# On aggressive early deflation in parallel variants of the QR algorithm* 

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#### Abstract

The QR algorithm computes the Schur form of a matrix and is by far the most popular approach for solving dense nonsymmetric eigenvalue problems. Multishift and aggressive early deflation (AED) techniques have led to significantly more efficient sequential implementations of the QR algorithm during the last decade. More recently, these techniques have been incorporated in a novel parallel QR algorithm on hybrid distributed memory HPC systems. While leading to significant performance improvements, it has turned out that AED may become a computational bottleneck as the number of processors increases. In this paper, we discuss a two-level approach for performing AED in a parallel environment, where the lower level consists of a novel combination of AED with the pipelined QR algorithm implemented in the ScaLAPACK routine PDLAHQR. Numerical experiments demonstrate that this new implementation further improves the performance of the parallel QR algorithm.


## 1 Introduction

The solution of matrix eigenvalue problems is a classical topic in numerical linear algebra, with applications in various areas of science and engineering. The QR algorithm developed by Francis and Kublanovskaya, see [9, 19] for recent historic accounts, has become the de facto standard for solving nonsymmetric and dense eigenvalue problems. Parallelizing the QR algorithm has turned out to be highly nontrivial matter [13]. To our knowledge, the ScaLAPACK [5] routine PDLAHQR implemented nearly 10 years ago based on work by Henry, Watkins, and Dongarra [14], is the only publicly available parallel implementation of the QR algorithm. Recently, a novel parallel QR algorithm [10] has been developed, which turns out to be more than a magnitude faster compared to PDLAHQR for sufficiently large problems. These improvements are attained by parallelizing

[^0]the multishift and aggressive early deflation (AED) techniques developed by Braman, Byers, and Mathias [6, 7] for the sequential QR algorithm.

Performed after each QR iteration, AED requires the computation of the Schur form for a trailing principle submatrix (the so called AED window) that is relatively small compared to the size of the whole matrix. In [10], a slightly modified version of the ScaLAPACK routine PDLAHQR is used for this purpose. Due to the small size of the AED window, the execution time spent on AED remains negligible for one or only a few processors but quickly becomes a dominating factor as the number of processors increases. In fact, for a $100000 \times 100000 \mathrm{ma}$ trix and 1024 processor cores, it was observed in [10] that $80 \%$ of the execution time of the QR algorithm was spent on AED. This provides a strong motivation to reconsider the way AED is performed in parallel. In this work, we propose to perform AED by a modification of the ScaLAPACK routine PDLAHQR, which also incorporates AED at this lower level, resulting in a two-level recursive approach for performing AED. The numerical experiments in Section 4 reveal that our new approach reduces the overall execution time of the parallel QR algorithm from [10] by up to $40 \%$.

## 2 Overview of the QR algorithm with AED

In the following, we assume some familiarity with modern variants of the QR algorithm and refer to $[15,18]$ for introductions. It is assumed that the matrix under consideration has already been reduced to (upper) Hessenberg form by, e.g., calling the ScaLAPACK routine PDGEHRD. Algorithm 1 provides a high-level description of the sequential and parallel QR algorithm for Hessenberg matrices, using multiple shifts and AED. Since this paper is mainly concerned with AED, we will only mention that the way the shifts are incorporated in the multishift QR sweep (Step 7) plays a crucial role in attaining good performance, see $[6,10$, 17] for details.

```
Algorithm 1 Multishift Hessenberg \(Q R\) Algorithm with \(A E D\)
    WHILE not converged
        Perform AED on the \(n_{\text {win }} \times n_{\text {win }}\) trailing principle submatrix.
        Apply the accumulated orthogonal transformation to the
        corresponding off-diagonal blocks.
        IF enough eigenvalues have been deflated in step 2
            GOTO step 2.
        END IF
        Perform a multishift QR sweep with undeflatable
        eigenvalues from Step 2 as shifts.
        Check for negligible subdiagonal elements.
    END WHILE
```

In the following, we summarize the AED technique proposed by Braman, Byers, and Mathias [7]. Given an $n \times n$ upper Hessenberg matrix $H$, we partition

$$
H=\begin{gathered}
n-n_{\text {win }}-1 \\
1 \\
n_{\text {win }}
\end{gathered}\left(\begin{array}{ccc}
n-n_{\text {win }}-1 & 1 & n_{\text {win }} \\
H_{11} & H_{12} & H_{13} \\
H_{21} & H_{22} & H_{23} \\
0 & H_{32} & H_{33}
\end{array}\right)
$$

where $n_{\text {win }}$ denotes the size of the AED window. Then a (real) Schur decomposition $H_{33}=V T V^{T}$ is performed, where $V$ is orthogonal and $T$ in upper quasi-triangular form. Setting

$$
U=\begin{gathered}
n-n_{\text {win }}-1 \\
1 \\
n_{\text {win }}
\end{gathered}\left(\begin{array}{ccc}
n-n_{\text {win }}-1 & 1 & n_{\text {win }} \\
I & & \\
& & 1
\end{array}\right)
$$

we obtain

$$
U^{T} H U=\left(\begin{array}{ccc}
H_{11} & H_{12} & H_{13} V \\
H_{21} & H_{22} & H_{23} V \\
0 & s & T
\end{array}\right)
$$

where $s \in \mathbb{R}^{n_{\text {win }}}$ is the so called spike, created from the subdiagonal entry contained in $H_{32}$. The eigenvalues of $T$ are checked subsequently for convergence and possibly deflated. The eigenvalue (or $2 \times 2$ block) in the bottom right corner of $T$ can be deflated if the magnitude of the last component (or the last two components) of the spike is negligibly small. Undeflatable eigenvalue are moved to the top left corner of $T$ by a swapping algorithm [4, 11]. After this transformation is completed, the next eigenvalue in the bottom right corner of $T$ is treated in the same way. The orthogonal transformations for swapping eigenvalues are accumulated in an orthogonal matrix $\tilde{V} \in \mathbb{R}^{n_{\text {win }} \times n_{\text {win }}}$. After all eigenvalues of $T$ have been processed, the entire matrix is reduced back to Hessenberg form and the off-diagonal blocks $H_{13}$ and $H_{23}$ are multiplied with the product of all involved orthogonal transformations. It is recommended to choose $n_{\text {win }}$ somewhat larger, e.g., by $50 \%$, than the number of shifts in the multishift QR iterations [6].

Dramatic performance gains from AED have been observed both for sequential and parallel variants of the QR algorithm. These gains can be achieved essentially no matter how the rest of the QR algorithm is implemented, in particular how many shifts are used in the multishift QR sweep [7]. In effect, any implementation of the QR algorithm may benefit from AED; a fact that we will use below to improve the ScaLAPACK routine PDLAHQR. A convergence analysis, partially explaining the success of AED, can be found in [16].

## 3 Parallel implementation of AED

Since the main aim of this paper is to improve the parallel QR algorithm and implementation described in [10], we first recall the structure of the main routines from this implementation, see Figure 1. The entry routine is PDHSEQR,

Fig. 1. Partial software structure for the parallel QR algorithm from [10].

which branches into PDLAQR1 for small to medium-sized matrices and PDLAQR0 for larger ones. The cut-off point for what is considered medium-sized will be explained in the numerical experiments, see Section 4. The main purpose of PDLAQRO is to call PDLAQR3 for performing AED and PDLAQR5 for performing multishift QR iterations. The former routine invokes PDLAQR1 for performing the Schur decomposition of the AED window. In [10], PDLAQR1 amounts to the ScaLAPACK routine PDLAHQR with minor modifications concerning the processing of $2 \times 2$ blocks in the real Schur form and the multithreaded application of small Householder reflectors. In the following, we will reconsider this choice for PDLAQR1.

### 3.1 Choice of algorithm for performing AED

A number of alternative choices are available for performing the Schur decomposition of the relatively small AED window:

- A recursive call to PDHSEQR or PDLAQRO, implementing the parallel QR algorithm with multishifts and AED.
- A call to PDLAQR1, a minor modification of ScaLAPACK's PDLAHQR.
- Assembling the AED window in local memory and a call to the sequential LAPACK [2] routine DLAHQR (or DLAQR4).

According to the numerical experiments in [10], a recursive call of PDLAQR0 may not be the optimal choice, mainly because of the fact that the way multishift QR iterations are implemented in PDLAQRO suffers from poor scalability for relatively small matrices. ScaLAPACK's PDLAHQR achieves better scalability but does not incorporate modern developments, such as AED, and therefore suffers from poor performance. The third alternative, calling a sequential algorithm, should be used for submatrices that are too small to justify the overhead incurred by parallelization. In our experimental setup this was the case for submatrices of size 384 or smaller.

In this work, we propose to modify PDLAQR1 further and add AED to the parallel pipelined QR algorithm implemented in ScaLAPACK's PDLAHQR. Since the main purpose of PDLAQR1 is to handle small to medium-sized submatrices, a
parallel implementation of AED, as in [10], will not be efficient on this level, since the size of the AED window is even smaller and does not allow for reasonable parallel performance in the Schur decomposition or the swapping of diagonal blocks. We have therefore chosen the third alternative for performing AED on the lowest level and invoke the sequential LAPACK routine DLAQR3 [8]. The accumulated orthogonal transformations returned by DLAQR3 are applied to the off-diagonal blocks in parallel. Therefore, $O(\sqrt{p})$ processors are used for updating the off-diagonal blocks. A high-level description of the resulting procedure is given in Algorithm 2.

```
Algorithm 2 Parallel pipelined \(Q R\) algorithm with \(A E D\) (new PDLAQR1)
    WHILE not converged
        Copy the \(\left(n_{\text {win }}+1\right) \times\left(n_{\text {win }}+1\right)\) trailing submatrix to local memory
        and perform sequential AED on an \(n_{\text {win }} \times n_{\text {win }}\) window.
        Apply the accumulated orthogonal transformations to the
        corresponding off-diagonal blocks in parallel.
        IF enough eigenvalues have been deflated
            GOTO step 2.
        END IF
        Compute the eigenvalues of a trailing submatrix.
        Perform a pipelined QR sweep with the eigenvalues computed
        in step 7 as shifts.
        Check for negligible subdiagonal elements.
10. END WHILE
```


### 3.2 Implementation details

In the following we discuss some implementation issues of Algorithm 2. The basis for our modification is PDLAQR1 from [10], referred to as the old PDLAQR1 in the following discussion. Following the notation established in the (Sca)LAPACK implementations of the QR algorithm, we let $\mathrm{NH}=\mathrm{IHI}-\mathrm{ILO}+1$ denote the dimension of the active $\mathrm{NH} \times \mathrm{NH}$ diagonal block and NS the number of shifts in the multishift QR sweep.

- In the special case when the active diagonal block is small enough, say $\mathrm{NH} \leq$ 384, we copy this block to local memory and call DLAHQR/DLAQR4 directly. The off-diagonal blocks are updated in parallel. This reduces communication while the required extra memory is negligible. We have observed that this modification has a non-negligible positive impact on the total execution time, especially during the final stages of the QR algorithm.
- The size of the deflation window, $n_{\text {win }}$, is determined by the return value of the LAPACK routine IPARMQ, see [8] for more details. In PDLAHQR/PDLAQR1, NS is mainly determined by the process grid and does not exceed 32 . This is usually smaller than the number of shifts suggested by IPARMQ. Also, typical
values of $n_{\text {win }}$ returned by IPARMQ are 96,192 and 384 , which is much larger than if we chose NS*3/2. Based on the observation that the optimal AED window size does not depend strongly on the number of shifts used in the QR sweeps, we prefer to stick to large $n_{\text {win }}$ rather than using NS*3/2. This increases the time spent on AED, but the overhead is compensated by fewer pipelined QR sweeps.
- The criterion for restarting another AED process rightaway, without an intermediate QR iteration, is the same as in LAPACK [8]:

1. The number of undeflatable eigenvalues is smaller than NS; or
2. the number of deflated eigenvalues is larger than $n_{\text {win }} \times 14 \%$.

Note that we choose the criterion in accordance with the window size suggested by IPARMQ.

- In contrast to Algorithm 1, undeflatable eigenvalues are not used as shifts in subsequent multishift QR sweep. This choice is based on numerical experiments with the following three shift strategies:

1. Use undeflatable eigenvalues obtained from AED as shifts.
2. Compute and use the eigenvalues of the NS $\times$ NS trailing submatrix after AED as shifts (by calling DLAHQR/DLAQR4).
3. Compute and use some of the eigenvalues of the $\left(n_{\text {win }}+1\right) \times\left(n_{\text {win }}+1\right)$ trailing submatrix after AED as shifts (by calling DLAHQR/DLAQR4).
An illustration of these strategies is given in Figure 2. Based on the experiments, we prefer the third strategy despite the fact that it is the computationally most expensive one. However, it provides shifts of better quality, mainly because of the larger window size, which was found to reduce the number of pipelined QR sweeps and to outweigh the increased cost for shift computation.

Fig. 2. Three shift strategies $\left(n_{\text {win }}=6\right.$, NS $=4$ )


- When performing AED within the new PDLAQR1, each processor receives a local copy of the trailing submatrix and calls DLAQR3 to execute the same
computations concurrently. This implies redundant work performed in parallel but it reduces communication since the orthogonal transformation matrix, to be applied in parallel in subsequent updates, is readily available on each processor. A similar approach is suggested in the parallel QZ algorithm by Adlerborn et al. [1]. If the trailing submatrix is not laid out across a border of the processor mesh, we call DGEMM to perform the updates. If the trailing submatrix is located on a $2 \times 2$ processor mesh, we organize the computation and communication manually for the update. Otherwise, PDGEMM is used for updating the off-diagonal blocks.


## 4 Numerical Experiments

All the experiments in this section were run on the 64 -bit low power Intel Xeon Linux cluster akka hosted by the High Performance Computing Center North (HPC2N). Akka consists of 672 dual socket quadcore L5420 2.5 GHz nodes, with 16 GB RAM per node, connected in a Cisco Infiniband network. The code is compiled by the PathScale compiler version 3.2 with the flags -02 -fPIC -TENV:frame_pointer=ON -OPT:Olimit=0. The software libraries OpenMPI 1.4.2, BLACS 1.1 patch3, ScaLAPACK/PBLAS 1.8.0, LAPACK 3.2.1 and GOTOBLAS2 1.13 [12] are linked with the code. No multithreaded features, in particular no mixture of OpenMP and MPI, were used. We chose NB $=50$ as the block size in the block cyclic distribution of ScaLAPACK. The test matrices are dense square matrices with entries randomly generated from a uniform distribution in $[0,1]$. The ScaLAPACK routine PDGEHRD is used to reduce these matrices initially to Hessenberg form. We only measure the time for the Hessenberg QR algorithm, i.e., the reduction from Hessenberg to real Schur form.

### 4.1 Improvement for PDLAQR1

We first consider the isolated performance of the new PDLAQR1 compared to the old PDLAQR1 from [10]. The sizes of the test matrices were chosen to fit the typical sizes of the AED windows suggested in [10]. Table 1 displays the measured execution time on various processor meshes. For determining the cross-over point for switching from PDLAQRO to PDLAQR1 in the main routine PDGSEQR, we also measured the execution time of PDLAQRO.

The new implementation of PDLAQR1 turns out to require much less time than the old one, with a few, practically nearly irrelevant exceptions. Also, the new PDLAQR1 scales slightly better than PDLAQR0, especially when the size of matrix is not large. It is worth emphasizing that the scaling of all implementations eventually deteriorates as the number of processor increases, simply because the involved matrices are not sufficiently large to create enough potential for parallelization.

Quite naturally, PDLAQR0 becomes faster than the new PDLAQR1 as the matrix size increases. The dashed line in Table 1 indicates the crossover point between both implementations. A rough model of this crossover point results is given by

Table 1. Execution time in seconds for old PDLAQR1 (1st line for each $n$ ), new PDLAQR1 (2nd line) and PDLAQRO (3rd line). The dashed line is the crossover point between the new PDLAQR1 and PDLAQRO.

| Matrix size | Processor mesh |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ( $n$ ) | $1 \times 1$ | $2 \times 2$ | $3 \times 3$ | $4 \times 4$ | $6 \times 6$ | $8 \times 8$ | $10 \times 10$ |
| 96 | 0.01 | 0.05 | 0.11 | 0.18 | 0.15 | 0.25 | 0.27 |
|  | 0.08 | 0.08 | 0.02 | 0.05 | 0.03 | 0.08 | 0.07 |
|  | 0.14 | 0.40 | 0.96 | 1.11 | 2.52 | 3.16 | 2.95 |
| 192 | 0.09 | 0.17 | 0.18 | 0.22 | 0.32 | 0.47 | 0.64 |
|  | 0.09 | 0.07 | 0.07 | 0.13 | 0.16 | 0.12 | 0.26 |
|  | 0.15 | 0.30 | 0.61 | 1.05 | 3.73 | 4.34 | 3.64 |
| 384 | 0.60 | 0.73 | 0.61 | 0.63 | 0.78 | 1.09 | 1.24 |
|  | 0.27 | 0.29 | 0.28 | 0.36 | 0.40 | 0.48 | 0.48 |
|  | 0.47 | 0.55 | 0.72 | 0.89 | 2.08 | 3.23 | 3.76 |
| 768 | 7.3 8- | $\overline{3} .5 \overline{3}$ | $\overline{2} .53{ }^{\text {² }}$ | 2.35 | 2.61 | 2.80 | 3.52 |
|  | 3.77 | 2.24 | 1.73 । | 1.57 | 1.73 | 2.17 | 2.25 |
|  | 1.83 | 1.51 | $1.61{ }^{1}$ | 1.68 | 2.70 | 3.03 | 3.31 |
| 1536 | 133.31 | 20.68 | $13.23{ }^{\text {L }}$ | -11. 127 | 9.79 | 10.48 | 13.05 |
|  | 35.94 | 9.27 | 6.54 | 5.521 | 5.11 | 5.31 | 6.33 |
|  | 12.34 | 6.61 | 5.63 | $4.86{ }^{1}$ | 6.26 | 6.76 | 6.84 |
| 3072 | 2313.61 | 139.05 | 96.73 | $66.06{ }^{\text {L }}$ | 50.64 | $\overline{4} 1.8 \overline{2}$ | $6 \overline{3} .22$ |
|  | 522.81 | 45.72 | 33.13 | 22.60 | 19.08 | 18.12 | 22.23 |
|  | 80.71 | 30.67 | 21.34 | 15.82 | 15.56 | 15.09 | 14.98 |
| 6144 |  | 1049.56 | 623.63 | 351.44 | 231.70 | 199.75 | 227.45 |
|  |  | 144.96 | 167.71 | 103.15 | 78.75 | 66.90 | 70.48 |
|  |  | 198.54 | 129.58 | 87.07 | 55.40 | 47.61 | 44.07 |

$n=220 \sqrt{p}$, which fits the observations reasonably well and has been incorporated in our implementation.

### 4.2 Overall Improvement

As the main motivation for the development of the new PDLAQR1 is its application to AED within the parallel QR algorithm, we have also measured the resulting reduction of the overall execution time of PDHSEQR. From the results presented in Table 2, it is clear that PDHSEQR with the new PDLAQR1 is almost always better than the old implementation. The improvement varies between $5 \%$ and $40 \%$. Among the measured configurations, there is one notable exception: $n=32000$ on a $6 \times 6$ processor grid. This is actually the only case for which PDLAQRO is called within the AED phase, which seems to indicate that the choice of the crossover point requires some additional fine tuning.

Note that the largest AED window in all these experiments is of size 1536. According to Table 1, we expect even more significant improvements for larger matrices, which have larger AED windows.

Table 2. Execution time in seconds for old PDHSEQR (1st line for each $n$ ), new PDHSEQR (2nd line). The third lines show the relative improvement.

| Processor mesh | Matrix size $(n)$ |  |  |  |
| :---: | ---: | ---: | ---: | ---: |
|  | 4000 | 8000 | 16000 | 32000 |
| $1 \times 1$ | 162.43 |  |  |  |
|  | 161.28 |  |  |  |
|  | $0.71 \%$ |  |  |  |
| $2 \times 2$ | 71.34 | 501.83 |  |  |
|  | 68.02 | 452.70 |  |  |
|  | $4.65 \%$ | $9.79 \%$ |  |  |
| $4 \times 4$ | 39.18 | 170.75 | 1232.40 |  |
|  | 30.68 | 158.66 | 1037.93 |  |
|  | $22.69 \%$ | $7.08 \%$ | $15.78 \%$ |  |
|  | 35.96 | 123.46 | 617.97 | 3442.08 |
|  | 24.62 | 96.23 | 509.38 | 3584.74 |
|  | $31.54 \%$ | $22.06 \%$ | $17.57 \%$ | $-4.14 \%$ |
| $10 \times 10$ | 33.09 | 97.20 | 435.52 | 2639.32 |
|  | 20.59 | 67.42 | 366.31 | 2016.93 |
|  | $37.78 \%$ | $31.64 \%$ | $15.89 \%$ | $24.58 \%$ |
|  | 36.05 | 101.75 | 355.38 | 2053.16 |
|  | 21.39 | 62.29 | 291.06 | 1646.30 |
|  | $41.67 \%$ | $39.58 \%$ | $18.10 \%$ | $19.82 \%$ |

## 5 Summary

We have reconsidered the way AED is performed in the parallel QR algorithm [10]. A recursive approach is suggested, in which the ScaLAPACK routine PDLAHQR is combined with AED to address medium-sized problems. The focus of this work has been on minimizing the total execution time instead of how to take utility of all the processors or how well the algorithm scales. Computational experiments demonstrate the efficiency of our approach, but also reveal potential for further improvements by a more careful fine tuning of the crossover point for switching between different implementations of the parallel QR algorithm.

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