Distance-Two Interpolation for Parallel Algebraic Multigrid

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Abstract. Algebraic multigrid (AMG) is one of the most efficient and scalable parallel algorithms for solving sparse linear systems on unstructured grids. However, for large three-dimensional problems, the coarse grids that are normally used in AMG often lead to growing complexity in terms of memory use and execution time per AMG V-cycle. Sparser coarse grids, such as those obtained by the Parallel Modified Independent Set coarsening algorithm (PMIS) [7], remedy this complexity growth, but lead to non-scalable AMG convergence factors when traditional distance-one interpolation methods are used. In this paper we study the scalability of AMG methods that combine PMIS coarse grids with long distance interpolation methods. AMG performance and scalability is compared for previously introduced interpolation methods as well as new variants of them for a variety of relevant test problems on parallel computers. It is shown that the increased interpolation accuracy largely restores the scalability of AMG convergence factors for PMIS-coarsened grids, and in combination with complexity reducing methods, such as interpolation truncation, one obtains a class of parallel AMG methods that enjoy excellent scalability properties on large parallel computers.

Key words. Algebraic multigrid, long range interpolation, parallel implementation, reduced complexity, truncation

AMS subject classifications. 65F10, 65N55, 65Y05

1. Introduction. Algebraic multigrid (AMG) [2, 5] is an efficient potentially scalable algorithm for sparse linear systems on unstructured grids. However, when applied to large three-dimensional problems, the classical algorithm often generates unreasonably large complexities with regard to memory use as well as computational operations. Recently, we suggested a new parallel coarsening algorithm, called the Parallel Modified Independent Set algorithm (PMIS) [7], which is based on a parallel independent set algorithm suggested in [10]. The use of this coarsening algorithm in combination with a slight modification of Ruge and Stueben’s classical interpolation scheme [11], leads to significantly lower complexities as well as significantly lower setup and cycle times. For various test problems, such as isotropic and grid aligned anisotropic diffusion operators, one obtains scalable results, particularly when AMG is used in combination with Krylov methods. However, AMG convergence factors are severely impacted for more complicated problems, such as problems with rotated anisotropies or highly discontinuous material properties. Since we realized that classical interpolation methods, which use only distance-one neighbors for their interpolatory set, were not sufficient for these coarse grids, we decided to investigate interpolation operators which also include distance-two neighbors. In this paper we focus on the following distance-two interpolation operators: we study three methods proposed in [12], namely, standard interpolation, multipass interpolation, and the
use of Jacobi interpolation to improve other interpolation operators, and we investi-
gate two extensions of classical interpolation, which we denote with ‘extended’ and
‘extended+i’ interpolation.

Our investigation shows that all of the long-distance interpolation strategies,
except for multipass interpolation, significantly improve AMG convergence factors
compared to classical interpolation. Multipass interpolation shows poor numerical
scalability, which, however, can be improved with a Krylov accelerator, but it has
very small computational complexity. All other long-distance interpolation operators
showed increased complexities. While the increase is not very significant for two-
dimensional problems, it is of concern in the three-dimensional case. Therefore we
also investigated complexity reducing strategies, such as the use of smaller sets of
interpolation points and interpolation truncation. The use of these strategies led to
AMG methods with significantly improved overall scalability.

The paper is organized as follows. In Section 2 we briefly describe AMG. In Sec-
tion 3 distance-one interpolation operators are presented, and Section 4 describes long
range interpolation operators. In Section 5 the computational cost of the interpola-
tion strategies is investigated, and in Section 6 some sequential numerical results are
given, which motivate the following sections. Section 7 presents various complexity
reducing strategies. Section 8 investigates the parallel implementation of the methods.
Section 9 presents parallel scaling results for a variety of test problems, and Section
10 contains the conclusions.

2. Algebraic Multigrid. In this section we give an outline of the basic prin-
ciples and techniques that comprise AMG, and we define terminology and notation.
Detailed explanations may be found in [4, 11, 12]. Consider a problem of the form

\[ Au = f, \quad (2.1) \]

where \( A \) is an \( n \times n \) matrix with entries \( a_{ij} \). For convenience, the indices are identified
with grid points, so that \( u_i \) denotes the value of \( u \) at point \( i \), and the grid is denoted by
\( \Omega = \{1, 2, \ldots, n\} \). In any multigrid method, the central idea is that “smooth error,” \( e \),
that is not eliminated by relaxation must be removed by coarse-grid correction. This
is done by solving the residual equation \( Ae = r \) on a coarser grid, then interpolating
the error back to the fine grid and using it to correct the fine-grid approximation.

Using superscripts to indicate level number, where 1 denotes the finest level so
that \( A^1 = A \) and \( \Omega^1 = \Omega \), AMG needs the following components: “grids” \( \Omega^1 \supset \Omega^2 \supset \ldots \supset \Omega^M \), grid operators \( A^1, A^2, \ldots, A^M \), interpolation operators \( P^k \), restriction
operators \( R^k \) (often \( R^k = (P^k)^T \)), and smoothers \( S^k \), where \( k = 1, 2, \ldots, M - 1 \).

Most of these components of AMG are determined in a first step, known as the
setup phase. During the setup phase, on each level \( k, k = 1, \ldots, M - 1 \), \( \Omega^{k+1} \) is
determined using a coarsening algorithm, \( P^k \) and \( R^k \) are defined and the \( A^{k+1} \) is
determined using the Galerkin condition \( A^{k+1} = R^k A^k P^k \). Once the setup phase is
completed, the solve phase, a recursively defined cycle, can be performed as follows:
Algorithm: $MGV(A^k, R^k, P^k, S^k, u^k, f^k)$. 
If $k = M$, solve $A^M u^M = f^M$ with a direct solver. 
Otherwise:
   - Apply smoother $S^k$ $\mu_1$ times to $A^k u^k = f^k$. 
   - Perform coarse grid correction: 
     - Set $r^k = f^k - A^k u^k$. 
     - Set $r^{k+1} = R^k r^k$. 
     - Apply $MGV(A^{k+1}, R^{k+1}, P^{k+1}, S^{k+1}, e^{k+1}, r^{k+1})$. 
     - Interpolate $e^k = P^k e^{k+1}$. 
   - Correct the solution by $u^k = u^k + e^k$. 
   - Apply smoother $S^k$ $\mu_2$ times to $A^k u^k = f^k$. 

In the remainder of the paper, index $k$ will be dropped for simplicity. The algorithm above describes a $V(\mu_1, \mu_2)$-cycle; other more complex cycles such as $W$-cycles are described in [4]. In every V-cycle, the error is reduced by a certain factor, which is called the convergence factor. A sequence of V-cycles is executed until the error is reduced below a specified tolerance. For a scalable AMG method, the convergence factor is bounded away from one as a function of the problem size $n$, and the computational work in both the setup and solve phases is linearly proportional to the problem size $n$. While AMG was originally developed in the context of symmetric M-matrix problems, AMG has been applied successfully to a much wider class of problems. We assume in this paper that $A$ has positive diagonal elements.

3. Distance-One Interpolation Strategies. In this section, we first give some definitions as well as some general remarks, and then recall the possibly simplest interpolation strategy, the so-called direct interpolation strategy [12]. This is followed by a description of the classical distance-one AMG interpolation method that was introduced by Ruge and Stuben [11].

3.1. Definitions and Remarks. One of the concepts used in the following sections is strength of connection. A point $j$ strongly influences a point $i$ or $i$ strongly depends on $j$ if

$$-a_{i,j} > \alpha \max_{k \neq i}(-a_{i,k}),$$

where $0 < \alpha < 1$. We set $\alpha = 0.25$ in the remainder of the paper.

We define the measure of a point $i$ as the number of points which strongly depend on $i$. When PMIS coarsening is used, a positive random number that is smaller than 1 is added to the measure to distinguish between neighboring points that strongly influence the same number of points. In the PMIS coarsening algorithm, points that do not strongly influence any other points are initialized as $F$-points.

Using this concept of strength of connection we define the following sets:

$$N_i = \{j | a_{ij} \neq 0\},$$
$$S_i = \{j \in N_i | j \text{ strongly influences } i\},$$
$$F^s_i = F \cap S_i,$$
$$C^s_i = C \cap S_i,$$
$$N^s_i = N_i \setminus (F^s_i \cap C^s_i).$$

In classical AMG [11], the interpolation of the error at the $F$-point $i$ takes the
form

\[ e_i = \sum_{j \in C_i} w_{ij} e_j, \quad (3.2) \]

where \( w_{ij} \) is an interpolation weight determining the contribution of the value \( e_j \) to \( e_i \), and \( C_i \subset C \) is the coarse interpolatory set of \( F \)-point \( i \). In most classical approaches to AMG interpolation, \( C_i \) is a subset of the nearest neighbors of grid point \( i \), i.e. \( C_i \subset N_i \), and longer-range interpolation is not considered.

The points to which \( i \) is connected, comprise three sets: \( C^s_i \), \( F^s_i \) and \( N^w_i \). Based on assumptions on small residuals for smooth error \([2, 4, 11, 12]\), an interpolation formula can be derived as follows. The assumption that algebraically smooth error has small residuals after relaxation

\[ A e \approx 0, \]

can be rewritten as

\[ a_{ii} e_i \approx - \sum_{j \in N_i} a_{ij} e_j, \quad (3.3) \]

or

\[ a_{ii} e_i \approx - \sum_{j \in C^s_i} a_{ij} e_j - \sum_{j \in F^s_i} a_{ij} e_j - \sum_{j \in N^w_i} a_{ij} e_j. \quad (3.4) \]

From this expression, various interpolation formulae can be derived. We use the terminology of \([12]\) for the various interpolation strategies.

3.2. Direct Interpolation. The so-called ‘direct interpolation’ strategy \([12]\) has one of the most simple interpolation formulae. The coarse interpolatory set is chosen as \( C_i = C^s_i \), and

\[ w_{ij} = - \frac{a_{ij}}{a_{ii} \sum_{k \in C^s_i} a_{ik}}, \quad j \in C^s_i. \quad (3.5) \]

This leads to an interpolation, which is often not accurate enough. Nevertheless, we mention this approach here, since various other interpolation operators which we consider are based on it. This method is denoted by ‘direct’ in the tables presented below. In \([12]\) it is also suggested to separate positive and negative coecients when determining the weights, a strategy which can help when one encounters large positive off-diagonal matrix coefficients. We do not consider this approach here, since the strategy did not lead to an improvement for the problems we consider here.

3.3. Classical Interpolation. A generally more accurate distance-one interpolation formula is the interpolation suggested by Ruge and Stüben in \([11]\), which we call ‘classical interpolation’ (‘clas’). Again, \( C_i = C^s_i \), but the contribution from strongly influencing \( F \)-points (the points in \( F^s_i \)) in (3.3) is taken into account more carefully. An appropriate approximation for the errors \( e_j \) of those strongly influencing \( F \)-points may be defined as

\[ e_j \approx \frac{\sum_{k \in C_i} a_{jk} e_k}{\sum_{k \in C_i} a_{jk}}. \quad (3.6) \]
This approximation can be justified by the observation that smooth error varies slowly in the direction of strong connection. The denominator simply ensures that constants are interpolated exactly. Replacing the \( e_j \) with a sum over the elements \( k \) of the coarse interpolatory set \( C_i \) corresponds to taking into account strong \( F - F \) connections using \( C \)-points that are common between the \( F \)-points. Note that, when the two \( F \)-points \( i \) and \( j \) do not have a common \( C \)-point in \( C_i \) and \( C_j \), the denominator in (3.6) is small or vanishing. Weak connections (from the points in \( N_i^w \)) are generally not important and errors \( e_j, j \in N_i^w \) are replaced by \( e_i \). This leads to the following formula for the interpolation weights:

\[
    w_{ij} = -\frac{1}{a_{ii} + \sum_{k \in N_i^w} a_{ik}} \left( a_{ij} + \sum_{k \in F_i} \frac{a_{ik}a_{kj}}{\sum_{m \in C_i} a_{km}} \right), \quad j \in C_i^s. \tag{3.7}
\]

In our experiments this interpolation is further modified as proposed in [9] to avoid extremely large interpolation weights that can lead to divergence.

Now the interpolation above was suggested based on a coarsening algorithm that ensured that two strongly connected \( F \)-points always have a common coarse neighbor. Since this condition is no longer guaranteed when using PMIS coarsening [7], it may happen that the term \( \sum_{m \in C_i} a_{km} \) in Eq. (3.7) vanishes. In our previous paper on the PMIS coarsening method [7], we modified interpolation formula (3.7) such that if this case occurs, \( a_{ik} \) is added to the diagonal term (the term \( a_{ii} + \sum_{k \in N_i^w} a_{ik} \) in Eq. (3.7)), i.e., the strongly influencing neighbor point \( k \) of \( i \) is treated like a weak connection of \( i \). In what follows, we denote the set of strongly connected neighbors \( k \) of \( i \) that are \( F \)-points but do not have a common \( C \)-point, i.e. \( C_i^s \cap C_k = \emptyset \), by \( F_i^{**} \). Combining this with the modification suggested in [9] we obtain the following interpolation formula:

\[
    w_{ij} = -\frac{1}{a_{ii} + \sum_{k \in N_i^w \cup F_i^{**}} a_{ik}} \left( a_{ij} + \sum_{k \in F_i^{**}} \frac{a_{ik}\tilde{a}_{kj}}{\sum_{m \in C_i} \tilde{a}_{km}} \right), \quad j \in C_i^s \tag{3.8}
\]

where

\[
    \tilde{a}_{ij} = \begin{cases} 
    0 & \text{if } \text{sign}(a_{ij}) = \text{sign}(a_{ii}) \\ 
    a_{ij} & \text{otherwise}. 
    \end{cases}
\]

In this paper we refer to formula (3.8) as ‘classical interpolation’. The numerical results that were presented in [7] showed that this interpolation formula, which is based on Ruge and Stüben’s original distance-one interpolation formula [11], resulted in AMG methods with acceptable performance when used with PMIS-coarsened grids for various problems, but only when the AMG cycle is accelerated by a Krylov subspace method. Without such acceleration, interpolation formula (3.8) is not accurate enough on PMIS-coarsened grids: AMG convergence factors deteriorate quickly as a function of problem size, and scalability is lost. For various problems, such as problems with rotated anisotropies or problems with large discontinuities, adding Krylov acceleration did not remedy the scalability problems.

One of the issues is that distance-one interpolation schemes do not treat situations like the one illustrated in Figure 3.1 correctly. Here we have an \( F \)-point with measure
smaller than 1 that has no coarse neighbors. This situation can occur for example if we have a fairly large strength threshold. Both for classical and direct interpolation, the interpolated error in this point will vanish, and coarse grid correction will not be able to reduce the error in this point.

Fig. 3.1. Example illustrating a situation occurring with PMIS coarsening, which will not correctly be treated by direct or classical interpolation. Black points denote $C$-points, white points denote $F$-points, and the arrow from $i$ to $l$ denotes that $i$ strongly depends on $l$.

A major topic of this paper is to investigate whether distance-two interpolation methods are able to restore grid-independent convergence to AMG cycles that use PMIS-coarsened grids, without compromising scalability in terms of memory use and execution time per AMG V-cycle.

4. Long-Range Interpolation Strategies. In this section, various long-distance interpolation methods are described. Parallel implementation of some of these interpolation methods and parallel scalability results on PMIS-coarsened grids are discussed later in this paper.

4.1. Multipass Interpolation. Multipass interpolation (‘mp’) is suggested in [12], and is useful for low complexity coarsening algorithms, particularly so-called aggressive coarsening [12]. We suggested it in [7] as a possible interpolation scheme to fix some of the problems that we saw when using our classical interpolation scheme (3.8). Multipass interpolation proceeds as follows:

1. Use direct interpolation for all $F$-points $i$, for which $C_i^* \neq \emptyset$. Place these points in set $F^*$.
2. For all $i \in F \setminus F^*$ with $F^* \cap F_i^* \neq 0$, replace, in Equation (3.3), for all $j \in F_i^* \cap F^*$, $e_j$ by $\sum_{k \in C_j} w_{jk} e_k$, where $C_j$ is the interpolatory set for $e_j$. Apply direct interpolation to the new equation. Add $i$ to $F^*$. Repeat step 2 until $F^* = F$.

Multipass interpolation is fairly cheap. However, it is not very powerful, since it is based on direct interpolation. If applied to PMIS, it still ends up being direct interpolation for most $F$-points. However, it fixes the situation illustrated in Figure 3.1. If we apply multipass interpolation, the point $i$ will be interpolated by the coarse neighbors (black points) of $F$-points $k$ and $l$.

4.2. Jacobi Interpolation. Another approach that remedies convergence issues caused by distance-one interpolation formulae is Jacobi interpolation [12]. This approach uses an existing interpolation formula and applies one or more Jacobi iteration steps to the $F$-point portion of the interpolation operator leading to a more accurate interpolation operator.

Assuming that $A$ and the interpolation operator $P^{(n)}$ are reordered according to the $C/F$-splitting and can be written in the following way:

$$A = \begin{pmatrix} A_{FF} & A_{FC} \\ A_{CF} & A_{CC} \end{pmatrix}, \quad P^{(n)} = \begin{pmatrix} P_{FC}^{(n)} \\ I_{CC} \end{pmatrix},$$

then Jacobi iteration on $A_{FF} e_F + A_{FC} e_C = 0$, with initial guess $e_F^{(0)} = P_{FC}^{(0)} e_C$, leads
to

$$P_{FC}^{(n)} = (I_{FF} - D_{FF}^{-1}A_{FF})P_{FC}^{(n-1)} - D_{FF}^{-1}A_{FC}, \quad (4.2)$$

where $D_{FF}$ is the diagonal matrix containing the diagonal of $A_{FF}$, and $I_{FF}$ and $I_{CC}$ are identity matrices.

If we apply this approach to a distance-one interpolation operator like classical interpolation, we obtain an improved long distance interpolation operator. This approach is also recommended to be used to improve multipass interpolation. We include results where classical interpolation is used followed by one step of Jacobi interpolation in our numerical experiments and denote them by 'clas+j'.

4.3. Standard Interpolation. Standard interpolation ('std') extends the interpolatory set that is used for direct interpolation [12]. This is done by extending the stencil obtained through (3.3) via substitution of every $e_j$ with $j \in F_i^n$ by $1/a_{jj} \sum_{k \in N_i} a_{jk} e_k$. This leads to the following formula

$$\hat{a}_{ii} e_i + \sum_{j \in \hat{N}_i} \hat{a}_{ij} e_j \approx 0$$

with the new neighborhood $\hat{N}_i = N_i \cup \bigcup_{j \in F_i^n} N_j$ and the new coarse point set $\hat{C}_i = C_i \cup \bigcup_{j \in F_i^n} C_j$. This can greatly increase the size of the interpolatory set.

See the left example in Figure 4.1. Consider point $i$. Using direct or classical interpolation, $i$ would only be interpolated by the two distance-one coarse points. However, when we include the coarse points of its strong fine neighbors $m$ and $n$, two additional interpolatory points $k$ and $l$ are added, leading to a potentially more accurate interpolation formula. Standard interpolation is now defined by applying direct interpolation to the new stencil, leading to

$$w_{ij} = -\frac{\hat{a}_{ij} \sum_{k \in \hat{N}_i} \hat{a}_{ik}}{\hat{a}_{ii} \sum_{k \in \hat{C}_i} \hat{a}_{ik}}. \quad (4.4)$$

4.4. Extended Interpolation. It is possible to extend the classical interpolation formula so that the interpolatory set includes $C$-points that are distance two away from the $F$-point to be interpolated, i.e. applying the classical interpolation formula, but using the same interpolatory set that is used in standard interpolation, see Figure 4.1: $\hat{C}_i = C_i \cup \bigcup_{j \in F_i^n} C_j$.

Using the same reasoning that leads to the classical interpolation formula (3.7), the following approximate statement can be made regarding the error at an $F$-point $i$:

$$\left( a_{ii} + \sum_{j \in N_i^n} a_{ij} \right) e_i \approx -\sum_{j \in \hat{C}_i} a_{ij} e_j - \sum_{j \in F_i^n} a_{ij} \sum_{k \in \hat{C}_i} a_{jk} e_k.$$  \quad (4.5)

It then follows immediately that the interpolation weights using the extended
Coarse interpolatory set $\hat{C}_i$ can be defined as

$$w_{ij} = \frac{1}{a_{ii} + \sum_{k \in N_i^w} a_{ik}} \left( a_{ij} + \sum_{k \in F_i^w} \sum_{m \in \hat{C}_i} \tilde{a}_{km} \right), \quad j \in \hat{C}_i. \quad (4.6)$$

This new interpolation formula deals efficiently with strong $F - F$ connections that do not share a common $C$-point. We call this interpolation strategy ‘extended interpolation’ (‘ext’).

**4.5. Extended+I Interpolation.** While extended interpolation remedies many problems that occur with classical interpolation, it does not always lead to the desired weights. Consider the case given in Figure 4.2. Here we have a 1-dimensional Laplace-problem generated by finite differences. Points 1 and 2 are strongly connected $F$-points, and points 0 and 3 are coarse points. Clearly 0, 3 is the interpolatory set for point 1 for the case of extended interpolation. If we apply formula (4.6) to this example to calculate $w_{1,0}$ and $w_{1,3}$, we obtain

$$w_{1,0} = 0.5, \quad w_{1,3} = 0.5.$$  

This is a better result than we would obtain for direct interpolation (3.5) and classical interpolation (3.8):

$$w_{1,0} = 1, \quad w_{1,3} = 0.$$
but worse than standard interpolation (4.4), for which we get the intuitively best interpolation weights:

\[ w_{1,0} = 2/3, \quad w_{1,3} = 1/3. \] (4.7)

This can be remedied if we include not only connections \( a_{jk} \) from strong fine neighbors \( j \) of \( i \) to points \( k \) of the interpolatory set but also connections \( a_{ji} \) from \( j \) to point \( i \) itself. An alternative to expression (3.6) for the error in strongly connected \( F \)-points, is then given by

\[ e_j \approx \frac{\sum_{k \in C_i \cup \{i\}} a_{jk} c_k}{\sum_{k \in C_i \cup \{i\}} a_{jk}}. \] (4.8)

This can be rewritten as

\[ e_j \approx \frac{\sum_{k \in C_i} a_{jk} c_k}{\sum_{k \in C_i \cup \{i\}} a_{jk}} + \frac{\sum_{k \in C_i \cup \{i\}} a_{ji} e_i}{\sum_{k \in C_i \cup \{i\}} a_{jk}}, \] (4.9)

which then, in a similar way as before, leads to interpolation weights

\[ w_{ij} = \frac{1}{a_{ii}} \left( a_{ij} + \sum_{k \in F_i^*} \frac{\bar{a}_{kj}}{\sum_{l \in C_i \cup \{i\}} \bar{a}_{kl}} \right), \quad j \in \hat{C}_i, \] (4.10)

with now

\[ \bar{a}_{ii} = a_{ii} + \sum_{n \in D_i^{sw}} a_{in} + \sum_{k \in F_i^*} a_{ik} \sum_{l \in C_i \cup \{i\}} \bar{a}_{kl}. \] (4.11)

We call this modified extended interpolation ‘extended+i’, and refer to it by ‘ext+i’ (or sometimes ‘e+i’ to save space) in the tables below. If we apply it to the example illustrated in Figure 4.2 we obtain weights (4.7).

5. Computational Cost of Interpolation Strategies. In this section we consider the cost of some of the interpolation operators described in the previous sections. We use the following notations:

- \( N_c \) total number of coarse points
- \( N_f \) total number of fine points
- \( n_k \) average number of distance-\( k \) neighbor points
- \( c_k \) average number of distance-\( k \) interpolatory points
- \( f_k \) average number of strong fine distance-\( k \) neighbors
- \( w_k \) average number of weak distance-\( k \) neighbors
- \( s_k \) average number of common distance-\( k \) interpolatory points
- \( f_w \) average number of strong neighbors treated weakly

Here \( f_w \) indicates the number of strong \( F \)-neighbors that are treated weakly, which occur only for classical interpolation (3.7). Also, \( s_k \) denotes the average number of \( C \)-points which are distance-one neighbors of \( j \in F_i^* \) and also distance-\( k \) interpolatory points for \( i \), the point to be interpolated. Note that \( s_k \) is usually smaller than \( c_k \) and at most equal to \( c_k \). Note also that \( n_k = f_k + c_k + w_k \).
In our considerations we assume a compressed sparse row data format, i.e. three arrays are used to store the matrix: a real array that contains the coefficients of the matrix, an integer array that contains the column indices for each coefficient and an integer array that contains pointers to the beginning of each row for the other two arrays. We also assume an additional integer array which indicates whether a point is an $F$- or a $C$-point.

For all interpolation operators mentioned before, it is necessary to determine at first the interpolatory set. At the same time the data structure for the interpolation operator can be determined. This can be accomplished by sweeping through each row that belongs to an $F$-point: coarse neighbors are identified via integer comparisons, and the pointer array for the interpolation operator is generated. For the distance-two interpolation schemes, it is also necessary to check neighbors of strong fine neighbors. This requires $n_1$ comparisons for direct and classical interpolation, and $(f_1 + 1)n_1$ comparisons for extended, extended+i and standard interpolation. The final data structure contains $N_c + N_f c_1$ coefficients for classical and direct interpolation, and $N_c + N_f (c_1 + c_2)$ coefficients for extended(+i) and standard interpolation.

Next, the interpolation data structure is filled.

For direct interpolation, all that is required is to sweep through a whole row once to compute $\alpha_i = -\sum_{k \in N_i} a_{ik}/\left(a_{ii} \sum_{k \in C_i} a_{ik}\right)$ and then multiply the relevant matrix elements $a_{ij}$ with $\alpha_i$. The sum in the denominator requires an additional $n_1$ comparisons, and the two summations require $n_1 + 1$ additions.

For classical, extended and extended+i interpolation one needs to first compute for each point $k \in F_i^s \setminus F_i^{s\ast}$, $\alpha_{ik} = a_{ik}/\left(\sum_{m \in D_{ik}} a_{km}\right)$ with the appropriate set $D_i^s$, which is different for each operator. For example, for classical interpolation, this requires $f_1 n_1$ comparisons. After this step, all these coefficients need to be processed again in order to add $\alpha_{ij} a_{kj}$ to the appropriate weights. This requires an additional $f_1 n_1$ comparisons. The number of additions, multiplications and divisions can be determined similarly.

For standard interpolation, at first the new stencil needs to be computed, leading to $f_1 n_1$ additions and multiplications. After this one proceeds just as for direct interpolation with a much larger stencil of size $n_1 + n_2$.

The number of floating point additions, multiplications and divisions to compute all interpolation weights for each $F$-point are given in Table 5.1. Note that a sum over $m$ elements is treated as $m$ additions, assuming that we are adding to a variable that was originally 0. Note also that occurrences of products of variables, such as $f_i c_i$ or $f_is_i$, are of order $n_i^2$, since $f_i$, $c_i$, $s_i$ are dependent on $n_i$. This is also reflected in the results given in Table 5.2 for two specific examples.

<table>
<thead>
<tr>
<th>Interp.</th>
<th>additions</th>
<th>multiplications</th>
<th>divisions</th>
<th>comparisons</th>
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</thead>
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<td>$c_1 + 1$</td>
<td>$1$</td>
<td>$2n_1$</td>
</tr>
<tr>
<td>clas</td>
<td>$2f_1 s_1 + w_1 + f_w$</td>
<td>$f_1 s_1 + c_1$</td>
<td>$f_1 - f_w + 1$</td>
<td>$(2f_1 + 1)n_1$</td>
</tr>
<tr>
<td>std</td>
<td>$(f_1 + 1)n_1 + n_2$</td>
<td>$f_1 n_1 + c_1 + c_2$</td>
<td>$f_1 + 1$</td>
<td>$(2f_1 + 2)n_1 - f_1 + n_2$</td>
</tr>
<tr>
<td>ext</td>
<td>$2f_1 (s_1 + s_2)$</td>
<td>$f_1 (s_1 + s_2)$</td>
<td>$f_1 + 1$</td>
<td>$(3f_1 + 1)n_1 - f_1$</td>
</tr>
<tr>
<td>ext+i</td>
<td>$2f_1 (s_1 + s_2 + 1)$</td>
<td>$f_1 (s_1 + s_2 + 1)$</td>
<td>$f_1 + 1$</td>
<td>$(3f_1 + 1)n_1 - f_1$</td>
</tr>
</tbody>
</table>

Table 5.1

Computational cost for various interpolation operators.
Let us look at some examples to get an idea about the actual cost involved. First consider a 5-point stencil as in Figure 4.1. Here, we have the following parameters: 
\[ c_1 = f_1 = 2, \quad w_1 = w_2 = 0, \quad n_1 = 4, \quad f_w = 2, \quad s_1 = 0, \quad c_2 = 2, \quad f_2 = 3, \quad n_2 = 5, \quad s_2 = 1.5. \]

Table 5.2 shows the resulting interpolation cost. Next, we look at an example with a bigger stencil, see Figure 4.1 and Table 5.2. The parameters are now 
\[ c_1 = 2, \quad f_1 = 6, \quad w_1 = w_2 = 0, \quad n_1 = 8, \quad f_w = 1, \quad s_1 = 1, \quad c_2 = 3, \quad f_2 = 12, \quad n_2 = 15, \quad s_2 = 1. \]

We clearly see that a larger stencil significantly increases the ratio of classical over direct interpolation, as well as that of distance-two over distance-one interpolations.

<table>
<thead>
<tr>
<th>Interpol.</th>
<th>Left example in Fig. 4.1</th>
<th>Right example in Fig. 4.1</th>
</tr>
</thead>
<tbody>
<tr>
<td>direct</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td>clas</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>std</td>
<td>17</td>
<td>12</td>
</tr>
<tr>
<td>ext</td>
<td>6</td>
<td>7</td>
</tr>
<tr>
<td>ext+i</td>
<td>10</td>
<td>9</td>
</tr>
</tbody>
</table>

Table 5.2

Cost for examples in Figure 4.1.

Table 5.3 shows the times for calculating these interpolation operators for matrices with stencils of various sizes. Two two-dimensional examples, one with a 5-point and one with a 9-point stencil, were examined on a 1000 × 1000 grid. The three-dimensional examples, with a 7-point and a 27-point stencil, were examined for an 80 × 80 × 80 grid. We have also included actual measurements of the average number of interpolatory points for these examples. As expected, larger stencils lead to a larger number of operations for each interpolation operator, with a much more significant increase for distance-two interpolation operators, particularly for the 3-dimensional problems. These effects are significant, especially since on coarser levels the stencils become larger and, thus, impact the total setup time.

<table>
<thead>
<tr>
<th>Stencil</th>
<th>(c_1)</th>
<th>(c_2)</th>
<th>direct</th>
<th>clas</th>
<th>std</th>
<th>ext</th>
<th>ext+i</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-pt</td>
<td>2.3</td>
<td>1.9</td>
<td>0.27</td>
<td>0.35</td>
<td>0.64</td>
<td>0.51</td>
<td>0.54</td>
</tr>
<tr>
<td>9-pt</td>
<td>1.8</td>
<td>2.8</td>
<td>0.36</td>
<td>1.11</td>
<td>2.16</td>
<td>2.09</td>
<td>2.48</td>
</tr>
<tr>
<td>7-pt</td>
<td>2.7</td>
<td>4.1</td>
<td>0.19</td>
<td>0.31</td>
<td>0.80</td>
<td>0.73</td>
<td>0.81</td>
</tr>
<tr>
<td>27-pt</td>
<td>2.3</td>
<td>7.2</td>
<td>0.40</td>
<td>3.72</td>
<td>8.00</td>
<td>7.43</td>
<td>8.32</td>
</tr>
</tbody>
</table>

Table 5.3

Average number of distance-one \((c_1)\) and distance-two \((c_2)\) interpolatory points and times for various interpolation operators.

6. Sequential Numerical Results. While the previous section examined the computational cost for the interpolation operator, we are of course mainly interested in the performance of the complete solver, which also includes coarsening, the generation of the coarse grid operator as well as the solve phase. We apply the new and old interpolation operators here to a variety of test problems from [7] to compare their efficiency. We did not include results using direct interpolation, since it performs worse than classical and multipass interpolation for the problems considered, nor
results using multipass interpolation followed by Jacobi interpolation, since these results were very similar to those obtained for 'clas+j'. All these tests were obtained using AMG as a solver with a strength threshold of $\alpha = 0.25$, and coarse-fine-Gauss-Seidel as a smoother. The iterations were stopped when the relative residual was smaller than $10^{-8}$.

<table>
<thead>
<tr>
<th>Method</th>
<th>$C_{op}$</th>
<th># its</th>
<th>time</th>
<th>$C_{op}$</th>
<th># its</th>
<th>time</th>
</tr>
</thead>
<tbody>
<tr>
<td>clas</td>
<td>1.92</td>
<td>244</td>
<td>151.60</td>
<td>1.24</td>
<td>157</td>
<td>100.44</td>
</tr>
<tr>
<td>clas+j</td>
<td>2.65</td>
<td>15</td>
<td>26.09</td>
<td>1.65</td>
<td>9</td>
<td>21.03</td>
</tr>
<tr>
<td>mp</td>
<td>1.92</td>
<td>244</td>
<td>152.34</td>
<td>1.24</td>
<td>183</td>
<td>115.72</td>
</tr>
<tr>
<td>ext</td>
<td>2.54</td>
<td>16</td>
<td>20.24</td>
<td>1.60</td>
<td>10</td>
<td>18.26</td>
</tr>
<tr>
<td>ext+i</td>
<td>2.57</td>
<td>11</td>
<td>16.93</td>
<td>1.60</td>
<td>10</td>
<td>18.40</td>
</tr>
<tr>
<td>std</td>
<td>2.56</td>
<td>16</td>
<td>20.63</td>
<td>1.60</td>
<td>17</td>
<td>23.06</td>
</tr>
</tbody>
</table>

**Table 6.1** AMG for the 5-point and the 9-point 2D Laplace problem on a 1000x1000 square with random right hand side using different interpolation operators.

<table>
<thead>
<tr>
<th>Method</th>
<th>$C_{op}$</th>
<th># its</th>
<th>time</th>
<th>$C_{op}$</th>
<th># its</th>
<th>time</th>
</tr>
</thead>
<tbody>
<tr>
<td>clas</td>
<td>1.90</td>
<td>168</td>
<td>38.60</td>
<td>1.82</td>
<td>&gt;1000</td>
<td></td>
</tr>
<tr>
<td>clas+j</td>
<td>2.39</td>
<td>29</td>
<td>10.50</td>
<td>3.40</td>
<td>424</td>
<td>131.85</td>
</tr>
<tr>
<td>mp</td>
<td>1.90</td>
<td>163</td>
<td>37.16</td>
<td>1.82</td>
<td>&gt;1000</td>
<td></td>
</tr>
<tr>
<td>ext</td>
<td>2.07</td>
<td>31</td>
<td>8.75</td>
<td>2.69</td>
<td>217</td>
<td>59.70</td>
</tr>
<tr>
<td>ext+i</td>
<td>2.07</td>
<td>11</td>
<td>4.05</td>
<td>2.89</td>
<td>97</td>
<td>29.78</td>
</tr>
<tr>
<td>std</td>
<td>2.07</td>
<td>13</td>
<td>4.53</td>
<td>2.89</td>
<td>148</td>
<td>43.68</td>
</tr>
</tbody>
</table>

**Table 6.2** AMG for a problem with a by 45 degrees and by 60 degrees rotated anisotropy on a 512x512 square using different interpolation operators.

Table 6.1 shows results for the two-dimensional Poisson problem $-\Delta u = f$ using a 5-point finite difference discretization and a 9-point finite element discretization. Table 6.2 shows results for the two-dimensional rotated anisotropic problem

$$-(c^2 + \epsilon_2 s^2)u_{xx} + 2(1-\epsilon)scu_{xy} - (s^2 + \epsilon^2)u_{yy} = 1$$

with $s = \sin \gamma$, $c = \cos \gamma$, and $\epsilon = 0.001$ with rotation angles $\gamma = 45^\circ$ and $\gamma = 60^\circ$.

The use of the distance-two interpolation operators combined with PMIS shows significant improvements over classical and multipass interpolation with regard to number of iterations as well as time. The best interpolation operator here is the ext+i interpolation, which has the lowest number of iterations and times in general. The difference is especially significant in the case of the problems with rotated anisotropies. The operator complexity is larger, however, as was to be expected.

This increase becomes more significant for three-dimensional problems. Here we consider the partial differential equation

$$-(a_{xx})x - (a_{xy})y - (a_{zz})z = f$$

on a $n \times n \times n$ cube. For the Laplace problem $a(x, y, z) = 1$, for the problem denoted by 'Jumps' we consider the function $a(x, y, z) = 1000$ for the interior cube $0.1 < \ldots < 1$.
Table 6.3

<table>
<thead>
<tr>
<th>Method</th>
<th>7-point C&lt;sub&gt;op&lt;/sub&gt;</th>
<th># its</th>
<th>time</th>
<th>27-point C&lt;sub&gt;op&lt;/sub&gt;</th>
<th># its</th>
<th>time</th>
<th>Jumps C&lt;sub&gt;op&lt;/sub&gt;</th>
<th># its</th>
<th>time</th>
</tr>
</thead>
<tbody>
<tr>
<td>clas</td>
<td>2.34</td>
<td>45</td>
<td>10.21</td>
<td>1.09</td>
<td>28</td>
<td>10.58</td>
<td>2.50</td>
<td>&gt;1000</td>
<td></td>
</tr>
<tr>
<td>clas+j</td>
<td>5.12</td>
<td>11</td>
<td>20.35</td>
<td>1.34</td>
<td>8</td>
<td>17.10</td>
<td>5.37</td>
<td>15</td>
<td>20.99</td>
</tr>
<tr>
<td>mp</td>
<td>2.35</td>
<td>47</td>
<td>10.40</td>
<td>1.10</td>
<td>30</td>
<td>9.39</td>
<td>2.50</td>
<td>80</td>
<td>17.37</td>
</tr>
<tr>
<td>ext</td>
<td>4.93</td>
<td>11</td>
<td>16.70</td>
<td>1.35</td>
<td>8</td>
<td>21.32</td>
<td>5.27</td>
<td>15</td>
<td>16.89</td>
</tr>
<tr>
<td>ext+i</td>
<td>4.27</td>
<td>9</td>
<td>14.48</td>
<td>1.35</td>
<td>8</td>
<td>21.55</td>
<td>5.10</td>
<td>11</td>
<td>15.96</td>
</tr>
<tr>
<td>std</td>
<td>4.20</td>
<td>10</td>
<td>12.78</td>
<td>1.38</td>
<td>10</td>
<td>18.58</td>
<td>5.21</td>
<td>18</td>
<td>17.47</td>
</tr>
</tbody>
</table>

AMG for a 7-point 3D Laplace problem, a problem with a 27-point stencil and a 3D structured PDE problem with jumps on a 60 × 60 × 60 cube with a random right hand side using different interpolation operators.

0.9, a(x, y, z) = 0.01 for 0 < x, y, z < 0.1 and the other cubes of size 0.1 × 0.1 × 0.1 that are located at the corners of the unit cube and a(x, y, z) = 1 elsewhere. The 27-point problem is a matrix with a 27-point stencil with the value 26 in the interior and -1 elsewhere and is being tested because we also wanted to consider a problem with a larger stencil.

While for these problems AMG convergence factors for distance-two interpolation improve significantly compared to classical and multi-pass interpolation, as can be seen in Table 6.3, overall times are worse for the 7pt 3D Laplace problem as well as the 27-point problem on a 60 × 60 × 60 grid. The only problem on the 60 × 60 × 60 grid that benefits from distance-two interpolation operators also with regard to time is the problem with jumps, which requires long-distance interpolation to even converge. Using distance-two interpolation operators leads to complexities about twice as large as those obtained when using classical or multi-pass interpolation, which work relatively well for the 7- and 27-point problem on the 60 × 60 × 60 grid. However, when we scale up the problem sizes, they show very good scalability in terms of AMG convergence factors, as can be seen in Figure 6.1, which shows the number of iterations for a 3D 7-pt Laplace problem on a n × n × n grid for increasing n. The anticipated large differences in numbers of iterations between distance-one and distance-two interpolations show up in the 2D results of Tables 6.1 and 6.2 on grids with 1000 points per direction, but are not particularly significant yet in the 3D results of Table 6.3 with only 60 points per direction. It is expected, however, that for the large problems that we want to solve on a parallel computer, distance-two interpolation operators will lead to overall better times than classical or multi-pass interpolation due to scalable AMG convergence factors, if the operator complexity can be kept under control. See Section 9 for actual test results.

7. Reducing Operator Complexity. While the methods described in the previous section largely restore grid-independent convergence to AMG cycles that use PMIS-coarsened grids, they also lead to much larger operator complexities for the V-cycles. Therefore, it is necessary to consider ways to reduce these complexities while (hopefully) retaining the improved convergence. In this section we describe a few ways of achieving this.

7.1. Choosing Smaller Interpolatory Sets. It is certainly possible to consider other interpolatory sets, which are larger than C<sub>i</sub><sup>s</sup>, but smaller than C<sub>i</sub>. Particularly, it appears that a good interpolatory set would be one that only extends C<sub>i</sub><sup>s</sup> for strong F – F connections without a common C-point, since in the other cases
point \(i\) is likely already surrounded by interpolatory points and an extension is not necessary. If we look at the right example in Figure 4.1, we see that neighbor \(k\) of \(i\) is the only fine neighbor that does not share a \(C\)-point with \(i\). Consequently, it may be sufficient to only include points \(n\) and \(l\) in the extended interpolatory set. Applying this approach to the extended interpolation leads to so-called \(F-F\) interpolation \([3]\). The size of the interpolatory set can be further decreased if we limit the number of \(C\)-points added when an \(F\)-point is encountered that does not have common coarse neighbors. This has been done in the so-called \(F-F1\) interpolation \([3]\), where only the first \(C\)-point is added. For the right example in Figure 4.1 this means that only point \(n\) or point \(l\) would be added to the interpolatory set for \(i\).

Choosing a smaller interpolatory set decreases \(c_2\) and with it \(s_2\), leading to fewer multiplications and additions for the extended interpolation methods. On the other hand, additional operations are needed to determine which coarse neighbors of strong \(F\)-points are common \(C\)-points. This means that the actual determination of the interpolation operator might not be faster than creating the extended interpolation operators. The real benefit, however, is achieved by the fact that the use of smaller interpolatory sets leads to smaller stencils for the coarse grid operator and hence to smaller overall operator complexities.

Applying these methods to some of our previous test problems, we get the results shown in Tables 7.1 and 7.2. Here, ‘x-cc’ denotes that interpolation ‘x’ is used, but the interpolatory set is only extended when there are no common \(C\)-points. Similarly, ‘x-ccs’ is just like ‘x-cc’, except that for every strong \(F\)-point without a common \(C\)-point only a single \(C\)-point is added.

The results show that 2-dimensional problems do not benefit from this strategy, since operator complexities are only slightly decreased, while the number of iterations increases. Therefore total times increase. However, 3-dimensional problems can be solved much faster due to significantly decreased setup times leading to only half the total times when the ‘ccs’-strategy is employed. Again, these beneficial effects are expected to be stronger on larger grids. Indeed, additional numerical tests (not presented here, see [3]) also show that the ‘x-cc’ and ‘x-ccs’ distance-two interpolations result in algorithms that are highly scalable as a function of problem size: \(C_{op}\) tends

![Fig. 6.1. Number of iterations for PMIS with various interpolation operators for a 3D 7-point Laplace problem on a \(n \times n \times n\)-grid.](image-url)
to a constant that is significantly smaller than the \(C_{op}\) value for the ‘x’ interpolations, and the number of iterations is nearly constant as a function of problem size, and only slightly larger than the number of iterations for the full ‘x’ interpolation formulas [3]. This shows that using distance-two interpolation formulas with reduced complexities restores the grid-independent convergence and scalability of AMG on PMIS-coarsened grids, without the need for GMRES acceleration. This makes these methods suitable algorithms for large problems on parallel computers, as is discussed below.

7.2. Interpolation Truncation. Another very effective way to reduce complexities is the use of interpolation truncation. There are essentially two ways we can truncate interpolation operators: we can choose a truncation factor \(\theta\) and eliminate every weight whose absolute value is smaller than this factor, i.e. for which \(|w_{ij}| < \theta\) [12], or we can limit the number of coefficients per row, i.e. choose only the \(k_{max}\) largest weights in absolute value. In both cases the new weights need to be re-scaled so that the total sums remain unchanged.

Both approaches can lead to significant reductions in setup times and operator complexities, particularly for 3-dimensional problems, but if too much is truncated, the number of iterations rises significantly, as one would expect.

We only report results for one interpolation formula (ext+i) for a 3D example here, see Table 7.3. However, similar results can be obtained using the other interpolation operators. For 2-dimensional problems, truncation leads to an increase in total time similarly as reported for interpolatory set restriction in the previous section.

8. Parallel Implementation. This section describes the parallel implementation and gives a rough idea of the cost involved, with particular focus on the increase
in communication required for the distance-two interpolation formulae compared to
distance-one interpolation. Since the core computation for the interpolation routines
is approximately the same as in the sequential case, we only focus on the additional
computations that are required for inter-communication between processors.

In parallel, each matrix is stored using a parallel data format, the ParCSR ma-
trix data structure, which is described and analyzed in detail in [8]. Matrices are
distributed across processors by contiguous blocks of rows, which are stored via two
compressed sparse row matrices, one storing the local entries and the other one storing
the off-processor entries. There is an additional array containing a mapping for the
off-processor neighbor points. The data structure also contains the information nec-
essary to retrieve information from distance-one off-processor neighbors. It however
does not contain information on off-processor distance-two neighbors, which com-
plicates the parallel implementation of distance-two interpolation operators. When
determining these neighbors, there are four scenarios that need to be considered, see
Figure 8.1. Consider point $i$, which is the point to be interpolated to, and is residing
on Processor 0. A distance-two neighbor can reside on the same processor as $i$, like
point $j$; it can be a distance-one neighbor to another point on Proc. 0, like point $l$,
and therefore be already contained in the off-processor mapping; it can be a new
point on a neighbor processor, like point $k$, or it can be located on a processor, which
is currently not a neighbor processor to Proc. 0, like point $m$.

![Fig. 8.1. Example of off-processor distance-two neighbors of point i. Black points are C-points, and white points are F-points.](image)

There are basically five additional parts that are required for the parallel im-
plementation, and for which we give rough estimates of the cost involved below.
Operations include floating point and integer operations as well as message passing and sends and receives required to communicate data across processors. We use the following notations: \( n_1 \) denotes the average number of distance-one neighbors per point, as defined previously, \( p \) is the total number of processors, \( q_i \) is the average number of distance-\( i \) neighbor processors per processor, \( N^o_i \) is the average number of distance-\( i \) off-processor points and equals the sum of the average number of distance-\( i \) off-processor \( C \)-points, \( C^o_i \), and distance-\( i \) off-processor \( F \)-points, \( F^o_i \). Note that the estimates of number of operations and number of processors involved given below are per processor.

1. Communication of C/F splitting for all off-processor neighbor points: This is required for all interpolation operators and takes \( O(N^o_1) + O(q_1) \) operations.

2. Obtaining the row information for off-processor distance-one \( F \)-points: This step is necessary for classical and distance-two interpolation, but not for direct interpolation, which only uses local matrix coefficients to generate the interpolation formula. It requires \( O(n_1 F^o_1 \log(N^o_1)) \) operations.

3. Determining off-processor distance-two points and additional communication information: This step is only required for distance-two interpolation operators. Finding the new off-processor points, which requires checking whether they are already contained in the map and describing the off-processor connections, takes \( O(n_1 F^o_1 \log(N^o_1)) \) operations. Sorting the new information takes \( O(N^o_2 \log(N^o_2)) \) operations. Getting the communication information for the new points using an assumed partition algorithm \([1]\), requires \( O(N^o_2) + O(\log p) + O((q_1 + q_2) \log(q_1 + q_2)) \) operations. Getting the additional C/F splitting information takes \( O(N^o_2) + O(q_1 + q_2) \) operations.

4. Communication of fine_to_coarse mappings: This step requires \( O(N^o_1) + O(q_1) \) operations for distance-one interpolation and \( O(N^o_1 + N^o_2) + O(q_1 + q_2) \) operations for the distance-two interpolation schemes.

5: Generating the interpolation matrix communication package: This step requires \( O(C^o_1) + O(\log p) + O(q_1 \log q_1) \) operations for distance-one interpolation and \( O(C^o_1 + C^o_2) + O(\log p) + O((q_1 + q_2) \log(q_1 + q_2)) \) operations for distance-two interpolation. Note that if truncation is used, \( C^o_i \) should be replaced by \( \hat{C}^o_i \) with \( \hat{C}^o_i < C^o_i \) for \( i = 1, 2 \).

Summarizing these results, direct interpolation requires the least amount of communication, followed by classical interpolation. Parallel implementation of distance-two interpolation requires more communication steps and additional data manipulation, and involves more data and neighbor processors. How significantly this overhead impacts the total time depends on many factors, such as the problem size per processor, the stencil size, the computer architecture and more. Parallel scalability results are presented below.

9. Parallel Numerical Results. In this section, we investigate weak scalability of the new interpolation operators by applying the resulting AMG methods to various problems.

The following problems were run on Thunder, an Intel Itanium2 machine with 1024 nodes of 4 processors each, located at Lawrence Livermore National Laboratory, unless we say otherwise. In this section, \( p \) denotes the number of processors used.

9.1. Two-Dimensional Problems. We first consider two-dimensional Laplace problems, which perform very poorly for PMIS with classical interpolation. The results we obtained for 5-point and 9-point stencils are very similar. Therefore we list only the results for the 9-point 2D Laplace problem here.
Table 9.1, which contains the number of iterations and total times for this problem, show that classical interpolation performs very poorly, and multipass interpolation even worse. Nevertheless, these methods lead to the lowest operator complexities: 1.24. All long-range interpolation schemes lead to good scalable convergence, with standard-interpolation performing slightly worse than classical followed by Jacobi, extended or extended+i interpolation, which are the overall fastest methods here with the best scalability. Operator complexities are highest for clas+j with 1.65 and about 1.6 for the other three interpolation operators. Also, when choosing the lower complexity versions e+i-cc and e+i-ccs, with complexities of 1.45 and 1.43, convergence deteriorates somewhat compared to e+i. Since for the 2-dimensional problems setup times are fairly small and the improvement in complexities is not very significant, this increase in number of iterations hurts the total times, and therefore there is no advantage in using low-complexity schemes for this problem. Truncated versions lead to even larger total times.

<table>
<thead>
<tr>
<th></th>
<th>clas</th>
<th>clas+j</th>
<th>mp</th>
<th>std</th>
<th>ext</th>
<th>e+i</th>
<th>e+i-cc</th>
<th>e+i-ccs</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>15(88)</td>
<td>3(9)</td>
<td>18(105)</td>
<td>4(15)</td>
<td>3(10)</td>
<td>3(10)</td>
<td>3(12)</td>
<td>3(13)</td>
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<tr>
<td>64</td>
<td>48(245)</td>
<td>4(11)</td>
<td>57(278)</td>
<td>6(20)</td>
<td>4(12)</td>
<td>4(12)</td>
<td>5(16)</td>
<td>5(19)</td>
</tr>
<tr>
<td>256</td>
<td>79(400)</td>
<td>5(12)</td>
<td>85(436)</td>
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<td>6(21)</td>
</tr>
<tr>
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<td>6(13)</td>
<td>n.c.</td>
<td>9(27)</td>
<td>6(14)</td>
<td>6(14)</td>
<td>7(21)</td>
<td>7(21)</td>
</tr>
</tbody>
</table>

**Table 9.1**

Times in seconds (number of iterations) for a 9-point 2D Laplace problem with 300 x 300 points per processor; ‘n.c.’ denotes ‘not converging within 500 iterations’.

Next, we consider the 2-dimensional problem with rotated anisotropy (6.1). The first problem here has an anisotropy of 0.001 rotated by 45 degrees, see Table 9.2. Operator complexities for classical and multipass interpolation are here 1.9; they are 2.4 for classical interpolation followed by Jacobi, and 2.1 for all remaining interpolation operators. Here, extended interpolation performs worse than standard and extended+i interpolation, which gives the best results.

<table>
<thead>
<tr>
<th></th>
<th>clas</th>
<th>clas+j</th>
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<th>ext</th>
<th>e+i</th>
<th>e+i-cc</th>
<th>e+i-ccs</th>
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<td>3(12)</td>
<td>3(12)</td>
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<td>11(39)</td>
<td>5(16)</td>
<td>7(21)</td>
<td>7(23)</td>
</tr>
<tr>
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<td>n.c.</td>
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<td>12(42)</td>
<td>6(18)</td>
<td>8(25)</td>
<td>8(27)</td>
</tr>
<tr>
<td>1024</td>
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<td>n.c.</td>
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<td>14(45)</td>
<td>8(21)</td>
<td>10(29)</td>
<td>11(31)</td>
</tr>
</tbody>
</table>

**Table 9.2**

Times in seconds (number of iterations) for a 2D problem with a 45 degrees rotated anisotropy of 0.001 with 300 x 300 points per processor; ‘n.c.’ denotes ‘not converging within 500 iterations’.

In Table 9.3 we consider the harder problem, where the anisotropy is rotated by 60 degrees. Operator complexities are now 1.8 for classical and multipass, 3.4 for clas+j, 2.9 for e+i and std, 2.7 for ext, 2.8 for e+i-cc and 2.5 for e+i-ccs. Here fastest convergence is obtained for the extended+i interpolation, followed by e+i-cc, e+i-ccs, std, and ext. The other interpolations fail to converge within 500 iterations. While long-range interpolation operators improve convergence, it is still not good enough, so this problem should be solved using Krylov subspace acceleration.

9.2. Three-Dimensional Structured Problems. We now consider 3-dimensional problems. Based on the sequential results in Section 6 we expect complexity reduction
schemes to make a difference here.

The first problem is a 7-point 3D Laplace problem on a structured cube with 40 × 40 × 40 unknowns per processor. Table 9.4 shows total times in seconds, and number of iterations. While classical interpolation solves the problem, the number of iterations increases rapidly with increasing number of processors and problem size. Multipass interpolation performs better for larger number of processors, but still shows unscaable convergence factors. Applying one step of Jacobi interpolation to classical interpolation leads to perfect scalability in terms of convergence factors, but unfortunately also to rising operator complexities (4.9-5.7), which are twice as large as for classical and multipass interpolation (2.3-2.4). Interestingly, while both standard and extended+i interpolation need less iterations for a small number of processors than extended interpolation, they show worse numerical scalability leading to far less iterations for extended interpolation for large number of processors. However extended interpolation leads to larger complexities (4.7-5.3) compared to extended+i (4.2-4.5) and standard interpolation (4.1-4.4). The complexity reducing strategies lead to the following complexities: ext-cc (4.5-4.9), ext-ccs (3.9-4.2), e+i-cc (4.0-4.3), and e+i-ccs (3.6-3.8). For the sake of saving space, we did not record the results for ext-cc or e+i-cc, but the times and number of iterations for these methods was in between those of ext and ext-ccs, or e+i and e+i-ccs, respectively. Interestingly, the complexity reducing strategies e+i-cc and e+i-ccs show not only better scalability with regard to time, but also better scalability of convergence factors than e+i interpolation in this case.

<table>
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<tr>
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<th>std</th>
<th>ext</th>
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<th>e+i-cc</th>
<th>e+i-ccs</th>
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</tr>
<tr>
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<td>n.c.</td>
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<td>96(330)</td>
<td>47(152)</td>
<td>50(196)</td>
<td>70(254)</td>
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<tr>
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<td>n.c.</td>
<td>95(305)</td>
<td>110(374)</td>
<td>56(176)</td>
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<td>84(299)</td>
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<tr>
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<td>n.c.</td>
<td>n.c.</td>
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<td>123(408)</td>
<td>62(193)</td>
<td>82(263)</td>
<td>100(347)</td>
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</tr>
</tbody>
</table>

*Table 9.3: Times in seconds (number of iterations) for a 2D problem with a by 60 degrees rotated anisotropy of 0.001 with 300 × 300 points per processor; ‘n.c.’ denotes ‘not converging within 500 iterations’.*

For this problem, complexity reducing strategies, thus, are paying off. Table 9.5 shows results for various truncated interpolation schemes. We used the truncation strategy which restricts the number of weights per row using either 4 or 5 for the maximal number of elements. While we present both results for ext and e+i, we present only the faster results for the remaining interpolation schemes for the sake of saving space. We used a truncation factor of 0.1 for clas+j. Operator complexities

<table>
<thead>
<tr>
<th>$p$</th>
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<th>clas+j</th>
<th>mp</th>
<th>std</th>
<th>ext</th>
<th>e+i</th>
<th>ext-ccs</th>
<th>e+i-ccs</th>
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<td>16(79)</td>
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</tr>
<tr>
<td>512</td>
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<td>26(12)</td>
<td>28(126)</td>
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<td>20(14)</td>
<td>17(15)</td>
<td>11(14)</td>
<td>11(11)</td>
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<tr>
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<td>39(175)</td>
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<td>30(13)</td>
<td>31(39)</td>
<td>15(15)</td>
<td>16(14)</td>
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<tr>
<td>1728</td>
<td>51(229)</td>
<td>63(12)</td>
<td>37(159)</td>
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<td>40(33)</td>
<td>22(15)</td>
<td>24(16)</td>
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</tbody>
</table>

*Table 9.4: Total times in seconds (number of iterations) for a 7 point 3D Laplace problem with 40 × 40 × 40 points per processor.*
were fairly consistent here across increasing numbers of processors: we obtained 2.9 for ext4, 3.2 for ext5, 2.8 for e+i4, 3.1 for e+i5, 3.2 for ext-cc5, 3.1 for e+i-cc5, 3.2 for std5 and 3.0 for clas+j0.1. Clearly, using four compared to five weights leads to lower complexities, but larger number of iterations. Total times are not significantly different. Comparing the fastest method, e+i-cc5, on 1728 processors to PMIS with classical interpolation, we see a factor of 11 in improvement with regard to number of iterations and a factor of 5 in improvement with regard to total time with a slight increase in complexity.

Table 9.6 shows results for the problem with jumps (6.2), for which PMIS with classical interpolation was shown to completely fail. Multipass interpolation converges here with highly degrading scalability but good complexities of 2.4. Applying Jacobi interpolation to classical interpolation leads to very good convergence, but, due to operator complexities between 5.1 and 5.7, it leads to a much more expensive setup and solve cycle. Applying a truncation factor of 0.1 as in the previous example leads to extremely bad convergence and is not helpful here. Standard interpolation converges very well for small number of processors, but diverges if $p$ is greater or equal to 64. Interestingly enough std4 converges, albeit not very well.

### Table 9.5

<table>
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<tr>
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<td>11(20)</td>
<td>28(46)</td>
<td>19(45)</td>
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</table>

Total times in seconds (number of iterations) for a 7 point 3D Laplace problem with $40 \times 40 \times 40$ points per processor.

### Table 9.6

<table>
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<th>ext-ccs</th>
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</tr>
<tr>
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<td>73(70)</td>
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<td>25(29)</td>
<td>29(23)</td>
<td>53(169)</td>
<td>17(36)</td>
</tr>
</tbody>
</table>

Total times in seconds (number of iterations) for a structured 3D problem with jumps with $40 \times 40 \times 40$ points per processor.

#### 9.3. Unstructured Problems.

In this section, we consider various linear systems on unstructured grids that have been generated by finite element discretizations. All of these problems were run on an Intel Xeon Linux cluster at Lawrence Livermore National Laboratory. The first problem is the three-dimensional diffusion problem

$$-a_1(x, y, z)u_{xx} - a_2(x, y, z)u_{yy} - a_3(x, y, z)u_{zz} = f$$

with Dirichlet boundary conditions on an unstructured cube. The material properties are discontinuous, and there are approximately 90,000 degrees of freedom per processor. See Figure 9.1 for an illustration of the grid used. There are five regions: four layers and the thin stick in the middle of the domain. This grid is further refined when larger number of processors are used. The functions $a_i(x, y, z), i = 1, 2, 3$ are constant within each of
the five regions of the domains with the following values \((4,0.2,1,1,10^4)\) for \(a_1(x,y,z)\), \((1.0, 2.3, 1.0^4)\) for \(a_2(x,y,z)\), and \((1.0, 0.01, 1,1,10^4)\) for \(a_3(x,y,z)\). We also include some results obtained with CLJP coarsening, which is a parallel coarsening scheme that was designed to ensure that two fine neighbors always have a common coarse neighbor and for which classical interpolation is therefore suitable [5, 9]. We include only results for the extended+i interpolation here, because the extended interpolation performs very similar. As a smoother we used hybrid Gauss-Seidel, which leads to a nonsymmetric preconditioner. Since in practice more complicated problems are usually solved using AMG as a preconditioner for Krylov subspace methods, we use AMG here as a preconditioner for GMRES\((10)\). Note that both classical and multipass interpolation do not converge within 1000 iterations for these problems if they are used without a Krylov subspace method, whereas both extended and extended+i interpolation, as well as classical interpolation on CLJP-coarsened grids, converge well without it, with a somewhat larger number of iterations and slightly slower total times.

Fig. 9.1. Grid for the elasticity problem.

The results in Table 9.7 show that the long-range interpolation operators, with the exception of multipass interpolation, restore the good convergence that was obtained with CLJP. CLJP has very large complexities, however. We also used a truncated version of classical interpolation, restricting the number of weights per fine point to at most 4 to control the complexities. While this hardly affected convergence factors, it significantly improved the total times to solution, see Figure 9.2, but still did not achieve perfect scalability. Total times for CLJP with clas4 interpolation are comparable to PMIS with classical interpolation due to the small complexities of PMIS in spite of its significantly worse convergence factors. The use of extended+i and e+i-cc interpolation leads to better scalability than the methods mentioned before due to their lower complexities if compared to CLJP, or their better convergence factors if compared to PMIS with classical interpolation. Multipass interpolation leads to even better timings, but the overall best time and scalability is achieved by applying truncation to 4 weights per fine point to extended+i interpolation.

The second problem is a 3-dimensional linear elasticity problem using the same domain as above. However, a smaller grid size is used, since this problem requires more
memory, leading to about 30,000 degrees of freedom per processor. The Poisson ratio chosen for the pile driver in the middle of the domain was chosen to be 0.4, the Poisson ratios in the surrounding regions were 0.1, 0.3, 0.3 and 0.2. Since this is a systems problem the unknown-based systems AMG method was used. For this problem, the conjugate gradient method was used as an accelerator, and hybrid symmetric Gauss-Seidel as a smoother. The results are given in Table 9.8 and Figure 9.3. Here also extended+i interpolation leads to the lowest run times and best scalability. CLJP ran out of memory for the 512 processor run.

### 10. Conclusions

We have studied the performance of AMG methods using the PMIS coarsening algorithm in combination with various interpolation operators. PMIS with classical, distance-one interpolation leads to an AMG method with low complexity, but has bad scalability in terms of AMG convergence factors. The use of distance-two interpolation operators restores this scalability. However, it leads to an increase in operator complexity. While this increase was fairly small for two-dimensional problems and was far outweighed by the much improved convergence, for three-dimensional problems complexities were often twice as large, and impacted scalability. To counter this complexity growth, we implemented various complexity...
Distance-two interpolation for parallel AMG

<table>
<thead>
<tr>
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<tr>
<td>C_{op}</td>
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<td>3.6-5.4</td>
</tr>
</tbody>
</table>

Table 9.8

Number of iterations for the 3D elasticity problem; range of operator complexities. AMG is used here as a preconditioner for conjugate gradient.

Reducing strategies, such as the use of smaller interpolatory sets and interpolation truncation. The resulting AMG methods, particularly the extended and extended+i interpolation in combination with truncation, lead to very good scalability for a variety of difficult PDE problems on large parallel computers.

Acknowledgments. We thank Tzanio Kolev for providing the unstructured problem generator and Jeff Painter for the Jacobi interpolation routine. This work was performed under the auspices of the U.S. Department of Energy by University of California Lawrence Livermore National Laboratory under contract No. W-7405-Eng-48.

REFERENCES


