts for the larger problem $n = 200$, and the relative error of the model was smaller for the larger problem, which is what one would expect from an accurate model of the leading term.

The coefficients of the leading $n^3$ term for each algorithm and their six major parts are given in the final row of Table 1. The first six rows show the major parts whose sum make up the coefficient of the leading term. The seventh and eighth rows show the arithmetic cost associated with $A$ and $B$, respectively, $Q$ and $Z$. The terms marked in boldface are those that differ from the corresponding terms in the preceding column.

**Table 1**

<table>
<thead>
<tr>
<th></th>
<th>Alg. I</th>
<th>Alg. II</th>
<th>Alg. III</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left $A$</td>
<td>$2d^3$</td>
<td>$d^3 + 3d^2 - 3d + 1$</td>
<td>$d^3 + 3d^2 - 6d + 5$</td>
</tr>
<tr>
<td>Right $A$</td>
<td>$3d^3$</td>
<td>$2d^3 + 3d^2 - 3d + 1$</td>
<td>$2d^3 + 3d^2 - 6d + 5$</td>
</tr>
<tr>
<td>Left $B$</td>
<td>$d^3$</td>
<td>$d^3$</td>
<td>$3d - 2$</td>
</tr>
<tr>
<td>Right $B$</td>
<td>$2d^3$</td>
<td>$2d^3$</td>
<td>$3d^2 - 1$</td>
</tr>
<tr>
<td>Right $Q$</td>
<td>$3d^3$</td>
<td>$3d^3$</td>
<td>$2d^3 + 3d - 1$</td>
</tr>
<tr>
<td>Right $Z$</td>
<td>$3d^3$</td>
<td>$3d^3$</td>
<td>$6d^2 - 3$</td>
</tr>
<tr>
<td>$\Sigma_{A,B}$</td>
<td>$8d^3$</td>
<td>$6d^3 + 6d^2 - 6d + 2$</td>
<td>$3d^3 + 9d^2 - 9d + 7$</td>
</tr>
<tr>
<td>$\Sigma_{Q,Z}$</td>
<td>$6d^3$</td>
<td>$6d^3$</td>
<td>$2d^3 + 6d^2 - 1$</td>
</tr>
<tr>
<td>$\Sigma$</td>
<td>$14d^3$</td>
<td>$12d^3 + 6d^2 - 6d + 2$</td>
<td>$5d^3 + 15d^2 - 9d + 6$</td>
</tr>
</tbody>
</table>

Table 2 shows the actual coefficients for a few concrete values of $d$ as well as the ratio of the costs of Algorithm II and Algorithm III relative to Algorithm I.

A conclusion from this analysis is that exploiting the structure of $A$ has a very small effect. The real benefit comes from exploiting the structures of $B$, $Q$, and $Z$ and the redundancy between $Q$ and $Z$. The benefit is larger for larger $d$. There is a reduction in the arithmetic cost for Algorithm III compared to Algorithm I and Algorithm II, but this reduction remains modest.
Table 2
The actual coefficient of the leading $n^3$ term in the arithmetic cost of three different HT reduction algorithms for some concrete values of $d$.

<table>
<thead>
<tr>
<th></th>
<th>$d = 2$</th>
<th>$d = 3$</th>
<th>$d = 4$</th>
<th>$d = 5$</th>
<th>$d \to \infty$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alg I</td>
<td>112</td>
<td>378</td>
<td>896</td>
<td>1750</td>
<td></td>
</tr>
<tr>
<td>Alg II</td>
<td>110</td>
<td>362</td>
<td>842</td>
<td>1622</td>
<td></td>
</tr>
<tr>
<td>Alg III</td>
<td>88</td>
<td>249</td>
<td>530</td>
<td>961</td>
<td></td>
</tr>
<tr>
<td>Alg II / Alg I</td>
<td>0.98</td>
<td>0.96</td>
<td>0.94</td>
<td>0.93</td>
<td>0.86</td>
</tr>
<tr>
<td>Alg III / Alg I</td>
<td>0.79</td>
<td>0.66</td>
<td>0.59</td>
<td>0.55</td>
<td>0.36</td>
</tr>
</tbody>
</table>

4. Optimizing for the cache. The algorithms mentioned thus far access memory with a large stride and perform only a few arithmetic operations for every accessed matrix entry. Following previous work on HT reduction for general inputs [9], we apply in this section a set of standard techniques that reduce the number of cache and translation lookaside buffer (TLB) misses as well as the volume of communication between cache and main memory.

The HTStructured algorithm already includes a key cache-optimization technique: pipelining the bulge-chasing steps from the reduction of one column.

4.1. Column striping. Assuming that the matrices are represented in the column-major storage format, a column rotation can be implemented such that it accesses two contiguous streams of memory, but a row rotation will necessarily access memory with a large stride. Since the structure-exploiting algorithm, due to the pipelining, consistently applies sequences of rotations instead of individual rotations, row rotations can be more efficiently applied by partitioning the columns into stripes and applying a sequence fully to each stripe in turn. In this way, any matrix entries that are brought into the cache by the application of one rotation is likely to still be present in the cache by the time the next rotation is applied. Such a column striping technique not only improves the cache reuse but also reduces the number of TLB misses [9].

4.2. The cache-blocked algorithm. The structure-exploiting algorithm has better cache reuse than the basic algorithm, yet it still applies only a constant number of floating point operations (approximately three to six) per accessed matrix entry and iteration of the outer loop. Since each iteration accesses most of the matrices, it follows that the arithmetic intensity$^1$ will be small for sufficiently large matrices. Significant improvements of the cache reuse therefore necessarily require the reordering of operations spanning several iterations of the outer loop. The main difficulty is to handle the large number of data dependencies that are caused by the intrinsic interleaving of row and column rotations.

Fortunately, the accumulation of rotations into $Q$ and $Z$ is performed exclusively using column rotations and can be effectively reorganized for improved cache reuse. However, many of the updates of $A$ and some of the updates of $B$ involve interleaved row and column updates that cannot be easily separated. Some improvements of the cache reuse are nonetheless possible.

Related works such as [9, 11, 12] describe how to apply a blocking technique to the accumulation of $Q$ and $Z$ as well as to those parts of $A$ and $B$ that are affected only by column rotations. Recent work by one of the authors demonstrates how to

---

$^1$With arithmetic intensity we mean the ratio of the number of floating point operations performed to the number of bytes transferred to and from memory.
Algorithm 3: $A, B, Q, Z \leftarrow$ HTBlocked$(n, d, A, B, Q, Z)$.

1. $\hat{c} \leftarrow 1$;
2. while $\hat{c} \leq dn - 2$ do
3.   Choose any $\hat{c} \in \{\hat{c}, \ldots, dn - 2\}$;
4.     /* For each iteration $\hat{c}$ in the current panel. */
5.     for $c \leftarrow \hat{c}$ to $\hat{c}$ do
6.        /* Phase I: Panel factorization */
7.        $r_1, r_2 \leftarrow c + 1, \min \{dn, c + n\}$;
8.        $G^Q_c \leftarrow$ LeftZero$(A, r_1, r_2, c)$;
9.        $G^Z_c \leftarrow$ RightZero$(G^Q_c, B, r_1, r_2, c_1, c_2)$;
10.       $G \leftarrow G^Z_c$;
11.      /* Chase bulges until they disappear. */
12.     for $k \leftarrow 1, 2, \ldots$ do
13.       /* Move from $B$ to $A$. */
14.       $r_1, r_2 \leftarrow c_1 + n, \min \{dn, c_2 + n\}$;
15.       $G \leftarrow$ RightLeftZero$(G, A, r_1, r_2, c_1, c_2)$;
16.      /* Exit if all bulges have disappeared. */
17.     if $r_1 > dn$ then exit loop;
18.     /* Generate and zero out bulges in $A$. */
19.    $G^{Q/Z}_{c,k} \leftarrow$ RightLeftZero$(G, A, r_1, r_2, c_1, c_2)$;
20.   $G \leftarrow G^{Q/Z}_{c,k}$;
21. /* Move from $A$ to $B$. */
22.   $c_1, c_2 \leftarrow r_1, r_2$;
23.  $k_c \leftarrow k - 1$;
24. /* Phase II: Delayed updates */
25. Accumulate $G^Q_c$ for $c \in \{\hat{c}, \ldots, \hat{c}\}$ into matrices $U^Q_i$ for $i \in \{1, \ldots, n_Q\}$;
26. Accumulate $G^Z_c$ for $c \in \{\hat{c}, \ldots, \hat{c}\}$ into matrices $U^Z_i$ for $i \in \{1, \ldots, n_z\}$;
27. Accumulate $G^{Q/Z}_{c,k}$ for $c \in \{\hat{c}, \ldots, \hat{c}\}$ into matrices $U^{Q/Z}_{i}$ for $i \in \{1, \ldots, n_{Q/Z}\}$;
28. Apply $U^{Q/Z}_{i}$ to $Q$ for all $i$;
29. Copy columns $\hat{c} + n$: $\min \{dn, \hat{c} + n\}$ from $Q$ to $Z$;
30. Apply $U^Q_i$ to $Q$ and $U^Z_i$ to $Z$ for all $i$;
31. Apply $U^{Q/Z}_{i}$ followed by $U^{Z}_{i}$ to rows $1: \hat{c}$ of $A$ for all $i$ and $i'$;
32. Apply $U^{Z}_{i}$ to rows $1: \hat{c}$ of $B$ for all $i$;
33. /* Advance to the next panel. */
34. $\hat{c} \leftarrow \hat{c} + 1$;
block also the updates of $A$ (and $B$) [10]. For the latter technique to be effective, however, the lower bandwidth should be much smaller than the size of the matrices, which is typically not the case here since the degree tends to be small relative to $n$ and hence the bandwidth, $n$, is large relative to the size, $dn$, of the matrices.

Let us now isolate the parts of the matrices which are updated only by column rotations from those parts which are updated by both row and column rotations. This will help us understand the blocked algorithm that follows. As mentioned above, all the accumulations into $Q$ and $Z$ are performed using column rotations. The processing of the $(2, 2)_c^B$ block of $B$, via the kernel LeftRightZero on line 8 (of Algorithm 2), involves row and column rotations. The remaining updates of $B$ are all based on column rotations. The processing of the blocks $(i, j)^A_c$ for $i = 2, 3, \ldots, N_c$ with $N_c$ defined in (3.1) and $j = i, \ldots, \min\{N_c, d\}$ involves both row and column rotations. The remaining updates of $A$, that is, to the blocks $(1, j)^A_c$ for $j = 2, 3, \ldots, N_c$, are all based on column rotations. In summary, only the submatrices $A(c + 1: dn, c + 1: dn)$ and $B(c + 1: c + n, c + 1: c + n)$ involve both row and column rotations; the rest involve only column rotations.

The cache-blocked Algorithm 3 improves on the structure-exploiting algorithm from section 3.7 in part by introducing blocking and in part by avoiding a redundancy that occurs in the accumulation of $Q$ and $Z$, the latter of which is explained in section 3.6. The blocked algorithm zeroes out columns of $A$ from left to right one column panel at a time. Each iteration of the outer loop on line 2 reduces one panel. Each block iteration consists of two phases. In Phase I (“panel factorization”), the columns $\hat{c}: \hat{c}$ are zeroed out, and the resulting bulges are chased using what amounts to the pipelined algorithm. This is accomplished by the outer for-loop on line 4. Only the submatrices $A(\hat{c} + 1: dn, \hat{c} + 1: dn)$ and $B(\hat{c} + 1: c + n, \hat{c} + 1: c + n)$ are updated in Phase I since the other updates are not required to construct the rotations. In Phase II (“delayed updates”), the rotations are regrouped and accumulated into small unitary matrices, which are then applied using matrix multiplication to $Q$, $Z$, $A(1: \hat{c}, \hat{c} + 1: dn)$, and $B(1: \hat{c}, \hat{c} + 1: \hat{c} + n)$.

4.3. Regrouping rotations for increased cache reuse. We increase the cache reuse in Phase II by adopting a known technique for applying multiple overlapping sequences of rotations [12]. This technique exploits the commutativity of rotations acting on disjoint sets of rows or columns and rearranges the rotations into diamond-shaped groups, each of which can be applied cache-efficiently.

However, we cannot apply this technique directly since the rotations obtained from one bulge-chasing sweep form not one but several nonoverlapping sequences. Fortunately, these shorter sequences can be merged by inserting identity rotations to artificially create one long sequence.

Consider the first three iterations of the outer for-loop over $c$ for the case $d = 4$, $n = 5$. Let $G_k^c$ denote the row rotation constructed in iteration $c$ to annihilate $a_{k,c}$ using $a_{k-1,c}$ as a pivot. This rotation acts on rows $k-1$ and $k$. Let $I_k$ denote an identity rotation acting on rows $k-1$ and $k$. Figure 10(a) illustrates the row rotations and their dependencies for the first three column reductions. Each vertex represents a rotation, and each arc represents a dependence caused by noncommutativity. Any topological ordering of the graph corresponds to a valid reordering of the rotations (and vice versa). The region with a dashed outline is an example of how to group rotations into a diamond-like shape. Such a group has the property that there is no directed path leaving and re-entering the region. This implies that the rotations in the group can be applied in sequence without any other rotation in between. In
particular, the rotations in a group can be explicitly accumulated into a unitary matrix and applied using matrix multiplication.

The graph in Figure 10(a) has “gaps” (e.g., rotation $G^3$ does not exist). In Figure 10(b), the gaps have been filled with identity rotations that have no effect other than to make the structure of the graph more regular. The graph in (b) is a restriction of the graph in (a) in the sense that any topological ordering of (b) contains a topological ordering of (a). The graph structure in Figure 10(b) is identical to the one in [12], and hence the technique described therein applies directly.

Another viewpoint is illustrated in Figure 11. The bottom part of the figure symbolizes a matrix, either $Q$ or $Z$, on top of which all the rotations are shown as rotated square brackets. Rotations closer to the matrix in Figure 11 should be applied first, unless they do not overlap, in which case they commute and can be applied in
any order. The figure also illustrates which of the rotations are shared by $Q$ and $Z$ (lightface brackets), and which are unique to $Q$ and $Z$ (boldface (red online) brackets). Hence, a rotation shown in boldface (red online) actually represents two rotations, one acting on $Q$ and the other on $Z$. The insertion of identity rotations and the regrouping of rotations are illustrated in Figure 12.

5. The parallel algorithm. Even though our proposed blocked algorithm requires less computation and communication through the memory hierarchy than the conventional algorithm, it is still quite costly in terms of the number of flops with an arithmetic complexity of $O(d^3n^3)$. We have therefore constructed a parallel algorithm for multicore-based systems implemented using threads.

The parallel algorithm is structured similarly to Algorithm 3 and consists of iterations over panels. Each iteration consists of two phases, each of which is executed in parallel. Successive phases are separated with a barrier synchronization. The parallelization of Phase II is described in section 5.1, and the (more technical) parallelization of Phase I is described in section 5.2.

5.1. Parallel formulation of Phase II. The parallel algorithm for Phase II consists of an accumulation step followed by an update step with a barrier synchronization in between. In the accumulation step, the rotations are regrouped, and each rotation group is accumulated into a small unitary matrix. The accumulations of different groups are independent, and the accumulation step is therefore perfectly parallel.

In the update step, the groups containing shared rotations are first applied to $Q$. This is then followed by the copying of a set of columns from $Q$ to $Z$ as part of the elimination of redundancy. The update step ends by applying the rotation groups that contain nonshared rotations to $Q$ and $Z$. Also the update step is perfectly parallel.
in the sense that the rows can be independently updated. However, the load is not uniformly distributed over the rows due to the sparsity structure of \( Q \) and \( Z \). We address this problem by calculating ahead of time the number of flops per row and partition the rows of \( Q \) and \( Z \) such that the flops are approximately balanced over the threads.

5.2. Parallel formulation of Phase I. The parallel algorithm for Phase I consists of multiple iterations, each of which interleaves row and column rotations with the zeroing out of matrix entries.

The steps in each iteration of the loop in Phase I (see Algorithm 3) can be rewritten in terms of coarse operations acting on the blocks of the natural block partitioning. In doing so we also take the opportunity to further increase the cache reuse by exploiting the fact that all the blocks in the upper triangle are subject to both a row sequence and a column sequence. By fusing the application of these two sequences into a single kernel we can reduce its communication volume by up to 50%.

The new kernel is defined as follows:

- **RightLeft**\((G, \tilde{G}, A, r_1, r_2, c_1, c_2)\).

  Applies a decreasing sequence of row rotations \( G \) and a decreasing sequence of column rotations \( \tilde{G} \) to the submatrix \( A(r_1 : r_2, c_1 : c_2) \). The number of row rotations is \( r_2 - r_1 \), and the number of column rotations is \( c_2 - c_1 \).

The new version of Phase I is given in Algorithm 4.

**Algorithm 4**: Phase I expressed using operations on blocks.

\[
\begin{align*}
1 & \text{for } c \leftarrow \hat{c} \text{ to } \check{c} \text{ do} \\
2 & \quad \text{LeftZero on } (2,1)^A_c; \\
3 & \quad \text{LeftRightZero on } (2,2)^B_c; \\
4 & \quad \text{for } i \leftarrow 3 \text{ to } N_c \text{ do} \\
5 & \quad \quad \text{RightLeftZero on } (i,i-1)^A_c; \\
6 & \quad \quad \text{for } j \leftarrow 3 \text{ to } N_c \text{ do} \\
7 & \quad \quad \quad \text{RightLeft on } (2,j)^A_c; \\
8 & \quad \quad \text{for } i \leftarrow 3 \text{ to } N_c \text{ do} \\
9 & \quad \quad \quad \quad \text{for } j \leftarrow i \text{ to } \min\{d,N_c\} \text{ do} \\
10 & \quad \quad \quad \quad \quad \text{RightLeft on } (i,j)^A_c;
\end{align*}
\]

With each operation in Algorithm 4 executed by a single thread, the number of tasks would be limited to \( O(d^2) \), and the average degree of concurrency would be only \( O(d) \). Therefore, a large degree would be necessary to saturate a typical multicore-based system. Since the degrees are predominantly small in practice, we further decompose each operation into finer grained tasks. This in turn requires an additional set of computational kernels, which we define in section 5.2.1. The task decomposition, dependencies, and scheduling of the tasks are described in section 5.2.2.

5.2.1. Task decomposition of Phase I. The task decomposition that we use is to a great extent induced by a partitioning of each block into square subblocks defined by a subblock size \( b \) and aligned with the top left corner of the block. This in turn induces a partitioning of the sequences of rotations into subsequences of length \( b \).

The reason for square subblocks is to ensure that a subblock of the row rotations that become column rotations align with the subblock partitioning of the columns.

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Fig. 13. Computational kernels employed in the parallel implementation of Phase I. Each subfigure illustrates one subblock and its immediate surroundings indicated by thick lines. Empty cells should be interpreted as zeros and the others as nonzeros. The colored (online) regions indicate which entries are updated by a sequence of row rotations (blue online), respectively, column rotations (red online). The arrows indicate the direction of and the order in which the sequences are applied.

The \textit{LeftZero} operation involves little work and is not decomposed into tasks in order to avoid the associated scheduling overhead. (Moreover, the operation is inherently sequential.) Instead it is merged with the \textit{LeftRightZero} operation that follows.

The \textit{RightLeft} operation takes two sequences as input: one row sequence and one column sequence. Figure 13(c) illustrates the kernel that forms the basis of the implementation. Shown is one of the subblocks and its immediate surroundings. The kernel applies to the subblock first the column sequence and then the row sequence. Note that the column (row) rotations affect one additional column (row) to the east (south). The region affected by the row rotations has also been shifted one column to the east since the leftmost column in the subblock is not fully up-to-date with respect to the column rotations. Special cases occur at the south and east boundaries of the block since the updates do not extend beyond the block. The task graph and the relationships between the tasks and the subblocks are illustrated in Figure 14(a). With $n_b = \lceil n/b \rceil$ subblocks in either dimension, the task graph contains $n_b^2$ tasks and has depth $2n_b - 1$.

The \textit{LeftRightZero} operation is built upon the two computational kernels illustrated in Figures 13(a)–(b), and its task graph is illustrated in Figure 14(b). The \textit{SubLeftRightZero} kernel applies to the subblocks on the diagonal, and the \textit{SubLeftRight} kernel applies to the subblocks above the diagonal. Again, the regions affected by row, respectively, column, rotations are not perfectly aligned to the sub-
blocks, as is evident from Figure 13. With \( n_b \) subblocks in either dimension, the task graph contains \( n_b(n_b + 1)/2 \) tasks and has depth \( 2n_b - 1 \).

The RightLeftZero operation is built upon the three computational kernels illustrated in Figures 13(d)–(f). Interestingly, even though the name and functionality of this operation are similar to LeftRightZero, they are not symmetric, and as a consequence their task decompositions are rather different. The task graph of RightLeftZero is illustrated in Figure 14(c), and the tasks on the diagonal use the kernel SubRightLeftZero. Each subblock in the strictly upper triangular part contains one SubRight task (closer to the bottom right corner) and one SubLeft task (closer to the upper left corner). With \( n_b \) subblocks in either dimension, the task graph contains \( n_b^2 \) tasks and has depth \( n_b + 1 \).

5.2.2. Scheduling Phase I to promote cache reuse. In the scheduling of Phase I we also attempt to increase the cache reuse. Specifically, the parallel implementation of Phase I begins by explicitly constructing the task graph for the first iteration. The tasks are then statically scheduled using the critical path heuristic (largest height first). We then reuse the same schedule for all subsequent iterations in the phase. One consequence of this approach is that each thread is statically assigned a subset of the tasks and by extension also a subset of the subblocks. This is what increases the cache reuse.

Figure 15 illustrates the task graph associated with one iteration of the first instance of Phase I for a polynomial of degree four. The subblock size is set to one-fourth of the block size. The labels on the nodes indicate the kernel used and are the same as the labels used in Figure 13. The graph contains 170 tasks and has depth 14, which yields a 12.143 average degree of concurrency. This should be contrasted to the case of doing no decomposition of the blocks at all (i.e., setting the subblock size equal to the block size). In that case the graph would have only 11 tasks and a depth of 5 for a 2.2 average degree of concurrency. (Since consecutive iterations partially overlap in practice, the average degree of concurrency is actually slightly larger in both cases.)

6. Experiments. We developed two separate implementations for different types of systems: a sequential implementation for workstations following Algorithm 3 and a thread-based implementation based on the parallel algorithm described in section 5. The sequential algorithm is intended for use on workstations via a MATLAB MEX interface and takes advantage of potential implicit parallelism by calling the MATLAB BLAS library. The parallel implementation is intended for use on multicore-based high-performance computing (HPC) systems.
Fig. 15. Task graph for one iteration of Phase I for a polynomial of degree four and with four subblocks in either dimension of each block. The node colors (online) and labels indicate the kernel used by the task and correspond to the subfigure labels in Figure 13.

Table 3

Comparison between the sequential mex code, the MATLAB hess function, and the parallel implementation on the workstation.

<table>
<thead>
<tr>
<th>n</th>
<th>d</th>
<th>( \text{MATLAB} )</th>
<th>( \text{Parallel} )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( p = 1 )</td>
<td>( p = 2 )</td>
<td>( p = 3 )</td>
</tr>
<tr>
<td>500</td>
<td>2</td>
<td>1.2</td>
<td>5.7</td>
</tr>
<tr>
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</tr>
<tr>
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<td>4</td>
<td>8.2</td>
<td>55</td>
</tr>
<tr>
<td>1000</td>
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<td>14</td>
<td>56</td>
</tr>
<tr>
<td>1000</td>
<td>3</td>
<td>40</td>
<td>192</td>
</tr>
<tr>
<td>1000</td>
<td>4</td>
<td>85</td>
<td>506</td>
</tr>
</tbody>
</table>

We measured the performance of our codes on two systems (one workstation and one HPC system) with different characteristics.

- **Workstation.** The workstation has an Intel Core i7-2600 processor with four cores and a nominal clock frequency of 3.4 GHz.
- **HPC system.** The HPC system is a cluster consisting of interconnected shared-memory nodes. Each node consists of four AMD Opteron 6238 processors each with 12 cores split into two NUMA domains. The nominal clock frequency is 2.6 GHz.

Both systems run a Linux-based operating system. All experiments were performed in double precision (64-bit) real floating point arithmetic. All execution times are given in seconds.

### 6.1. Results from the workstation.

On the workstation we measured the execution time of our mex code with implicit parallelism via the parallel BLAS used in MATLAB and compared it against the HT reduction function hess in MATLAB, our parallel code with \( p \in \{1, 2, 3, 4\} \) threads, and a prototype implementation, DGCHR3, of the blocked one-stage HT reduction algorithm described in [9]. The results were obtained using MATLAB version R2011b for Linux.

The results are summarized in Table 3 for \( n \in \{500, 1000\} \) and \( d \in \{2, 3, 4\} \). The times are given in seconds. The column labeled hess/mex shows the ratio of the execution time of the hess function to the execution time of our mex code. The
ALGORITHMS FOR HESSENBERRY-TRIANGULAR REDUCTION

Table 4

Execution time of the prototype implementation DGGHR3 of the blocked one-stage HT reduction algorithm described in [9] linked with multithreaded BLAS on the workstation.

<table>
<thead>
<tr>
<th>n</th>
<th>d</th>
<th>p = 1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>500</td>
<td>2</td>
<td>2.3</td>
<td>1.9</td>
<td>1.8</td>
<td>1.8</td>
</tr>
<tr>
<td>500</td>
<td>3</td>
<td>7.3</td>
<td>6.0</td>
<td>5.7</td>
<td>5.6</td>
</tr>
<tr>
<td>500</td>
<td>4</td>
<td>17</td>
<td>14</td>
<td>13</td>
<td>13</td>
</tr>
<tr>
<td>1000</td>
<td>2</td>
<td>17</td>
<td>14</td>
<td>13</td>
<td>13</td>
</tr>
<tr>
<td>1000</td>
<td>3</td>
<td>56</td>
<td>46</td>
<td>43</td>
<td>41</td>
</tr>
<tr>
<td>1000</td>
<td>4</td>
<td>133</td>
<td>107</td>
<td>99</td>
<td>97</td>
</tr>
</tbody>
</table>

Table 5

The execution time of the LAPACK generic HT reduction routine DGGHRD following a custom structure-exploiting reduction of B to upper triangular form on the HPC system.

<table>
<thead>
<tr>
<th>n</th>
<th>500</th>
<th>1000</th>
<th>1500</th>
<th>2000</th>
</tr>
</thead>
<tbody>
<tr>
<td>d</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>d</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>d</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>d</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>d</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>d</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>d</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>d</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>

The size of the improvement depends on both n and d. The observed improvements of the mex code range from 4.0 (for n = 1000, d = 2) to 6.8 (for n = 500, d = 4). The observed improvements of the parallel code with four threads range from 6.3 (for n = 500, d = 2) to 12.3 (for n = 1000, d = 4). As expected, the increasing sparsity of the linearization for large degrees leads to increased improvements over the generic hess function. A larger size of the coefficient matrices does seem to slightly decrease the improvement of the mex code while, on the other hand, increasing slightly the improvement of the parallel code.

We obtained a prototype implementation, DGGHR3, of the blocked one-stage algorithm presented in [9], which represents the state of the art for generic HT reduction. The execution times for a varying number of threads are shown in Table 4, which can be directly compared column by column to Table 3 (parallel results). DGGHR3 significantly outperforms the hess function in MATLAB and performs only slightly worse than our structured code. The ratios of the execution times of DGGHR3 to our parallel code range from 0.9 (n = 1000, d = 2, p = 1) to 2.8 (n = 500, d = 4, p = 4) with a mean of 1.8.

6.2. Results from the HPC system. On the HPC system we measured the performance of our parallel code and compared it against the LAPACK DGGHRD generic HT reduction routine as well as DGGHR3. The matrix B must be upper triangular on input, so we performed a structure-exploiting preprocessing step that reduces the top left n × n dense block of B to upper triangular form before calling DGGHRD or DGGHR3.

6.2.1. Total execution time. The DGGHRD routine in LAPACK is based on Level 1 BLAS operations without explicit parallelism and thereby does not scale on multicore-based systems. Table 5 summarizes the total execution times for LAPACK linked with OpenBLAS 0.2.8 (sequential) for n ∈ {500, 1000, 1500, 2000} and d ∈ {2, 3, 4}.

These measurements should be compared to our parallel code solving problems of the same size with 1 ≤ p ≤ 24 threads. The time measurements include both the preprocessing step and the bulge-chasing part of the reduction (only the latter of which is parallel) and are summarized in Table 6. The row with p = 1 can be directly...
The parallel execution time (including both the preprocessing step and the bulge-chasing part) of our polynomial HT reduction algorithm on the HPC system.

<table>
<thead>
<tr>
<th>$n$</th>
<th>500</th>
<th>1000</th>
<th>1500</th>
<th>2000</th>
</tr>
</thead>
<tbody>
<tr>
<td>$p$</td>
<td>54</td>
<td>53</td>
<td>33</td>
<td>32</td>
</tr>
<tr>
<td>$d$</td>
<td>22</td>
<td>22</td>
<td>32</td>
<td>32</td>
</tr>
<tr>
<td></td>
<td>43</td>
<td>74</td>
<td>56</td>
<td>97</td>
</tr>
</tbody>
</table>

The parallel execution time on the HPC system of the DGGHR3 prototype implementation of the blocked one-stage algorithm described in [9] (including both the preprocessing step and the actual HT reduction).

<table>
<thead>
<tr>
<th>$n$</th>
<th>500</th>
<th>1000</th>
<th>1500</th>
<th>2000</th>
</tr>
</thead>
<tbody>
<tr>
<td>$p$</td>
<td>35</td>
<td>12</td>
<td>26</td>
<td>35</td>
</tr>
<tr>
<td>$d$</td>
<td>36</td>
<td>12</td>
<td>26</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>27</td>
<td>86</td>
<td>92</td>
<td>12</td>
</tr>
</tbody>
</table>

The largest observed improvement of the parallel code over DGGHRD is 28 (for $n = 1000, d = 4, p = 16$). The largest relative speedup of the parallel code (i.e., relative improvement over $p = 1$) ranges from 2.4 ($n = 500, d = 2, p = 12$) to 6.7 ($n = 2000, d = 4, p = 16$).

Table 7 shows the parallel execution time of the DGGHR3 prototype implementation with up to 12 threads (one full processor) in the multithreaded version of the BLAS library. In all cases except for small values of $n$, the code reaches its scalability limit. The code scales modestly, which is expected considering that the code is only implicitly parallelized via the BLAS.

Comparing the minimum execution time of DGGHR3 with the minimum execution time of our parallel code, we find that the relative improvement of our code ranges from 3 to 4 for $n = 2000, d = 4, p = 12$. The largest relative speedup of the parallel code (i.e., relative improvement over $p = 1$) ranges from 2.4 ($n = 500, d = 2, p = 12$) to 6.7 ($n = 2000, d = 4, p = 16$).
6.2.2. Analysis of the parallel bulge-chasing algorithm. The parallel bulge-chasing (section 5) is the most time consuming part of the reduction. In this section, we analyze the behavior of the internals of the parallel implementation.

The code was instrumented to (approximately) count the number of floating point operations performed in each phase of each panel iteration. These counts are plotted against the panel iteration number in Figure 16 for the case $n = 2000, d = 4$. (Each panel consists of 32 columns for a total of $nd/32 = 250$ iterations.)

Despite most of the flops being accounted for by Phase II, it is typically Phase I that consumes the majority of the execution time due to its low arithmetic intensity and greater complexity. Figure 17 shows the performance in Gflops, i.e., the ratio of the floating point operation counts from Figure 16 and the measured execution times for a varying number of threads. The graphs on the left side correspond to Phase I, and the graphs on the right correspond to Phase II. It is observed from the latter that

---

**Fig. 16.** Distribution of flops over the two phases as a function of the iteration number ($n = 2000, d = 4$).

**Fig. 17.** Performance of Phase I (left) and Phase II (right) for $n = 2000, d = 4$, and a varying number of threads on the HPC system.

from $2.6 (n = 2000, d = 2, p = 8$ threads in DGGHR3 and $p = 18$ threads in our parallel code) to $5.1 (n = 500, d = 4, p = 4$ threads in DGGHR3 and $p = 12$ threads in our parallel code).
Phase II scales up to \( p = 18 \). The graphs for Phase I also indicate good scaling up to \( p = 18 \), but there is also a sharp decrease in the performance toward the second half of the iterations.

We investigated the source of the performance degradations observed in Phase I by measuring the per-thread time spent in computational kernels per instance of Phase I. More specifically, consider any instance of Phase I, and let \( T_k \) denote the time that thread \( k \) spent computing in that instance. Then the total cost of computation is given by \( \sum_{k=1}^{p} T_k \). If we assume that the work could have been perfectly distributed over the \( p \) threads, then \( \frac{1}{p} \sum_{k=1}^{p} T_k \) is a lower bound for the parallel execution time. The most heavily loaded thread computes for a total of \( \max_k T_k \) time units. It follows that the ratio

\[
L = \frac{\sum_{k} T_k}{p \cdot \max_k T_k}
\]

in the range \((0, 1]\) is a measure of the load imbalance with \( L = 1 \) indicating perfect load balance. In general, the most heavily loaded thread has \( 1/L \) times the optimal load. The amount of load imbalance for the case \( n = 2000, d = 4 \), and a varying number of threads is shown in Figure 18. From the strong correlation between the increasing load imbalance and the decreasing performance of Phase I shown in Figure 17 (left), we conclude that the load imbalance is a primary explaining factor and runtime autotuning of the subblock size might improve the performance further.

7. Conclusion and future work. We have identified a special Fiedler linearization of matrix polynomials of arbitrary degree, which shares many of the properties of standard companion linearizations but also allows (a) the deflation of some of the zero and infinite eigenvalues without affecting the block structure of the linearization, and (b) the exploitation of the sparsity structure in the HT reduction. We have presented a backward stable high-performance parallel algorithm for the HT reduction of such linearization. Our algorithm exploits the sparsity structure of the linearization and the redundancy in the computation of the transformation matrices, and thereby requires fewer flops than existing algorithms for unstructured pairs of matrices, including the state-of-the-art HT reduction in [9]. Special care has been taken to improve cache reuse. Experiments on both a workstation and an HPC system have demonstrated
that our structure-exploiting parallel implementation can outperform both the general LAPACK routine \texttt{DGGHRD} and the prototype implementation \texttt{DGGHR3} of a general blocked algorithm.

Future work includes runtime autotuning of the subblock size to better balance the load, alternative parallelization schemes that address the trade-off between cache reuse and concurrency differently, and a hybrid approach that switches to a (parallel variant of) \texttt{DGGHR3} during the last iterations where the structure is less pronounced.

\section*{Appendix A. Deflation of infinite and zero eigenvalues on special Fiedler linearization.}

If either one or both of \( P_0 \) and \( P_d \) are rank deficient, then it is possible to deflate structurally induced zero and infinite eigenvalues prior to the HT reduction. This is desirable since these eigenvalues might otherwise go partially undetected due to round-off errors accumulated during the QZ algorithm. We show in this appendix how to transform the \( dn \times dn \) Fiedler pencil \( L(\lambda) \) in (1.2) into block upper triangular form

\begin{equation}
A + \lambda B = 
\begin{bmatrix}
A_{11} & A_{12} & A_{13} \\
0 & A_{22} & A_{23} \\
0 & 0 & 0_{n-r_0}
\end{bmatrix} + \lambda 
\begin{bmatrix}
0_{n-r_2} & B_{12} & B_{13} \\
0 & B_{22} & B_{23} \\
0 & 0 & 0_{3n}
\end{bmatrix},
\end{equation}

(A.1)

where the \( dn \times dn \) matrices \( A \) and \( B \) are partitioned conformably. Here \( r_0 \) denotes the rank of \( P_0 \), and \( r_d \) denotes the rank of \( P_d \). Note that if \( A_{11} \) (or \( B_{33} \)) is singular, then \( \det P(\lambda) = \gamma \det L(\lambda) \equiv 0 \) and hence \( P(\lambda) \) is singular. When \( A_{11} \) and \( B_{33} \) are nonsingular, (A.1) reveals \( n-r_0 \) zero eigenvalues and \( n-r_d \) infinite eigenvalues. The remaining eigenvalues are those of the \( m \times m \) pencil \( A_{22} + \lambda B_{22} \), where \( m = (d-2)n + r - 1 + r_d \). Now to use our HT reduction algorithm, \( A_{22} + \lambda B_{22} \) must retain most of the sparsity structure of the Fiedler linearization (1.2). We show that our deflation yields a deflated pencil \( A_{22} + \lambda B_{22} \) of the form (when \( d = 4 \))

\begin{equation}
\begin{bmatrix}
X_{n, n} & X_{n, r_0} & X_{n, r_0} & X_{n, r_0} \\
-I_{r_0} & 0 & 0 & 0 \\
-I_{r_0} & 0 & 0 & 0 \\
0 & 0 & 0 & 0
\end{bmatrix} + \lambda 
\begin{bmatrix}
X_{n, n} & 0 & 0 & 0 \\
0 & I_{r_0} & 0 & 0 \\
0 & 0 & I_{r_0} & 0 \\
0 & 0 & 0 & I_{r_0}
\end{bmatrix},
\end{equation}

(A.2)

where \( X_{m, n} \) denotes a dense matrix of size \( m \times n \) and \( I_n \) is an identity matrix of size \( n \times n \).

We illustrate the proposed deflation procedure for the case \( d = 4 \), which is large enough to clearly exhibit the general pattern. The procedure can be readily generalized to any degree \( d \geq 3 \). However, the extreme case \( d = 2 \) requires a little bit of special treatment, which we defer to the end of this appendix.

The appendix is outlined as follows. Fiedler linearizations and our special choice are defined in section A.1. The deflation procedure consists of two steps:

1. deflation of zero eigenvalues induced by \( P_0 \) described in section A.2,
2. deflation of infinite eigenvalues induced by \( P_d \) described in section A.3.

The special treatment of the case \( d = 2 \) is described in section A.4.

\subsection*{A.1. Fiedler linearizations.}

With the notation

\begin{align*}
M_0 &= \text{diag}(I_{n(d-1)}, -P_0), & M_d &= \text{diag}(P_d, I_{(d-1)n}), \\
M_j &= \text{diag} \left( I_{n(d-j-1)}, \begin{bmatrix} -P_j & I_n \\ I_n & 0 \end{bmatrix}, I_{n(j-1)} \right), & j &= 1 : d - 1,
\end{align*}
and any permutation \( \sigma = (i_0, i_1, \ldots, i_{d-1}) \) of the indices \((0, 1, \ldots, d-1)\), we define the associated Fiedler pencil by [1]

\[
L_\sigma(\lambda) = -M_{i_0}M_{i_1} \cdots M_{i_{d-1}} + \lambda M_d.
\]

The pencils \( L_\sigma(\lambda) \) are all linearizations of \( P(\lambda) \) [1]. The Fiedler pencil we are interested in corresponds to the permutation \( \sigma = (d-1, \ldots, 3, 2, 0, 1) \), that is,

\[
L_\sigma(\lambda) = -M_{d-1} \cdots M_2 M_1 M_0 + \lambda M_d
\]

if \( d > 2 \)

\[
\begin{bmatrix}
P_{d-1} & P_{d-2} & \cdots & P_1 & -I_n \\
-I_n & \ddots & \ddots & \ddots & \ddots \\
& \ddots & \ddots & \ddots & \ddots \\
& & -I_n & P_0 & \\
& & & P_0 &
\end{bmatrix}
+ \lambda
\begin{bmatrix}
P_d \\
I \\
\vdots \\
I
\end{bmatrix}
\cdot
\]

(A.3)

A.2. Deflation of zero eigenvalues induced by \( P_0 \). If \( P_0 \) has full rank, then we have nothing to do in this step. Therefore, assume that \( P_0 \) has rank \( r_0 < n \) and compress its rows upward by premultiplication with a carefully crafted unitary matrix \( U^* \); i.e., choose a unitary matrix \( U \) such that

\[
U^* P_0 = \begin{bmatrix}
\tilde{P}_0 \\
0
\end{bmatrix},
\]

where \( \tilde{P}_0 \) has \( r_0 \) rows and full rank. (This can be accomplished, for instance, by using a singular value decomposition of \( P_0 \) or a QR factorization with column pivoting.) Transform \((A, B)\), where \( A + \lambda B \) is the Fiedler linearization described in (A.3), by premultiplying the last block row with \( U^* \). This changes the bottom right block of \( B \) from \( I \) to \( U^* \). Restore \( B \) by postmultiplying the last block column with \( U \). This also changes the top right block of \( A \) from \( -I \) to \( -U \), but this we accept. The pair \((A, B)\) now takes the form

\[
A = \begin{bmatrix}
P_1 & P_2 & P_3 & -U_1 & -U_2 \\
-I & 0 & 0 & 0 & 0 \\
0 & -I & 0 & 0 & 0 \\
0 & 0 & \tilde{P}_0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0
\end{bmatrix}, \quad B = \begin{bmatrix}
P_1 & 0 & 0 & 0 & 0 \\
0 & I & 0 & 0 & 0 \\
0 & 0 & I & 0 & 0 \\
0 & 0 & 0 & I & 0 \\
0 & 0 & 0 & 0 & I
\end{bmatrix},
\]

where the last block row and column have \( n - r_0 \) rows and columns, respectively. This effectively deflates \( n - r_0 \) zero eigenvalues at the bottom right corner. After removing the last \( n - r_0 \) rows and columns (boldface above), we obtain a reduced problem that takes the form

\[
\begin{bmatrix}
P_3 & P_2 & P_1 & (-U_1) \\
-I & 0 & 0 & (0) \\
0 & -I & 0 & (0) \\
(0) & (0) & \tilde{P}_0 & 0
\end{bmatrix}, \quad B = \begin{bmatrix}
P_1 & 0 & 0 & (0) \\
0 & I & 0 & (0) \\
0 & 0 & I & (0) \\
(0) & (0) & (0) & I
\end{bmatrix},
\]

where each lightface block has size \( n \times n \), the boldface block has size \( r_0 \times r_0 \), and each boldface block with parentheses has size \( n \times r_0 \) or \( r_0 \times n \). This completes our description of the first step of the deflation procedure.
A.3. Deflation of infinite eigenvalues induced by $P_d$. The input to this step takes the form (A.4) for $d = 4$, and the aim is to deflate infinite eigenvalues induced by a rank deficiency in $P_d$. If $P_d$ has full rank, then there is nothing to do. Therefore, assume that $P_d$ has rank $r_d < n$ and compress the columns of $P_d$ toward the right by postmultiplication with a carefully crafted unitary matrix $\hat{U}$; i.e., choose a unitary matrix $\hat{U}$ such that

$$P_d \hat{U} = \begin{bmatrix} 0 & \hat{P}_d \end{bmatrix},$$

where $\hat{P}_d$ has $r_d$ columns and full rank. Transform $(A, B)$ by postmultiplying the first block column with $\hat{U}$. This changes the $(2, 1)$ block of $A$ from $-I$ to $-\hat{U}$. Restore the structure of $A$ by premultiplying the second block row with $\hat{U}^*$. This simultaneously changes the $(2, 2)$ block of $B$ from $I$ to $\hat{U}^*$. Continue restoring identities until finally the $(d, d-1)$ block of $A$ changes from $\hat{P}_0$ to $\hat{P}_0 \hat{U}$ and the process ends. The transformed pair $(A, B)$ now takes the form

$$A = \begin{bmatrix} \hat{P}_{3,1} & \hat{P}_{3,3} & \hat{P}_{2,1} & \hat{P}_{2,3} & \hat{P}_{1,1} & \hat{P}_{1,3} & -U_{1,1} & -U_{1,2} \\ -I & 0 & 0 & 0 & 0 & 0 \\ 0 & -I & 0 & 0 & 0 & 0 \\ 0 & 0 & -I & 0 & 0 & 0 \\ 0 & 0 & 0 & -I & 0 & 0 \\ 0 & 0 & 0 & 0 & \hat{P}_{0,1} & \hat{P}_{0,2} & 0 \end{bmatrix},$$

$$B = \begin{bmatrix} 0 & \hat{P}_{4,1} & 0 & 0 & 0 & 0 \\ 0 & \hat{P}_{4,2} & 0 & 0 & 0 & 0 \\ 0 & 0 & I & 0 & 0 & 0 \\ 0 & 0 & 0 & I & 0 & 0 \\ 0 & 0 & 0 & 0 & I & 0 \\ 0 & 0 & 0 & 0 & 0 & I \end{bmatrix}.$$

With two exceptions, the odd-numbered block rows and columns have $n - r_d$ rows and columns, and the even-numbered block rows and columns have $r_d$ rows and columns. The two exceptions are the last block row and column, each of which has $r_0$ rows and $r_0$ columns.

Next take the top left $3 \times 1$ block submatrix of $A$ and transform it to upper triangular form by means of a $QR$ factorization,

$$\begin{bmatrix} \hat{P}_{3,1} \\ \hat{P}_{3,2} \\ -I \end{bmatrix} = Q \begin{bmatrix} R \\ 0 \\ 0 \end{bmatrix}.$$

Then transform the first three block rows of $A$ and $B$ by a premultiplication with $Q^*$. 

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This changes the structure of \((A, B)\) to
\[
A = \begin{bmatrix}
R & X & X & X & X & X \\
0 & X & X & X & X & X \\
0 & -I & 0 & 0 & 0 & 0 \\
0 & 0 & -I & 0 & 0 & 0 \\
0 & 0 & 0 & -I & 0 & 0 \\
0 & 0 & 0 & 0 & X & X \\
0 & 0 & 0 & 0 & X & X \\
\end{bmatrix},
\]
\[
B = \begin{bmatrix}
0 & X & X & 0 & 0 & 0 \\
0 & X & X & 0 & 0 & 0 \\
0 & 0 & 0 & I & 0 & 0 \\
0 & 0 & 0 & 0 & I & 0 \\
0 & 0 & 0 & 0 & 0 & I \\
0 & 0 & 0 & 0 & 0 & I \\
\end{bmatrix},
\]
where \(X\) denotes a dense matrix of appropriate size. This deflates \(n - r_d\) infinite eigenvalues at the top left corner. What remains after removing the first \(n - r_d\) rows and columns is a reduced problem with the structure
\[
(A.6) \quad A = \begin{bmatrix}
\hat{P}_3 & \hat{P}_2 & \hat{P}_1 & X \\
- I & 0 & 0 & 0 \\
0 & -I & 0 & 0 \\
0 & 0 & -I & 0 \\
\end{bmatrix}, \quad B = \begin{bmatrix}
\hat{P}_3 & (0) & 0 & 0 \\
(0) & I & 0 & 0 \\
(0) & 0 & I & 0 \\
(0) & 0 & 0 & I \\
\end{bmatrix}.
\]
The sizes of the blocks in (A.6) are the same as the sizes of the blocks in (A.4), except for the boldface blocks, which have shrunk. The boldface blocks without parentheses have size \(r_d \times r_d\), and one dimension of each boldface block with parentheses has reduced from \(n\) to \(r_d\). It is straightforward to verify that the block sizes agree with the output specification (A.2). This completes our description of the second step of the deflation procedure.

**A.4. The extreme case \(d = 2\).** The extreme case \(d = 2\) requires a little bit of additional care. The first step (section A.2) of the deflation procedure can be applied in the same way as before. The output from the first step takes for \(d = 2\) the form
\[
A = \begin{bmatrix}
P_1 & (- U_1) \\
(P_0) & 0 \\
\end{bmatrix}, \quad B = \begin{bmatrix}
P_2 & (0) \\
(0) & I \\
\end{bmatrix},
\]
where the last block row and column have \(r_0\) rows and columns. What concerns us is what transpires in the second step after having compressed the columns of \(P_2\). According to section A.3 we should next compute a QR factorization of the top left \(3 \times 1\) block submatrix of \(A\). But for \(d = 2\) the third block in that submatrix is not \(- I\) as in (A.5) but instead a dense block \(\hat{P}_{0,1}\) of size \(r_0 \times (n - r_2)\). For \(d > 2\) that block would have been equal to \(- I_{n - r_d}\). What is important for the effect of the QR factorization on the sparsity structure is the number of leading nonzero rows in that
Algorithm 5: Deflation procedure. Deflates zero and infinite eigenvalues induced by rank deficiencies in $P_0$ and/or $P_d$.

1. Construct the matrix pair $(A, B)$ as in (A.3);
2. Compute the ranks $r_0$ and $r_d$ of $P_0$ and $P_d$, respectively;
3. Construct a unitary matrix $U_0$ such that $U_0^*P_0 = \begin{bmatrix} \tilde{P}_0 \end{bmatrix}$ and $\tilde{P}_0$ has full rank;
4. Construct a unitary matrix $U_d$ such that $P_dU_d = \begin{bmatrix} \tilde{P}_d \end{bmatrix}$ and $\tilde{P}_d$ has full rank;
5. Overwrite the $(d, d - 1)$ block of $A$ with $U_0^*P_0$;
6. Overwrite the top right corner of $A$ with $U_0$;
7. Deflate $n - r_0$ zero eigenvalues from the bottom right corner and remove the last $n - r_0$ rows and columns from $A$ and $B$;
8. Overwrite the $(1, 1)$ block of $B$ with $P_dU_d$;
9. for $k \leftarrow 1, 2, \ldots, d - 1$ do
10. \hspace{1em} Overwrite the $(1, k)$ block of $A$ with $P_{d-k}U_d$;
11. \hspace{1em} Overwrite the $(d, d - 1)$ block of $A$ with $(U_0^*P_0)U_d$;
12. if $d = 2$ and $r_0 > n - r_2$ then
13. \hspace{2em} Let $X$ denote the submatrix $A(n + 1: n + r_0, 1: n - r_2)$;
14. \hspace{2em} Compute a QR factorization $X = QR$;
15. \hspace{2em} Update $A$ by premultiplying rows $n + 1: n + r_0$ by $Q^*$;
16. \hspace{2em} Update $A$ by postmultiplying columns $n + 1: n + r_0$ by $Q$;
17. \hspace{1em} Let $X$ denote the submatrix $A(1: 2n - r_d, 1: n - r_d)$;
18. \hspace{1em} Compute a QR factorization $X = QR$;
19. \hspace{1em} Update $A$ and $B$ by premultiplying rows $1: 2n - r_d$ by $Q^*$;
20. \hspace{1em} Deflate $n - r_d$ infinite eigenvalues at the top left corner and remove the first $n - r_d$ rows and columns from $A$ and $B$;

3 × 1 block submatrix. Special care for $d = 2$ is therefore needed if $r_0 > n - r_2$. In that case the number of leading nonzero rows is greater than for the case $d > 2$, and thus a direct application of the second step would lead to more fill. To avoid this problem we first reduce the dense (tall and skinny) $P_{0, 1}$ block to upper triangular form, which in itself has no effect on the generic sparsity structure. But it crucially reduces the number of nonzero leading rows in $P_{0, 1}$ from $r_0$ to the desired $n - r_2$ and thereby overcomes the problem.

A.5. Deflation algorithm. Algorithm 5 summarizes the discussion in the appendix by formalizing the deflation procedure.

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