

# A ROOT-NODE BASED ALGEBRAIC MULTIGRID METHOD\*

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## Abstract.

This paper provides a unified and detailed presentation of root-node style algebraic multigrid (AMG). Algebraic multigrid is a popular and effective iterative method for solving large, sparse linear systems that arise from discretizing partial differential equations. However, while AMG is designed for symmetric positive definite matrices (SPD), certain SPD problems, such as anisotropic diffusion, are still not adequately addressed by existing methods. Non-SPD problems pose an even greater challenge, and in practice AMG is often not considered as a solver for such problems.

The focus of this paper is on so-called root-node AMG, which can be viewed as a combination of classical and aggregation-based multigrid. An algorithm for root-node is outlined and a filtering strategy is developed, which is able to control the cost of using root-node AMG, particularly on difficult problems. New theoretical motivation is provided for root-node and energy-minimization as applied to symmetric as well non-symmetric systems. Numerical results are then presented demonstrating the robust ability of root-node to solve non-symmetric problems, systems-based problems, and difficult SPD problems, including strongly anisotropic diffusion, convection-diffusion, and upwind steady-state transport, in a scalable manner. New, detailed estimates of the computational cost of the setup and solve phase are given for each example, providing additional support for root-node AMG over alternative methods.

**Key words.** multigrid, algebraic multigrid, root-node, energy minimization, interpolation smoothing, anisotropic diffusion

**AMS subject classifications.** 65F10, 65M22, 65M55

**1. Introduction.** Algebraic multigrid (AMG) methods such as classical AMG (CF AMG<sup>1</sup>) [11, 42] and smoothed aggregation (SA AMG) [48] are efficient solution techniques for large, sparse linear systems. Algebraic multigrid was developed specifically for symmetric positive-definite (SPD) systems that arise from the discretization of elliptic partial differential equations (PDEs), and software packages such as BoomerAMG [28] in the hypre library [26] demonstrate its parallel scalability to hundreds of thousands of cores [3].

AMG targets solving a sparse linear system (typically SPD)

$$(1) \quad Ax = b$$

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<sup>1</sup>Classical AMG or so-called Ruge-Stüben AMG use a splitting of the degrees-of-freedom (DOFs) into coarse *C-points* and fine *F-points*, leading to the abbreviation CF.

with  $O(n)$  work, where  $\mathbf{x}, \mathbf{b} \in \mathbb{R}^n$  and  $A \in \mathbb{R}^{n \times n}$ . This optimality is achieved through two complementary parts of a multigrid method, relaxation and coarse-grid correction, which together uniformly damp error of all frequencies (see Section 2).

However, there are certain SPD systems and many non-symmetric systems for which AMG continues to struggle. Problems with strongly anisotropic components, and problems arising in particle transport, advective flow calculations, and strongly varying material properties, among others, challenge the standard approaches to AMG, thus highlighting the need for more robust methods. There have been a number of efforts in recent years to improve the convergence and scope of applicability of AMG. Adaptive methods focus on improving the multigrid hierarchy through trial cycles in the setup phase [17, 22, 16, 18]. Other methods focus on modified or improved strength measures when forming coarse grids [39, 22, 9, 36, 10, 13, 12, 7, 15, 31]. Furthermore, generalizing interpolation through energy minimization [41] and other methods [27, 20, 23, 50] is used to improve the accuracy of interpolation between grid levels. Nevertheless, many problems remain difficult for AMG to solve, while many ‘robust’ AMG methods suffer from high computational cost.

An AMG solver consists of a hierarchy of matrices,  $\{A_\ell\}$ , with the initial matrix on level  $\ell = 0$ ,  $A_0 := A$ , and progressively smaller matrices on levels  $\ell = 1, 2, \dots, L$ . Interpolation and restriction operators, also known as transfer operators, transfer vectors between different levels of the hierarchy. For a given matrix,  $A_\ell$ , the next ‘coarser’ matrix in the hierarchy (level  $\ell + 1$ ) is generally developed in one of two different ways: using a CF-splitting of points (CF AMG) or using an aggregation of points (SA AMG). A CF-splitting splits the set of all DOFs of matrix  $A_\ell$  into a coarse set of C-points and a fine set of F-points, with C-points corresponding to DOFs on the coarse grid. Transfer operators are then defined using the CF-splitting, where values at C-points are restricted and interpolated by injection and values at F-points use a linear combination of connected neighboring points. In SA AMG, a measure of the strength-of-connection (SOC) between nodes is used to form ‘aggregates’, which are disjoint sets of strongly connected nodes, where each aggregate represents one node on the coarse grid, and transfer operators are formed based on aggregates. A more detailed look at SA AMG and CF AMG is given in Section 2.

Root-node AMG (RN AMG) uses a hybrid approach, wherein SA-type strength-of-connection and aggregation routines are used to form aggregates. In each aggregate, one node is chosen to be the ‘root-node’, which corresponds to a C-point, and other nodes are designated as F-points. Transfer operators are then formed based on this CF-splitting together with aggregation.

Root-node AMG was initially identified in [41] as a small part of a general, energy-minimizing framework to form interpolation operators in AMG. The work in [45] implemented the RN approach and demonstrated its potential as an effective and scalable solver for strongly anisotropic, non-grid-aligned diffusion operators — problems which have proven difficult for other multigrid methods. However, certain problems required a large computational cost, especially in the setup of the method.

This paper provides, for the first time, a unified and detailed presentation of RN AMG and how it combines many of the benefits of CF AMG and SA AMG. Root-node AMG allows for classical point-wise decisions in the setup, to help control complexity<sup>2</sup> and provide theoretical motivation, as well as aggregation-style construction, to facilitate the design of a multigrid solver based on the spectral behavior of the

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<sup>2</sup>“Complexity” or “cost” refer to the overall computational cost of the method in terms of floating point operations.

problem. New theoretical motivation is given for the pairing of RN AMG with energy minimization, in both the symmetric and non-symmetric cases, proving the equivalence of the energy-minimization process to minimizing the difference with an optimal form of interpolation. Moreover, a new interpolation filtering strategy is developed to limit the complexity of the method, which proves critical for problems that require large sparsity patterns in transfer operators. Last, a numerical survey is presented to highlight the robustness and flexibility of RN AMG in comparison to CF AMG and SA AMG, including scalable convergence for strongly anisotropic diffusion problems and a discontinuous, upwind discretization of the steady-state transport equation. For each test problem, a detailed measure of computational cost or complexity is provided, a novel addition to AMG literature, which provides a complete picture of a method when coupled with convergence factors.

In Section 2, background information on algebraic multigrid methods is discussed, including current limitations and the basic motivation for a RN-type algorithm. The RN AMG method and algorithm are presented in Section 3, along with a discussion of computational complexity and a filtering process proposed to address cases of high complexity. Section 4 provides theoretical motivation for RN AMG. Numerical results are provided in Section 5, and conclusions and future work discussed in Section 6.

**2. Background.** Multigrid methods, such as SA AMG and CF AMG, involve two phases: (i) the setup phase, where a multilevel solver hierarchy is constructed, and (ii) the solve phase, where the constructed solver hierarchy is applied to solve the linear system (1) to a desired tolerance. Smoothed-aggregation AMG and CF AMG are distinguished by the setup phase; that is, once a hierarchy is constructed, both methods execute the solve phase in the same fashion.

In the following, a multigrid hierarchy consists of a set of matrices, starting with an initial, fine matrix  $A_0 \equiv A \in \mathbb{R}^{n \times n}$ . Matrices for additional levels in the hierarchy,  $A_\ell \in \mathbb{R}^{n_\ell \times n_\ell}$ , are then constructed based on interpolation and restriction operators,  $P_\ell : \mathbb{R}^{n_{\ell+1}} \rightarrow \mathbb{R}^{n_\ell}$  and  $R_\ell : \mathbb{R}^{n_\ell} \rightarrow \mathbb{R}^{n_{\ell+1}}$ , respectively, via  $A_{\ell+1} = R_\ell A_\ell P_\ell$ , where  $n_{\ell+1} < n_\ell$ . In the case of an SPD matrix, restriction  $R = P^T$ .

The AMG solve phase iterates using two complementary parts: relaxation — e.g., weighted Jacobi — to reduce the high-energy error that is associated with large eigenvalues in the operator, and coarse-grid correction, which targets algebraically smooth error,  $A_0 \mathbf{e} \approx 0$ , associated with the small eigenvalues. A two-grid solve proceeds as follows. Pre-relaxation on  $A_0 \mathbf{x}_0 = \mathbf{b}_0$  is applied, and the resulting residual is then restricted to the coarse grid,  $\mathbf{b}_1 = R_0(\mathbf{b}_0 - A_0 \mathbf{x}_0)$ , which serves as the right-hand side for the coarse-grid equation,  $A_1 \mathbf{x}_1 = \mathbf{b}_1$ , where  $A_1 = R_0 A_0 P_0$ . The solution,  $\mathbf{x}_1$ , provides a coarse-grid error correction that is interpolated back to the fine grid,  $\mathbf{x}_0 \leftarrow \mathbf{x}_0 + P_0 \mathbf{x}_1$ . Last, post-relaxation is applied to the updated  $\mathbf{x}_0$ . Together, these three steps form a two-level multigrid  $V(\nu_{\text{pre}}, \nu_{\text{post}})$ -cycle, where  $\nu_{\text{pre}}$  refers to the number of pre-relaxations and  $\nu_{\text{post}}$  to the number of post-relaxations. A full solve then consists of using successive V-cycles to iterate on a vector until the relative residual norm,  $\|\mathbf{r}_0\|/\|\mathbf{b}_0\|$  is less than some tolerance — e.g.,  $10^{-8}$ .

The effectiveness of this complementary process is explained by considering the corresponding two-grid error propagation operator. Let  $\mathbf{e}^{(0)}$  be the initial error in approximating the solution to (1) and let  $\mathbf{e}^{(0)} \leftarrow G \mathbf{e}^{(0)}$  represent the error propagator for the relaxation method — e.g.,  $G = I - \omega D^{-1} A$  in the case of weighted Jacobi. Then (dropping subscripts), the error after a two-grid cycle is given by

$$(2) \quad \mathbf{e}^{(1)} \leftarrow G^{\nu_2} \left( I - P(RAP)^{-1} RA \right) G^{\nu_1} \mathbf{e}^{(0)},$$

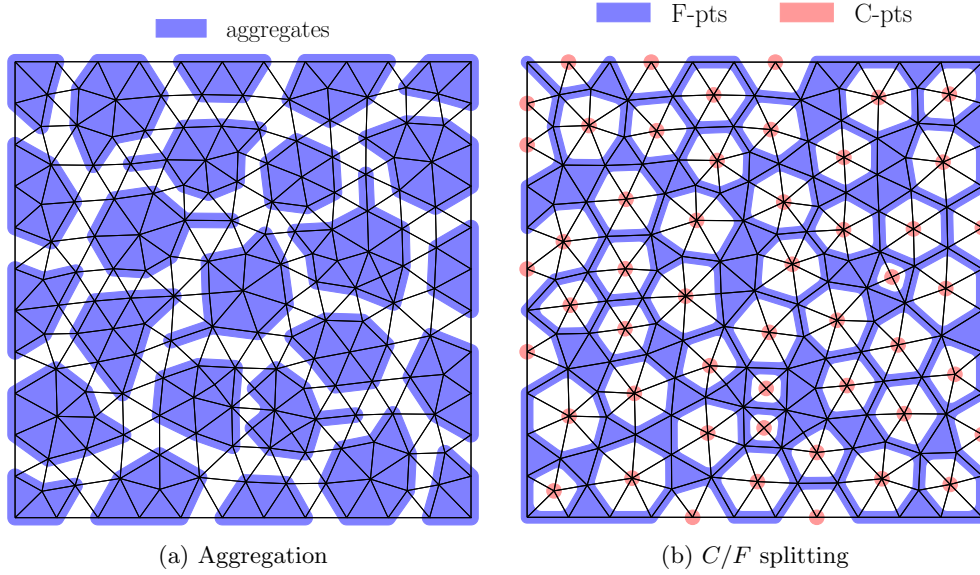


Fig. 1: Example SA AMG and CF AMG coarsening for a linear finite element approximation to a Laplace operator. The fine level problem has 191 DOFs. In this example, aggregation yields 25 coarse DOFs (aggregates), while a CF-splitting yields 51 coarse DOFs (C-points).

where  $P(RAP)^{-1}RA$  is a projection onto  $\mathcal{R}(P)$ . In the case of SPD  $A$ ,  $R = P^T$ , and this is an  $A$ -orthogonal projection. In either case, if  $Ge^{(0)} \in \mathcal{R}(P)$ , then the iteration is exact. In other words, if interpolation is complementary and accurate for modes not effectively reduced by relaxation, then error reduction with (2) will be large.

Algebraic multigrid methods attempt to automatically determine interpolation and coarse-grid operators ( $P_\ell$  and  $A_\ell$ ) that yield optimal error reduction with (2). The two standard approaches are SA AMG and CF AMG, which are outlined in Sections 2.1 and 2.2, respectively. Broadly, SA AMG defines a coarse problem through an *aggregation* of nodes (see Figure 1a), while CF AMG defines a coarse problem through a splitting of the DOFs into coarse C-points and fine F-points (see Figure 1b). Each offer advantages as noted in the following descriptions.

**2.1. AMG based on smoothed aggregation.** The effectiveness of SA AMG relies on *a priori* knowledge of algebraically smooth error in the form of candidate vectors,  $B$ . These vectors generally represent the lowest energy modes of the governing PDE with no boundary conditions — e.g., the constant for diffusion and rigid-body-modes for elasticity [48]. Using  $B$  (possibly determined *a priori*), the setup phase first uses a strength measure on the connectivity of nodes to define a SOC matrix,  $S$ , which is used to identify so-called aggregates (see Figure 1a). The goal of the strength measure is to ensure that algebraically smooth error at each DOF in an aggregate strongly correlates with algebraically smooth error at other DOFs in that aggregate. This holds for model problems, where algebraically smooth error over each aggregate is well represented by the restriction of the candidate vectors to the aggregate. Consequently, this *injection* of the candidate vectors over each aggregate is

what leads to the initial representation of interpolation, termed the tentative interpolation operator  $T$ , with candidate vectors exactly in the range of  $T$ . Each aggregate corresponds to one block in the block-diagonal operator  $T$ , with one column for each candidate and one row for each point in the aggregate.

As an example, consider the 1D-Laplace operator on an eight-node mesh, using standard finite differences, with candidate vectors  $B = [\mathbf{1}, \mathbf{x}]$ . This yields three aggregates,  $\mathcal{A}_i, i = 0, 1, 2$ , as shown in Figure 2a. The aggregation pattern matrix,  $C$ , coupled with the candidate vectors,  $B$ , yield the following

$$(3) \quad C = \begin{bmatrix} * & & & & & & & \\ * & & & & & & & \\ & * & & & & & & \\ & * & & & & & & \\ & * & & & & & & \\ & & * & & & & & \\ & & * & & & & & \\ & & * & & & & & \end{bmatrix}, \quad B = \begin{bmatrix} 1 & 1/9 \\ 1 & 2/9 \\ 1 & 3/9 \\ 1 & 4/9 \\ 1 & 5/9 \\ 1 & 6/9 \\ 1 & 7/9 \\ 1 & 8/9 \end{bmatrix} \rightarrow T = \begin{bmatrix} 1 & -1 & & & & & & \\ 1 & 1 & & & & & & \\ & & 1 & -1 & & & & \\ & & 1 & 0 & & & & \\ & & 1 & 1 & & & & \\ & & & & 1 & -1 & & \\ & & & & 1 & 0 & & \\ & & & & 1 & 1 & & \end{bmatrix} D,$$

where  $D = \text{diag}([2, 2, 3, 2, 3, 2])^{-1/2}$  is a diagonal matrix to normalize each column in the  $l^2$ -norm. Each block of  $T$  in (3) is an orthogonal basis for the restriction of  $B$  to each aggregate.

In the case of a single candidate vector,  $B = [\mathbf{1}]$ ,  $T$  consists of columns 0, 2, and 4 in (3). These columns are plotted in Figure 2b (dashed lines). It is important to note that each column is nonzero only on its respective aggregate. To improve the accuracy of interpolation for algebraically smooth modes — i.e., to make  $\mathcal{R}(P)$  more complementary to relaxation — the columns of the tentative interpolation operator are smoothed — e.g., with weighted Jacobi — to form  $P$  (Figure 2b, solid lines). As the range of  $P$  becomes richer, so does the nonzero footprint in the operator. Indeed, nonzero elements of the middle column of  $P$  in Figure 2b (solid green) now overlap into the neighboring aggregates.

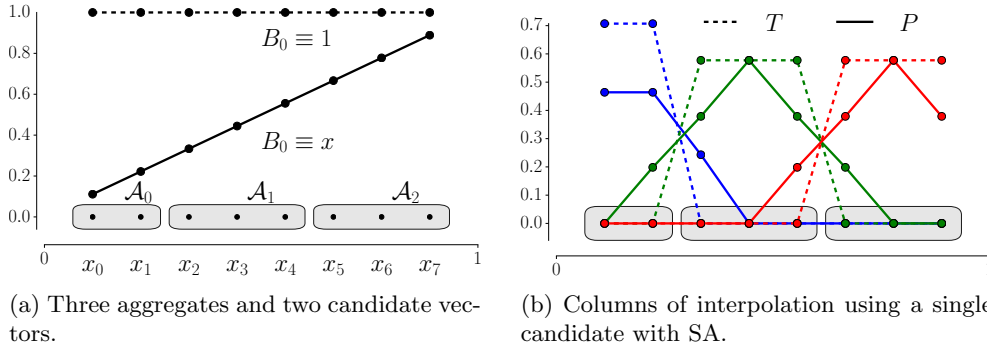


Fig. 2: 1-D Laplace example using SA AMG.

Given a set of candidate vectors  $B$  on the current grid that are exactly represented in the range of  $T$ , coarse-grid candidate vectors are constructed as the pre-image of  $B$  under  $T$ . The motivation is that the pre-image of low energy vectors for  $\mathcal{A}_\ell$  should be low-energy vectors for  $\mathcal{A}_{\ell+1}$ . Let the orthogonal projection onto  $\mathcal{R}(T)$  be given

by  $\pi_T = T(T^T T)^{-1} T^T$ . Since the columns of  $T$  are orthonormal,  $\pi_T = T T^T$ , and requiring  $\pi_T B = B$ , that is  $B \in \mathcal{R}(T)$ , results in a coarse-grid pre-image of  $B$  under  $T$  given by  $B_c = T^T B$ . Since  $B$  is assumed to be a low energy mode, forming  $P$  by applying smoothing iterations to  $T$  to improve multilevel convergence [46] keeps  $B$  close to the range of  $P$ .

A general SA AMG setup algorithm is given in Algorithm 1.

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<b>Algorithm 1:</b> SA_setup()	
<hr/>	
<b>Input:</b>	$A_0$ : fine-grid operator $B_0$ : fine-grid candidate vectors max_size: threshold for max size of coarsest problem
<b>Output:</b>	$A_1, \dots, A_L,$ $P_0, \dots, P_{L-1}$
<hr/>	
1	$\ell = 0$
2	<b>while</b> size( $A_\ell$ ) > max_size
3	$S_\ell = \text{strength}(A_\ell)$ <span style="float: right;">{Strength-of-connection}</span>
4	$\mathcal{A}_\ell = \text{aggregate}(S_\ell)$ <span style="float: right;">{Aggregation}</span>
5	$T_\ell, B_{\ell+1} = \text{inject}(\mathcal{A}_\ell, B_\ell)$ <span style="float: right;">{Form tentative interpolation and coarse candidates}</span>
6	$P_\ell = \text{smooth}(A_\ell, T_\ell)$ <span style="float: right;">{Smooth <math>T_\ell</math>}</span>
7	$A_{\ell+1} = P_\ell^T A_\ell P_\ell$ <span style="float: right;">{Coarse-grid operator}</span>
8	$\ell = \ell + 1$

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**2.2. AMG based on coarse-fine splittings.** In contrast to SA AMG, CF AMG builds a multilevel hierarchy through a CF-splitting. On each level  $\ell = 0, 1, \dots, L$ , the index set of DOFs,  $\Omega^\ell = \{0, \dots, n_\ell\}$ , is split into  $\Omega^\ell = \mathcal{C}_\ell \cup \mathcal{F}_\ell$ , where  $\mathcal{C}_\ell \cap \mathcal{F}_\ell = \emptyset$ . The set  $\mathcal{C}_\ell$  defines the coarse-level DOFs so that  $n_{\ell+1} = |\mathcal{C}_\ell|$ . Similar to SA AMG, a strength measure,  $S$ , is used to determine the splitting so that algebraically smooth error at F-points are well-approximated by evaluating neighboring C-points. Then interpolation,  $P : \mathbb{R}^{n_{\ell+1}} \rightarrow \mathbb{R}^{n_\ell}$ , is formed as

$$(4) \quad P = \left[ \begin{array}{c} W \\ I \end{array} \right] \begin{array}{l} \} \text{F-points} \\ \} \text{C-points} \end{array},$$

where  $W \in \mathbb{R}^{|\mathcal{F}_\ell| \times |\mathcal{C}_\ell|}$  is a sparse matrix, with entries chosen to approximate smooth error at F-points (rows of  $W$ ) as a linear combination of C-points (columns of  $W$ ).

The form of interpolation in (4) highlights two attributes of interpolation in CF AMG that distinguishes it from SA AMG. The first is that the source of complexity and accuracy is clear: the number of nonzeros in  $W$  controls both the accuracy and cost of interpolation. This is explored further in Section 2.4. Second, coarse values are *injected* to the finer grid through the identity in the bottom block. This in turn ensures linear independence of the columns of  $P$ , an important feature not guaranteed in SA AMG.

A general CF AMG setup algorithm is given in Algorithm 2.

**2.3. Benefits and limitations.** In CF AMG, it is assumed that the constant vector is representative of the near null space of the underlying problem, as in the case of a Poisson problem. As a result, convergence can degrade if this is not an accurate



**Algorithm 2:** CF\_setup()

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**Input:**  $A_0$ : fine-grid operator  
 $\text{max\_size}$ : threshold for max size of coarsest problem

**Output:**  $A_1, \dots, A_L,$   
 $P_0, \dots, P_{L-1}$

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1  $\ell = 0$ 
2 while  $\text{size}(A_\ell) > \text{max\_size}$ 
3    $S_\ell = \text{strength}(A_\ell)$                                 {Strength-of-connection}
4    $\mathcal{C}_\ell, \mathcal{F}_\ell = \text{splitting}(S_\ell)$                 {C/F-splitting}
5    $W = \text{weights}(S_\ell, A_\ell, \mathcal{C}_\ell, \mathcal{F}_\ell)$             {Interpolation weights}
6    $P_\ell = \begin{bmatrix} W \\ I \end{bmatrix}$                             {Form interpolation}
7    $A_{\ell+1} = P_\ell^T A_\ell P_\ell$                             {Coarse-grid operator}
8    $\ell = \ell + 1$ 

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assumption. Moreover, the interpolation formulas for CF AMG are static and offer no immediate ways to improve them for more difficult problems.

One benefit of CF AMG is that it provides a structure for controlling sparsity through the determination of weights (see line 5 in Algorithm 2). Each row of  $W$  (see (4)) represents an interpolation formula for a given F-point from surrounding C-points. As a result, this leads to simple filtering strategies in  $P$  [23] that eliminate small entries in each row. In SA AMG, interpolation is based on having algebraically smooth columns of  $P$ . Consequently, existing methods to explicitly filter entries in  $P$  are limited, as removing entries can greatly reduce smoothness [20]. The structure of interpolation operators in CF AMG also allows for theoretical results that are not feasible for arbitrary transfer operators as in SA AMG [25, 49, 38], and which are used in Section 4 to motivate RN AMG.

For problems in which a single global vector adequately represents the algebraically smooth error, SA AMG and CF AMG can each be effective. One benefit of SA AMG is that the method allows for multiple candidate vectors to help define the range of interpolation in order to improve the coarse-grid correction. However, the computational cost of iterating tends to increase substantially when additional candidate vectors are included. Since each candidate vector occupies a column in  $P$  for each aggregate (see (3)), additional candidate vectors quickly increase the number of DOFs and nonzeros in coarse-level operators.

As an example, consider a two-level multigrid method for a 2D-Laplacian discretized with linear, quadrilateral finite elements over a  $50 \times 50$  uniform grid. In this case,  $A_0 \in \mathbb{R}^{2500 \times 2500}$ , with 21904 nonzero elements. Using the symmetric strength matrix — i.e.  $S_{ij} = 1$  if  $A_{ij}/\sqrt{A_{ii}A_{jj}} > 0.25$  [48] — along with standard, greedy aggregation [48] yields 289 aggregates. This results in  $P_0 \in \mathbb{R}^{2500 \times 298}$  in the case of a single candidate vector, and a coarse-grid operator  $A_1 \in \mathbb{R}^{289 \times 289}$  with 2401 nonzeros. Using two candidate vectors,  $P \in \mathbb{R}^{2500 \times 578}$ , and the coarse-grid operator,  $A_1 \in \mathbb{R}^{578 \times 578}$ , has 9604 nonzeros. In this two-grid example, there is approximately a 30% increase in the total number of matrix nonzeros in the hierarchy, which will

correspond to a comparable increase in the cost of each iteration<sup>3</sup>. A key feature of the RN AMG method introduced in Section 3 is that the growth in complexity is mitigated when incorporating multiple candidate vectors in the range of  $P$ .

Additionally, SA AMG provides a simple way to improve interpolation operators through the interpolation smoothing process. While classical SA AMG only uses one weighted-Jacobi iteration to improve  $P$ , multiple iterations as well as smoothing with a *filtered* operator to further improve  $P$  have been used [20, 33]. Nevertheless, because each traditional smoothing iteration expands the sparsity pattern of  $P$ , this process is limited, a problem which is overcome in RN AMG through *a priori* fixed sparsity patterns of transfer operators.

**2.4. Computational cost.** The computational kernel in the multigrid setup and solve phases is a sparse matrix-vector product (SpMV). Thus, a representative measure of the cost of an AMG solver is the number of floating point operations relative to one SpMV with the initial matrix. This measure is referred to as a *work unit* (WU), where one WU is the cost of computing a SpMV on the finest level. Both CF AMG and SA AMG often yield minimal setup costs or setup complexity (SC), but as more features are introduced — e.g., improved SOC methods and energy minimization — the SC may grow. In contrast to the fixed cost of setup, the solve cost depends on the number of iterations or cycles taken, which in turn depends on the stopping residual tolerance. Consequently, the *cycle* complexity (CC), denoted  $\chi_{CC}$ , is defined as the number of WUs required for each multigrid cycle, and is used to measure the solve cost. A similar measure is the *operator* complexity (OC), denoted  $\chi_{OC}$ , which models the cost of a multigrid hierarchy as the ratio of the total number of nonzeros on all levels to the number of nonzeros on the finest-level:

$$(5) \quad \chi_{OC} = \sum_{\ell} \frac{|A_{\ell}|}{|A_0|},$$

where  $|C|$  denotes the number of nonzeros in some sparse matrix  $C$ . Note that this is equivalent to the total cost to perform one SpMV on each level of the hierarchy. Using this, the CC is often considered to scale with OC. For example, in the case of a V(2,2) cycle,  $\chi_{CC} \approx 4\chi_{OC}$ . However, a more detailed model for CC includes the residual computation and coarse-grid correction steps. While it is not typical to account for these parts of the solve phase, they often contribute significantly to the CC, especially for the richer interpolation sparsity patterns examined later. To this end, the CC for a  $V(\nu_{\text{pre}}, \nu_{\text{post}})$ -cycle is defined here as

$$(6) \quad \chi_{CC} = \sum_{\ell} \frac{(\nu_{\text{pre}} + \nu_{\text{post}} + 1)|A_{\ell}| + |P_{\ell}| + |R_{\ell}|}{|A_0|},$$

which reflects pre- and post-relaxation, a residual calculation, and one interpolation and restriction per level (see solve phase discussion in Section 2).

Detailed estimates of the complexity measures are often neglected in numerical results. One contribution of this work is that precise estimates of the SC, OC, and CC are provided for the numerical results presented in Section 5. Coupled with the convergence factor, this information is used to assess the effectiveness of the solver. The SC estimates have been used to expose the expensive parts of the algorithm and motivated the complexity reduction techniques introduced in Section 5.1.

<sup>3</sup>See [30] for an example where a scalar diffusion-like problem requires multiple candidate vectors.



**3. Root-node method.** The general algorithm for constructing a RN AMG hierarchy with  $L + 1$  levels, using energy-minimizing interpolation smoothing, is given in Algorithm 3. The following subsections detail each algorithmic step, comparing and contrasting with SA AMG and CF AMG.

**3.1. Candidate vectors  $B$ .** As with SA AMG, *a priori* knowledge of the algebraically smooth error is assumed as input in the form of a set of candidate vectors  $B$ . These vectors are critical for ensuring accurate interpolation of important algebraically smooth modes. In the case of  $A$  being symmetric, one set of candidate vectors,  $B$ , is sufficient. In the case of non-symmetric problems, a restriction operator  $R$  is formed independently (whereas  $R = P^T$  in the case of symmetry) through the *left* candidate vectors  $\hat{B}$ , which target smooth error in  $A^T$ . Generally if candidate vectors are not known or provided, a constant vector is used, as the constant is geometrically smooth, and a good choice for many problems. Finally, to ensure smooth (including at the boundaries) candidate vectors, a small number of relaxation sweeps are applied to  $A_\ell B_\ell = \mathbf{0}$  (line 13). This improves the algebraic smoothness of  $B_\ell$ , especially near boundaries of a domain. Even for textbook examples such as a Laplacian, the standard candidate  $B_\ell = \mathbf{1}$  can yield a poor approximation to algebraically smooth error near Dirichlet boundaries.

**3.2. Strength matrix  $S$ .** The first level-specific step is the construction of an  $n_\ell \times n_\ell$  SOC matrix,  $S_\ell$  (line 3), which indicates strong connections between DOFs in the problem. This matrix is used for the aggregation of DOFs, and for the construction of a sparsity pattern,  $\mathcal{N}$ , for  $P$ . This work considers the classical strength measure [42], symmetric strength measure [48], and so-called evolution strength measure [39], although other measures have also been proposed [10, 12]. The classical and symmetric strength measures essentially look at the magnitude of an off-diagonal entry when determining if it represents a strong connection. In contrast, the evolution measure computes strength around a DOF  $i$  by locally evolving a unit vector centered at  $i$  with a few sweeps of weighted-Jacobi. This creates a locally smooth vector which is then post-processed to determine which matrix entries in row  $i$  are strong. For instance, for anisotropic diffusion, the directions in which the unit vector diffuses most quickly, are selected as strong connections.

After finishing, each method produces a matrix  $S_\ell$  where individual entries represent the strength-of-connection in the graph of  $A_\ell$ . One modification to  $S_\ell$  that is used here is to normalize each row so that

$$(7) \quad \max_{j \neq i} (S_\ell)_{ij} = (S_\ell)_{ii} = 1.$$

That is, all elements are non-negative, and for each row in  $S_\ell$ , the diagonal and the largest off-diagonal, i.e., the strongest connection, both equal 1. This scaling and the non-binary nature of  $S_\ell$  are important when computing the sparsity pattern in line 7.

For vector-based problems,  $A_\ell$  has a block structure of block-size  $m \times m$  and for common cases such as elasticity, each block corresponds to the DOFs associated with different variables but defined at the same spatial node. It is typical to group each block into a single so-called *supernode* [48], followed by aggregation only at the supernode level. This requires condensing the SOC with an **amalgamate** step in line 5, which reduces  $S_\ell$  to an  $n_\ell/m \times n_\ell/m$  matrix. The amalgamated entry is equal to the maximal entry of its associated  $m \times m$  block in  $S_\ell$ . This allows for aggregation based on supernodes.

**Algorithm 3:** `root_node_setup`


---

**Input:**  $A_0$ : fine-grid operator  
 $B_0$ : fine-grid candidate vectors for  $A$   
 $\hat{B}_0$ : fine-grid candidate vectors for  $A^T$  (if  $A \neq A^T$ )  
 $d$ : interpolation sparsity pattern width  
**vector**: flag indicating vector-based problem  
**max\_size**: threshold for max size of coarsest problem  
**prefilter**: Pre-filtering of interpolation sparsity pattern  
**postfilter**: Post-filtering of interpolation sparsity pattern

**Output:**  $A_1, \dots, A_L$ ,  
 $P_0, \dots, P_{L-1}$ ,  
 $R_0, \dots, R_{L-1}$

```

1  $\ell = 0$ 
2 while  $\text{size}(A_\ell) > \text{max\_size}$ 
3    $S_\ell = \text{strength}(A_\ell)$                                 {Strength-of-connection of matrix}
4   if vector
5      $S_\ell = \text{amalgamate}(S_\ell)$                         {Amalgate from degree-of-freedom to nodal}
6    $C_\ell, \text{roots} = \text{aggregate}(S_\ell)$                   {Construct aggregates and root-nodes}

7    $N_\ell \leftarrow S_\ell^d C_\ell$                             {Form interpolation sparsity pattern}
8   if prefilter
9      $N_\ell = \text{filter}(N_\ell)$                             {Eliminate small entries}
10  if vector
11     $N_\ell, \text{roots} = \text{unamalgamate}(N_\ell, \text{roots})$     {From nodal to degree-of-freedom}
12   $N_\ell = \text{root\_node\_pattern}(N_\ell)$                   {Convert to root-node pattern}

13   $B_\ell = \text{smooth}(A_\ell, B_\ell)$                             {Improve candidates with relaxation}
14   $T_\ell, B_{\ell+1} = \text{inject}(C_\ell, N_\ell, B_\ell, \text{roots})$  {Form tentative interpolation and  $B_{\ell+1}$ }
15   $P_\ell = \text{improve}(A_\ell, T_\ell, B_\ell, B_{\ell+1})$           {Create smooth  $P_\ell$  with  $P_\ell B_{\ell+1} = B_\ell$ }
16  if postfilter
17     $P_\ell = \text{filter}(P_\ell)$                             {Eliminate small entries}
18     $P_\ell = \text{enforce}(P_\ell, B_\ell, B_{\ell+1})$             {Enforce mode constraint with eqn. (12)}
19     $P_\ell = \text{improve}(A_\ell, P_\ell, B_\ell, B_{\ell+1})$         {Re-smooth  $P$  with one iteration}
20  if  $\text{symmetric}(A_\ell)$ 
21     $R_\ell = P_\ell^T$ 
22  else
23     $\hat{B}_\ell = \text{smooth}(A_\ell^T, \hat{B}_\ell)$                     {Improve candidates for the non-symmetric case}
24     $\hat{T}_\ell, \hat{B}_{\ell+1} = \text{inject}(C_\ell, N_\ell, \hat{B}_\ell, \text{roots})$  {Form tentative restriction and  $\hat{B}_{\ell+1}$ }
25     $R_\ell^T = \text{improve}(A_\ell^T, \hat{T}_\ell, \hat{B}_\ell, \hat{B}_{\ell+1})$     {Create smooth  $R_\ell$  with  $R_\ell^T \hat{B}_{\ell+1} = \hat{B}_\ell$ }
26    if postfilter
27       $R_\ell^T = \text{filter}(R_\ell^T)$                         {Eliminate small entries}
28       $R_\ell^T = \text{enforce}(R_\ell^T, \hat{B}_\ell, \hat{B}_{\ell+1})$         {Enforce mode constraint with eqn. (12)}
29       $R_\ell^T = \text{improve}(A_\ell^T, R_\ell^T, \hat{B}_\ell, \hat{B}_{\ell+1})$  {Re-smooth  $R$  with one iteration}
30   $A_{\ell+1} = R_\ell A_\ell P_\ell$                                 {Form coarse-grid}
31   $\ell = \ell + 1$ 

```

---

**3.3. Aggregation.** The next operation is forming an aggregation,  $\mathcal{A}_\ell$ , as in SA AMG (see Algorithm 1), and an associated list of root-nodes (line 6). The classical greedy aggregation algorithm [48] is used, wherein an unaggregated vertex in  $S_\ell$  is selected (as the root-node) and all neighboring vertices with strong edge weights are collected to form an aggregate. An aggregation pattern matrix  $C_\ell$  is then defined as a partition of unity, such that  $C_\ell(i, j) = 1$  if node  $i$  belongs to aggregate  $j$ , and zero otherwise. This pattern will be used in forming the sparsity pattern for  $P_\ell$ . Other aggregation routines such as pairwise aggregation [22, 36] have been considered, but have not demonstrated improvements in RN AMG performance, and the choice of root node is less clear than in a greedy aggregation routine.

Example aggregates for standard isotropic diffusion problems are given in Figure 3. For the systems case,  $m$  DOFs at each supernode implies that each root-node also contains  $m$  DOFs.

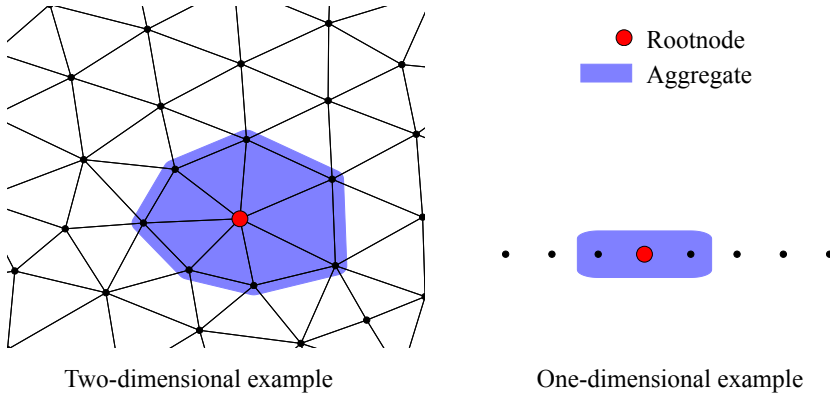


Fig. 3: Sample aggregates on the finest level for standard isotropic diffusion in one and two dimensions.

**3.4. Arbitrary sparsity pattern  $\mathcal{N}$ .** Aggregation gives an interpolation structure, with each root-node corresponding to one block column of  $T$  and  $P$  as in (3), but not a sparsity pattern for  $P$ . Thus, in Lines 7–12, the sparsity pattern  $\mathcal{N}_\ell$  for interpolation is built, where  $(i, j) \notin \mathcal{N}_\ell \Leftrightarrow N_\ell(i, j) = 0$ . Nonzero elements are based on growing the aggregation pattern matrix,  $C_\ell$ , based on the strength matrix,  $S_\ell$ , through multiplication  $N_\ell = S_\ell^d C_\ell$ , for some *degree* or *distance*  $d$ . Using  $d$  applications of  $S_\ell$  extends the interpolation stencil for a given F-point to a distance of  $d$  in the strength matrix. Large values of  $d$  allow the sparsity pattern to grow in the direction of strong connections, allowing for long-distance interpolation. This approach differs from [41, 45] in that a normalized SOC matrix (7) is used with  $C_\ell$ , and as a result,  $N_\ell$  can be *filtered* (line 9) by examining the magnitude of the entries: larger entries indicate a stronger path from the root-node  $i$ . As a result, the normalization of  $S_\ell$  imbues a relative size across columns so that the product with the binary aggregation matrix  $C_\ell$  yields individual entries related to strength-of-connection.

In the case of vector problems, the **unamalgamate** step in line 11 reverses amalgamation and converts the list of root-nodes into a list of DOFs. If the amalgamated list of roots is of length  $n_{\ell+1}/m$ , then the unamalgamated list is of length  $n_\ell$ . Similarly, the unamalgamated entry  $N_\ell(i, j)$  is equal to the amalgamated entry  $N_\ell(\lfloor i/m \rfloor, \lfloor j/m \rfloor)$ .

The last step in computing the sparsity is `root_node_pattern` in line 12. This function enforces the root-node pattern of interpolation in equation (4) by traversing  $N_\ell$  to change each root-node row to be the corresponding row of the identity — i.e., the restriction to each root-node for the coarse-grid does not involve other root-nodes, and each root-node is interpolated by value back to the fine grid. Other than this root-node requirement, RN AMG allows for arbitrary sparsity patterns and enables selective control of the number of interpolation points. This flexibility is used in the next section, where the sparsity pattern is filtered (dropping entries) in a way that targets only *strong* long-distance connections in  $P_\ell$ .

It should be noted that the sparsity pattern for  $R^T$  is built in the same manner as  $P$ : by expanding a tentative operator with a SOC matrix based on  $A$ . Another option is to construct the sparsity pattern for  $R$  based on a SOC from  $A^T$ ; however, numerical results consistently indicate a degradation in convergence factors when doing so. Choosing the *optimal* sparsity pattern for both  $P$  and for  $R$  remains an important and open research question.

**3.5. Filtering sparsity pattern.** A filtering step, which removes nonzeros in  $N_\ell$ , is used after the construction of the sparsity pattern for  $P$ . A large degree  $d$  (see line 7) is often needed [45, 30] to construct effective interpolation operators; filtering can limit the additional cost due to the growth in the sparsity pattern. In particular, filtering allows for long-distance interpolation in the direction of strong connections, while limiting complexity.

Pre-filtering is used to filter the sparsity pattern matrix  $N_\ell$  before energy-minimization. Here, entries are eliminated in  $N_\ell$  prior to constructing  $P_\ell$  based on the size of entries in  $N_\ell = S_\ell^d C_\ell$ , which indicate the strength-of-connection between two DOFs. Because pre-filtering is only based on SOC and not the fully formed interpolation operator, it is possible that influential entries are inadvertently removed, thus degrading convergence. However, in practice trimming the sparsity pattern of  $P_\ell$  before initiating the construction significantly lowers the SC in many cases, with minimal impact on AMG convergence.

Given an initial sparsity pattern, entries are filtered as in [23], by either retaining the  $k$  largest values in a row or by applying a drop tolerance  $\theta$ . Algorithm 4 describes this process in detail, where  $\max(G, i, k)$  is the  $k$ th largest off-diagonal entry in row  $i$ . The idea of pre-filtering has shown to be effective for model problems using a polynomial approximation to  $A_{ff}^{-1}$  in [14]. In contrast, the pre-filtering used here is less expensive and relies on values already computed by the root-node algorithm.

---

**Algorithm 4: filter( $G$ )**


---

**Input:**  $G$ : matrix to be filtered  
 $\theta$ : filtering drop-tolerance  
 $k$ : filtering threshold

**Output:**  $G$

```

1 if  $k$ 
2   for  $|G(i, j)| < \max(G, i, k)$  do
3      $G(i, j) \leftarrow 0$ 
4 if  $\theta$ 
5   for  $|G(i, j)| < \theta \max(G, i, 1)$  do
6      $G(i, j) \leftarrow 0$ 

```

---

**3.6. Interpolation construction.** A *tentative* interpolation operator,  $T$ , is constructed from the sparsity pattern. The full interpolation is then formed based on a constrained energy-minimization with the following principles:

- i.  $T$  and  $P$  satisfy interpolation constraints of provided algebraically smooth candidates. That is,

$$(8) \quad TB_c = B \quad \text{and} \quad PB_c = B,$$

for candidates  $B$ , and coarse level candidates  $B_c$ .

- ii. The **improve** procedure reduces the *energy* of each  $j$ th column of  $P$ :

$$(9) \quad \|P_{(j)}\| \leq \|T_{(j)}\|,$$

for some  $A$ -induced norm  $\|\cdot\|$ . For instance, interpolation smoothing in SA AMG is one example.

- iii. Given sparsity pattern  $\mathcal{N}$ ,

$$(10) \quad T_{ij} = 0 \quad \text{if } (i, j) \notin \mathcal{N} \quad \text{and} \quad P_{ij} = 0 \quad \text{if } (i, j) \notin \mathcal{N}.$$

The energy-minimization approach [41] is used here, which satisfies these principles. Energy-minimization is an iterative smoothing process that improves  $P$ , through several passes. As a result, growth in the sparsity pattern of  $P$  necessitates a constraint on the sparsity pattern constraint,  $\mathcal{N}$ . However, in enforcing  $\mathcal{N}$  by dropping entries in  $P$ , the constraints are no longer satisfied. In response, the constraints are enforced as an additional step.

Root-node AMG proceeds by taking the aggregation and list of root-nodes to construct the coarse-grid candidates by injection,  $B_{\ell+1}(i, j) = B_\ell(k, j)$ , where  $k$  is the  $i$ th root-node. If  $m$  is the block size of the original matrix ( $m = 1$  for a scalar problem), then *only* the first  $m$  candidates are injected over each aggregate to form an initial  $T_\ell$  (line 14). As a result, each root-node represents  $m$  DOFs on the coarse grid. An additional step is performed on  $T_\ell$ , normalizing each column so that the coarse-grid variables inject to the fine-grid root-nodes. This process yields the following form

$$(11) \quad T_\ell = \left[ \begin{array}{c} W_\ell \\ I \end{array} \right] \begin{array}{l} \} \text{Non Root-nodes} \\ \} \text{Root-nodes} \end{array}.$$

For  $m = 1$ ,  $T_\ell$  has non-overlapping columns; for  $m > 1$ ,  $W_\ell$  is block diagonal, as  $T$  in (3). With the identity over C-points in  $T_\ell$ , RN AMG resembles that of CF AMG (cf. (4)).

If there are more than  $m$  candidates, the remaining candidates are projected into  $\text{ran}(T_\ell)$  in the Euclidean inner-product. In RN AMG, it is assumed that the sparsity pattern has sufficient DOFs that this is an underdetermined problem. This results in the minimal norm update to each row of  $T_\ell$  such that  $T_\ell B_{\ell+1} = B_\ell$  and  $T_\ell$  obeys the sparsity pattern  $\mathcal{N}_\ell$ . More specifically, an update  $u$  to only the allowed nonzero portion of row  $i$  of  $T_\ell$  (called  $t$ ) is computed by solving

$$(12) \quad (t + u)B_{\ell+1} = B_\ell \quad \Leftrightarrow \quad B_{\ell+1}^T u^T = B_\ell^T - B_{\ell+1}^T t^T,$$

for  $u$  using least-squares. Note that only  $W_\ell$  is modified in (12), that is, injection over C-points is maintained and  $W_\ell$  is expanded to interpolate candidate vectors.

Next, interpolation  $P_\ell$  is formed using **improve** in line 15. Energy-minimization forms  $P_\ell$  as a succession of energy-minimization updates to  $T_\ell$ . Each update  $U$  is

computed to reduce the energy of each column via a Krylov process and to also satisfy  $UB_{\ell+1} = 0$ . As a result, (i)  $(T_\ell + U)B_{\ell+1} = T_\ell B_{\ell+1}$  (constraints are satisfied exactly), and (ii)  $\|T_\ell + U\| \leq \|T - \ell\|$  (energy is reduced). The projection operator enforcing the constraints is analogous to (12) (see [41]).

**3.7. Filtering interpolation.** After  $P_\ell$  is formed, a post-filtering process (similar to pre-filtering in Section 3.5) is applied in line 17 to reduce complexity. Post-filtering removes elements directly from  $P_\ell$  after smoothing, but this leads to a  $P_\ell$  that violates the mode interpolation constraints (8). Thus, the function `enforce` is used to re-apply these constraints via (12). Finally, an additional iteration of `improve` is used to account for large increases in energy caused by removing entries. One advantage of post-filtering is that element removal is based on the actual smoothed interpolation stencil entries, and is thus less likely to inadvertently degrade convergence compared with pre-filtering — i.e. by removing important entries from  $P$ . Post-filtering generally results in a lower complexity in the Galerkin coarse-grid operator (line 30) and all subsequent coarser grid operations. A similar filtering approach is also effective for classical AMG methods [23]. However, post-filtering does not reduce the OC and SC as effectively as pre-filtering, in particular because energy-minimization (one of the dominant costs in SC) is applied to a larger sparsity pattern, which is then trimmed afterwards.

**3.8. Coarse-grid construction.** The final step in Algorithm 3 is the construction of the coarse-grid operator through a (Petrov) Galerkin triple-matrix product  $A_{\ell+1} = R_\ell A_\ell P_\ell$ . For symmetric  $A_\ell$ , restriction is the transpose of interpolation. For non-symmetric  $A_\ell$ , the interpolation construction process is duplicated for  $R_\ell^T$  using  $A_\ell^T$  (the  $\hat{\cdot}$  notation is used to denote the quantities used to compute  $R_\ell$ ) [18].

**3.9. Discussion: Compare and contrast RN AMG with SA AMG.** Similar to SA AMG, RN AMG facilitates the use of multiple, arbitrary candidate vectors, but handles the complexity challenges by not adding columns to  $P$  for each additional candidate. The dimensions of  $T$  and  $P$  are fixed, with the number of columns in  $T$  and  $P$  being equal to the number of aggregates. The candidates  $B$  are projected exactly into  $\text{ran}(P)$  and  $\text{ran}(T)$ , assuming a sparsity pattern with enough entries. With SA AMG, for each candidate vector added to  $B$ , a new column is added to  $T$  and  $P$  for every aggregate (see (3)). This difference allows RN AMG to exhibit significantly lower complexity than SA AMG in some instances.

Root-node AMG also uses interpolation smoothing, like SA AMG, so that  $P$  is iteratively improved. However, standard interpolation smoothing does *not* satisfy the exact candidate interpolation constraints in general, although it does attempt to target the same three principles outlined in Section 3.6 (including accurate but not exact candidate vector interpolation). A consequence of satisfying the constraints is that the candidate vectors are exactly represented in the range of interpolation. To measure this, consider the error in interpolating  $B$  found through an orthogonal projection of  $B$  into the range of interpolation:

$$(13) \quad e_B = (I - PP^\dagger)B,$$

where  $P^\dagger$  is the pseudo-inverse of  $P$ . The point-wise values of the error  $e_B$  are shown in Figure 4, where RN AMG achieves much lower error, directly satisfying the constraints (globally).

Additionally, standard interpolation smoothing does not allow for multiple smoothing passes (and hence longer-distance interpolation) without suffering from complex-



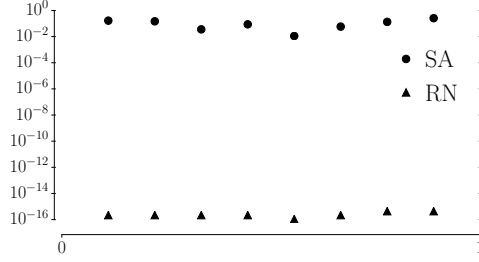


Fig. 4: Error  $e_B$  in satisfying the constraints  $B \equiv \mathbf{1} \in \text{ran}(P)$  for a 1D Poisson problem with eight DOFs using central finite differences.

ity issues and fill-in in  $P$ . The arbitrary sparsity pattern allows RN AMG to have longer-distance interpolation and multiple smoothing iterations, while also effectively managing complexity.

An SA AMG-type method supporting vector problems can be derived from Algorithm 3 by removing the root-node specific lines and by using the `inject` and `smooth` functions from Algorithm 1 to form  $T_\ell$  and  $P_\ell$ . This form of SA AMG also supports non-symmetric problems [43].

### 3.10. Discussion: Compare and contrast RN AMG with CF AMG.

Compared with CF AMG, RN AMG adopts the structure of  $P$  from (4), which aids the use of pre- and post-filtering since the columns of interpolation are normalized (with the identity). (In Section 4, additional theoretical motivation for using this form is investigated.) To understand this, consider how the CF AMG structure (11) scales interpolation around each root-node for a simple 1D Laplace example on eight nodes with finite differencing and  $B = \mathbf{1}$ . In this case, three aggregates are formed as in Figure 2a. Each column of  $T$  is shown in Figure 5a, where a relative balance in weighting across aggregates is observed, in contrast to SA AMG, which is shown in Figure 2b. This balance in weighting is evident again in the final  $P$  in Figure 5b, again because the identity form is preserved according to equation (11).

This scaling aids the filtering strategy, which eliminates relatively small interpolation weights in the rows of  $P$ , because it provides the rationale for comparing the magnitude of entries across columns. In other words, RN AMG stipulates that a large interpolation weight is one, representing injection from the coarse to fine grid, visible as the peaks in Figure 5b.

In contrast to CF AMG, there are the many similarities that RN AMG shares with SA AMG, with the ability to iteratively improve  $P$  and guarantee accurate interpolation of the user-defined candidate vectors  $B$  being key differences. Regarding similarities to more recent adaptive CF-style AMG methods, bootstrap AMG (BAMG) [8] also fits multiple candidate vectors into the range of interpolation, but does so by generating a set of many candidates and over-determining each row of  $P$ . In this respect, RN AMG is closer to SA AMG, as each row of  $P$  is under-determined and only fits a small number of candidate vectors into the range of  $P$ , using interpolation smoothing to fully determine each interpolation entry.

In contrast to both CF AMG and SA AMG, RN AMG allows for automatically expanded sparsity patterns  $\mathcal{N}_\ell$ , which can be filtered, to facilitate long-distance interpolation. This is critical for robustness and performance, as shown in the examples in Section 5, and is a unique feature of RN AMG.

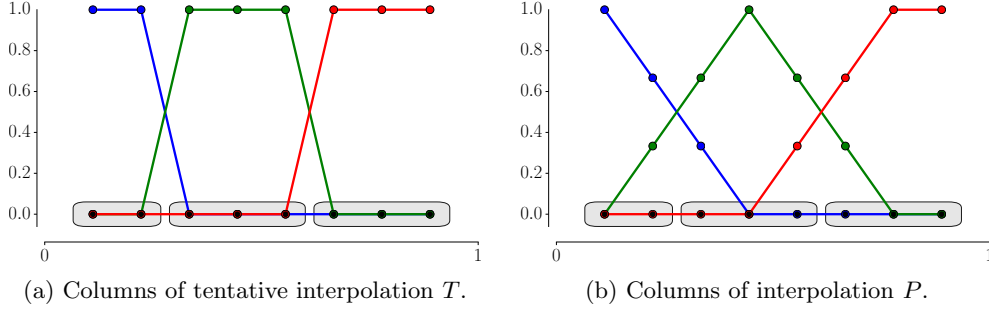


Fig. 5: Interpolation using a single candidate  $B = \mathbf{1}$  in RN AMG.

**4. Theoretical motivation for root-node.** In this section, theoretical motivation for RN AMG is introduced. Initially, it is assumed that  $A$  is SPD in order to connect with classical AMG theory. Based on relations established in the symmetric setting, some results are extended to non-symmetric systems in Section 4.2.

**4.1. The symmetric case.** Let  $A \in \mathbb{R}^{n \times n}$  be SPD,  $\|\cdot\|$  and  $\|\cdot\|_A$  represent the  $l^2$ - and  $A$ -norms, respectively, and  $P : \mathbb{R}^{n_c} \rightarrow \mathbb{R}^n$  be an interpolation operator defining a coarse space of size  $n_c$ . The error propagation operator for a two-grid method is given by

$$(14) \quad E_{TG} = I - P(P^T A P)^{-1} P^T A,$$

and a multilevel version,  $E_{MG}$  is similarly defined (see [32]). Here, bounds on the  $A$ -norm of  $E_{TG}$  and  $E_{MG}$  are constructed. The *weak approximation property* (WAP) gives necessary and sufficient conditions for two-grid convergence as follows: there exists  $K \in \mathbb{R}$  such that for any vector  $\mathbf{u} \in \mathbb{R}^n$ ,

$$(15) \quad \min_{\mathbf{w}_c \in \mathbb{R}^{n_c}} \|\mathbf{u} - P\mathbf{w}_c\|^2 \leq \frac{K}{\|A\|} \|\mathbf{u}\|_A^2,$$

wherein  $\|E_{TG}\|_A = 1 - \frac{1}{K_{TG}}$  and  $1 \leq K_{TG} \leq K$  [49]. For simplicity, (15) is based on Richardson relaxation. A WAP with respect to a general relaxation scheme along with a tight bound on  $K_{TG}$  can be found in [25, 49]. The *strong approximation property* (SAP) establishes multilevel convergence with a stronger condition: there exists  $K \in \mathbb{R}$  such that for any vector  $\mathbf{u} \in \mathbb{R}^n$ ,

$$(16) \quad \min_{\mathbf{w}_c \in \mathbb{R}^{n_c}} \|\mathbf{u} - P\mathbf{w}_c\|_A^2 \leq \frac{K}{\|A\|} \|A\mathbf{u}\|^2.$$

If (16) holds on each level of the hierarchy, then  $\|E_{MG}\|_A = 1 - \frac{1}{K_{MG}}$  and  $1 \leq K_{MG} \leq 1 + K \frac{\|M\|}{\|A\|}$ , where  $M$  is the chosen relaxation scheme of the form  $\mathbf{x}_{k+1} = \mathbf{x}_k + M^{-1} \mathbf{r}_k$ , for residual  $\mathbf{r}$  [49].

Since  $A$  is assumed to be SPD, its eigenvectors form an  $l^2$ - and  $A$ -orthonormal basis for the space  $\mathbb{R}^n$ . Thus, if the WAP and SAP hold for all eigenvectors, they hold for all vectors, and it follows that the WAP requires eigenvectors be interpolated with accuracy on the order of the corresponding eigenvalue, and the SAP requires interpolation accuracy on the order of the eigenvalue squared. This leads to an equivalence of satisfying the WAP based on  $A^2$  and the SAP for  $A$  as follows:

LEMMA 1 (Lemma 5.20 [49]). *Let  $A \in \mathbb{R}^{n \times n}$  be SPD and  $P \in \mathbb{R}^{n \times n_c}$ . Then*

$$(17) \quad \min_{\mathbf{w}_c \in \mathbb{R}^{n_c}} \|\mathbf{u} - P\mathbf{w}_c\|^2 \leq \frac{K^2}{\|A^2\|} \|\mathbf{u}\|_{A^2}^2 \quad \text{for all } \mathbf{u},$$

*if and only if*

$$(18) \quad \min_{w_c} \|\mathbf{u} - P\mathbf{w}_c\|_A^2 \leq \frac{K}{\|A\|} \|A\mathbf{u}\|^2 \quad \text{for all } \mathbf{u}.$$

The accuracy demands of the WAP and SAP with respect to eigenvalues indicates that the range of  $P$  should contain eigenvectors of  $A$  associated with small eigenvalues (or so-called *algebraically smooth* modes). In building AMG hierarchies, this generally takes one of two forms, (i) ensuring that known low-energy modes are exactly represented in the range of  $P$ , and (ii) minimizing columns of  $P$  in the  $A$ -norm so that the range of  $P$  corresponds to algebraically smooth vectors. Recall that RN AMG combines each of these approaches in a constrained energy-minimization [14, 41, 45, 40].

Given a CF-splitting of the current grid, consider a matrix ordering of the form  $A = \begin{bmatrix} A_{ff} & A_{fc} \\ A_{cf} & A_{cc} \end{bmatrix}$ , where  $A_{ff}$  corresponds to F-point-to-F-point connections,  $A_{fc}$  to F-point-to-C-point connections, and so on. In a CF AMG context, interpolation is then assumed to have the form  $P = \begin{pmatrix} W \\ I \end{pmatrix}$ , where the lower identity block interpolates and restricts C-points by injection and  $W$  interpolates and restricts F-points based on linear combinations. The minimizing coarse-grid vector,  $\mathbf{w}_c$ , in the WAP and the SAP is given by  $l^2$ -orthogonal and  $A$ -orthogonal projections of the vector  $\mathbf{u}$  onto the range of  $P$ , respectively. In practice, such projections are generally too expensive to form explicitly; thus, computable measures are also of interest. One option consistent with CF AMG is to let  $\mathbf{w}_c = \mathbf{u}_c$ , that is define  $\mathbf{w}_c$  as the restriction of  $\mathbf{u}$  to C-points. This provides a bound on the WAP, as  $\min_{\mathbf{w}_c \in \mathbb{R}^{n_c}} \|\mathbf{u} - P\mathbf{w}_c\|^2 \leq \|\mathbf{u} - P\mathbf{u}_c\|^2$  for all  $\mathbf{u}$ , and thus

$$(19) \quad \mu(P) := \max_{\mathbf{u} \neq 0} \frac{\|\mathbf{u} - P\mathbf{u}_c\|^2}{\|\mathbf{u}\|_A^2} \geq K_{TG}.$$

Assuming  $P = \begin{pmatrix} W \\ I \end{pmatrix}$ , the optimal interpolation operator under  $\mu(P)$  is given by

$$(20) \quad \begin{aligned} P_{\text{ideal}} &= \operatorname{argmin}_P \max_{\mathbf{u}} \frac{\|\mathbf{u} - P\mathbf{u}_c\|^2}{\|\mathbf{u}\|_A^2} \\ &= \begin{bmatrix} -A_{ff}^{-1}A_{fc} \\ I \end{bmatrix}, \end{aligned}$$

where  $P_{\text{ideal}}$  is referred to as “ideal interpolation.”

In addition to  $P_{\text{ideal}}$  being optimal with respect to the measure  $\mu(P)$  (and thus satisfying the WAP for some  $K \geq K_{TG}$ ), if  $A_{ff}$  is well-conditioned, then  $\|A_{ff}^{-1}\|$  is bounded by a small constant, in which case  $P_{\text{ideal}}$  also satisfies the SAP:

$$\min_{\mathbf{w}_c \in \mathcal{V}_c} \|\mathbf{u} - P_{\text{ideal}}\mathbf{w}_c\|_A^2 \leq \|\mathbf{u} - P_{\text{ideal}}\mathbf{u}_c\|_A^2 \leq \|A_{ff}^{-1}\| \|A\mathbf{u}\|^2 \leq \frac{K}{\|A\|} \|A\mathbf{u}\|^2,$$

for some  $K$ . While  $P_{\text{ideal}}$  indicates an effective interpolation scheme,  $A_{ff}^{-1}A_{fc}$  is often a dense matrix and difficult to compute. However, if  $A_{ff}$  is well-conditioned, its entries decay exponentially fast away from the diagonal [14], suggesting that a sparse approximation can be formed. In addition to directly satisfying constraint vectors, the energy-minimization piece of root-node interpolation constructs a sparse approximation to  $P_{\text{ideal}}$ , with the goal of retaining the convergence properties of ideal interpolation, while limiting coarse-grid complexity. In general, aggregation-based AMG is motivated through energy-minimization principles over the columns of the interpolation operator [47, 46, 35, 19, 21, 29]. The root-node approach, however, is supported by a more complete theoretical motivation, as well as practical benefits including the use of aggregation or CF-splittings for coarsening, and better conditioning of the coarse-grid operator in the non-symmetric setting (Section 4.2).

Observe that  $AP_{\text{ideal}} = \begin{pmatrix} 0 \\ S \end{pmatrix}$ , where  $S = A_{cc} - A_{cf}A_{ff}^{-1}A_{fc}$  is the Schur complement of  $A$ . Given that  $AP_{\text{ideal}} = 0$  over F-points, this motivates minimizing columns of  $P = \begin{pmatrix} W \\ I \end{pmatrix}$  in the  $A$ -norm to approximate the action of  $P_{\text{ideal}}$ . The identity block over C-points along with any constraints enforced ensure that columns of  $W$  are nonzero (the solution to minimizing a general  $P$  in the  $A$ -norm without constraints is  $P = \mathbf{0}$ ). Coupled with a predetermined sparsity pattern and constraints  $PB_c = B$ , where  $B$  is a set of column-wise constraint vectors to be in the range of  $P$  and  $B_c$   $B$  restricted to C-points, energy minimization in the  $A$ -norm is exactly the conjugate gradient variant of energy minimization proposed in [41]. Lemma 2 shows the relationship between  $P_{\text{ideal}}$  and energy minimization. That is, the CG variant of energy minimization used to form  $P$  over a given sparsity pattern is equivalent to minimizing the difference between columns of  $P$  and  $P_{\text{ideal}}$  in the  $A$ -norm, over a given sparsity pattern.

LEMMA 2. Let  $A \in \mathbb{R}^{n \times n}$  be SPD,  $P_{\text{ideal}} \in \mathbb{R}^{n \times n_c}$  be given by (20), and  $\mathbf{e}_\ell$  the  $\ell$ th canonical basis vector, where  $\mathbf{p}_\ell = P\mathbf{e}_\ell$  is the  $\ell$ th column of  $P$ . Denote by  $\mathcal{N}^F$  a sparsity pattern for any matrix  $W \in \mathbb{R}^{n_f \times n_c}$  where  $W_{ij} \neq 0$  if  $(i, j) \in \mathcal{N}^F$ . Then define the set  $\mathcal{P}_\ell$  as the  $\ell$ th column of any matrix with the structure of  $P_{\text{ideal}}$ , restricted to sparsity pattern  $\mathcal{N}^F$ , that is

$$(21) \quad \mathcal{P}_\ell = \left\{ P\mathbf{e}_\ell : P = \begin{bmatrix} W \\ I \end{bmatrix}, \text{ where } W \in \mathbb{R}^{n_f \times n_c} \text{ and } W_{ij} = 0 \text{ if } (i, j) \notin \mathcal{N}_{ij}^F \right\}.$$

Then for  $l = 1, \dots, n_c$ ,

$$(22) \quad \operatorname{argmin}_{\mathbf{p}_\ell \in \mathcal{P}_\ell} \|\mathbf{p}_\ell\|_A = \operatorname{argmin}_{\mathbf{p}_\ell \in \mathcal{P}_\ell} \|\mathbf{p}_\ell - P_{\text{ideal}}\mathbf{e}_\ell\|_A.$$

Equivalently, minimizing the columns of  $P$  in the  $A$ -norm is equivalent to minimizing the difference between columns of  $P$  and  $P_{\text{ideal}}$  in the  $A$ -norm.

*Proof.* Equivalence is established by demonstrating identical weak forms for the two minimization problems in (22). Consider  $\mathbf{p}_\ell \in \mathcal{P}_\ell$  and define  $N_\ell$  to be the diagonal matrix that enforces sparsity pattern  $\mathcal{N}^F$  on the  $\ell$ th column of  $W$ ,  $\mathbf{w}_\ell$ . That is,  $\mathbf{w}_\ell = W\mathbf{e}_\ell = N_\ell W\mathbf{e}_\ell$ , where the  $k$ th entry of  $\mathbf{w}_\ell$  equals zero if  $(k, \ell) \notin \mathcal{N}^F$ .

1. Consider minimizing the  $\ell$ th column of  $P$ , given by  $\mathbf{p}_\ell = \begin{bmatrix} \mathbf{w}_\ell \\ \mathbf{e}_\ell \end{bmatrix}$ , in the  $A$ -norm.

To this end, define the functional  $G(\mathbf{w}_\ell) = \langle A\mathbf{p}_\ell, \mathbf{p}_\ell \rangle = \left\langle A \begin{bmatrix} \mathbf{w}_\ell \\ \mathbf{e}_\ell \end{bmatrix}, \begin{bmatrix} \mathbf{w}_\ell \\ \mathbf{e}_\ell \end{bmatrix} \right\rangle$ ,

with first variation

$$G'(\mathbf{w}_\ell; \mathbf{v}) = 2 \left\langle N_\ell A_{ff} N_\ell \mathbf{w}_\ell - N_\ell A_{fc} \mathbf{e}_\ell, \mathbf{v} \right\rangle$$

for  $\mathbf{v} = N_\ell \mathbf{v}$ . The weak form for of minimizing  $G$  is then given by

$$N_\ell A_{ff} N_\ell \mathbf{w}_\ell = N_\ell A_{fc} \mathbf{e}_\ell.$$

2. Consider minimizing the difference between the  $l$ th column of  $P$  and  $P_{\text{ideal}}$  in the  $A$ -norm. That is, define the functional

$$\begin{aligned} H(\mathbf{w}_\ell) &= \left\langle A(P - P_{\text{ideal}}) \mathbf{e}_\ell, (P - P_{\text{ideal}}) \mathbf{e}_\ell \right\rangle \\ &= \left\langle A_{ff}(N_\ell \mathbf{w}_\ell - A_{ff}^{-1} A_{fc} \mathbf{e}_\ell), N_\ell \mathbf{w}_\ell - A_{ff}^{-1} A_{fc} \mathbf{e}_\ell \right\rangle. \end{aligned}$$

Taking the first variation yields

$$H'(\mathbf{w}_\ell; \mathbf{v}) = \left\langle N_\ell A_{ff} N_\ell \mathbf{w}_\ell - N_\ell A_{fc} \mathbf{e}_\ell, \mathbf{v} \right\rangle,$$

with  $\mathbf{v} = N_\ell \mathbf{v}$ , resulting in the weak form

$$N_\ell A_{ff} N_\ell \mathbf{w}_\ell = N_\ell A_{fc} \mathbf{e}_\ell. \quad \square$$

*Remark 3.* A similar result to Lemma 2 can be found in [14] in the Frobenius norm.

Although  $P_{\text{ideal}}$  in (20) is motivated through (and optimal in the sense of (19)) the WAP (15), and with an appropriate CF-splitting satisfies the SAP (16),  $P_{\text{ideal}}$  is not optimal in any sense with respect to the SAP. However, a similar derivation leads to an equivalent “ideal interpolation” operator with respect to the SAP, as introduced in Lemma 4.

LEMMA 4. Let  $A \in \mathbb{R}^{n \times n}$  be SPD and  $P \in \mathbb{R}^{n \times n_c}$  take the form  $P = \begin{pmatrix} W \\ I \end{pmatrix}$ , that is  $C$ -points are interpolated by injection. Then, consider the SAP under the assumption that the pre-image of any vector  $\mathbf{u}$  under  $P$  is given by  $\mathbf{u}_c$ ,

$$\hat{\mu}(P) := \max_{\mathbf{u} \neq 0} \frac{\|\mathbf{u} - P \mathbf{u}_c\|_A^2}{\|A \mathbf{u}\|^2}.$$

Then for any smoothing scheme,  $M$ ,  $K_{MG} \leq 1 + \hat{\mu}(P) \frac{\|M\|}{\|A\|}$ , and the optimal  $P$  with respect to minimizing  $\hat{\mu}$  is given by

$$\operatorname{argmin}_P \hat{\mu}(P) = \left[ \begin{array}{c} (A_{ff}^2 + A_{fc} A_{cf})^{-1} (A_{ff} A_{fc} + A_{fc} A_{cc}) \\ I \end{array} \right],$$

which is exactly ideal interpolation (20) for  $A^2$ .

*Proof.* First note from the SAP,

$$\begin{aligned} K_{MG} &\leq 1 + \left( \max_{\mathbf{u} \neq 0} \min_{\mathbf{w}_c} \frac{\|\mathbf{u} - P \mathbf{w}_c\|_A^2}{\|A \mathbf{u}\|^2} \right) \frac{\|M\|}{\|A\|} \\ &\leq 1 + \left( \max_{\mathbf{u} \neq 0} \frac{\|\mathbf{u} - P \mathbf{u}_c\|_A^2}{\|A \mathbf{u}\|^2} \right) \frac{\|M\|}{\|A\|}. \end{aligned}$$

Based on the proof of Theorem 3.1 and Corollary 3.2 in [25], it follows that

$$\operatorname{argmin}_P \max_{\mathbf{u} \neq 0} \frac{\|\mathbf{u} - P\mathbf{u}_c\|_A^2}{\|A\mathbf{u}\|^2} = \operatorname{argmin}_P \max_{\mathbf{u} \neq 0} \frac{\|\mathbf{u} - P\mathbf{u}_c\|^2}{\|A\mathbf{u}\|^2} = \operatorname{argmin}_P \max_{\mathbf{u} \neq 0} \frac{\|\mathbf{u} - P\mathbf{u}_c\|^2}{\|\mathbf{u}\|_{A^2}^2}.$$

The final equation is  $\mu(P)$  (19) as applied to  $A^2$ , and thus the minimum is attained by ideal interpolation (20) as applied to  $A^2$ .  $\square$

*Remark 5.* An equivalent result holds for  $A^*A$  and  $\sqrt{A^*A}$  (as opposed to  $A^2$  and  $A$ ), which is used in the following section on non-symmetric operators.

**4.2. The non-symmetric case.** The difficulties of convergence theory for non-symmetric  $A$  lie in the fact that  $A$  no longer defines a norm, raising the question of an appropriate norm to measure convergence. The spectral radius bounds asymptotic convergence factors [37, 50]; however, consider the following error propagation matrix as an example:

$$E = \begin{pmatrix} \epsilon & k \\ 0 & \epsilon \end{pmatrix},$$

for  $\epsilon < 1$ . Irrespective of  $k$ ,  $\rho(E) = \epsilon$ , while if  $k = 0$ , then the convergence factor in any reasonable norm should be  $\epsilon$ . But, if  $k = 10^8$ , let  $\mathbf{e}_0 = [0, 1]^T$  and the convergence factor in the  $l^2$ -norm after one iteration is

$$\rho_1 = \frac{\|\mathbf{e}_1\|}{\|\mathbf{e}_0\|} = \|M\mathbf{e}_0\| = \sqrt{1000^2 + \epsilon^2} \approx 1000.$$

This demonstrates the downside of considering convergence for non-symmetric AMG methods based on spectral radius, namely that iterations may diverge, and the spectral radius bound on the convergence factor is not necessarily achieved in practice. In the case of SPD  $A$ , the two-grid AMG error propagation operator,  $E_{TG}$ , is  $A$ -symmetric and  $\|E_{TG}\|_A = \rho(E_{TG})$ . In [18], energy-norm convergence is extended to non-symmetric problems through  $\sqrt{A^*A}$ - and  $\sqrt{AA^*}$ -norms (using the square root to maintain the order of the problem). Here, for non-singular  $A$ , energy minimization and RN AMG are related to two-grid theory in the  $\sqrt{A^*A}$ -norm.

Energy-minimization techniques for non-symmetric problems have also been proposed in the  $A^*A$ -norm for  $P$  and  $AA^*$ -norm for  $R$ , as opposed to the  $A$ -norm as referenced in Lemma 2 (see for example, fGMRES [41, (2.7)] and CGNR [41, (2.34)]), and are the basis of forming transfer operators in RN AMG for non-symmetric problems. Building on Lemmas 1 and 2 gives the following result in Lemma 6. Coupled with Conjecture 7, two-grid convergence follows from [18]. Lemma 6 provides a meaningful theoretical motivation for energy minimization as applied to non-symmetric problems.

**LEMMA 6.** *The solution to energy-minimization in the  $A^*A$ - and  $AA^*$ -norms satisfy the non-symmetric strong approximation property in the  $\sqrt{A^*A}$ - and  $\sqrt{AA^*}$ -norms, respectively, that is, for all  $\mathbf{v}$  there exists a  $\mathbf{v}_{c1}, \mathbf{v}_{c2}$  such that*

$$\begin{aligned} \left\| \mathbf{v} - P_{\text{ideal}}^{A^*A} \mathbf{v}_{c1} \right\|_{\sqrt{A^*A}}^2 &\leq \frac{K}{\|A^*A\|} \|\mathbf{v}\|_{A^*A}^2, \\ \left\| \mathbf{v} - P_{\text{ideal}}^{AA^*} \mathbf{v}_{c2} \right\|_{\sqrt{AA^*}}^2 &\leq \frac{K}{\|AA^*\|} \|\mathbf{v}\|_{AA^*}^2. \end{aligned}$$

*Proof.* The proof follows immediately from the equivalence of the WAP( $A^2$ ) and SAP( $A$ ) (Lemma 1) and the convergence of energy-minimization to  $P_{\text{ideal}}$ , in this case  $P_{\text{ideal}}$  for  $A^*A$  and  $AA^*$  (Lemma 2).  $\square$



*Conjecture 7* (Stability). Let  $A$  be nonsingular. The non-orthogonal coarse-grid correction given by transfer operators  $R = (P_{\text{ideal}}^{AA^*})^T$  and  $P = P_{\text{ideal}}^{A^*A}$  is *stable*, that is

$$(23) \quad \left\| P(RAP)^{-1}RA \right\|_{\sqrt{A^*A}} = \left\| I - P(RAP)^{-1}RA \right\|_{\sqrt{A^*A}} = C,$$

for some constant  $C$ , independent of mesh spacing.

*Remark 8.* In the symmetric case,  $R = P^T$  and  $C = 1$ , as the coarse-grid correction is an  $A$ -orthogonal projection. In the non-symmetric case, the stability assumption necessary for two-grid convergence as shown in [18] is primarily to ensure a non-singular and reasonably conditioned coarse-grid operator,  $RAP$ . Conjecture 7 appears to hold in general, but expanding the ideal operators and forming  $RAP$  or the full projection (23) does not provide a clear method to bound its norm. However, in practice the root-node approach over traditional aggregation offers greater stability of the non-orthogonal projection through enforcing the identity over C-points in transfer operators. Specifically, when forming the coarse grid, the identity block in  $R$  and  $P$  help ensure the non-singularity of  $RAP$ .

**5. Numerical results.** In the following, the open source software package PyAMG [6] is used to implement RN AMG<sup>4</sup> and SA AMG<sup>5</sup>. In each routine, cumulative estimates of the SC and CC (see Section 2.4) are tracked in WUs. All costs in the setup phase are included, such as computing the triple matrix product  $RAP$ . In addition, the CC estimates include pre- and post-smoothing, computing and restricting the residual, and coarse-grid correction, but the coarsest-grid direct solve is not included. Measured CC multiplied by the time to perform one SpMV is highly accurate when compared with the wall-clock time of the solve phase. Although measured SC does not track as closely with wall-clock time (likely due to a more complicated setting with memory allocation, conditionals, etc), they still provide a good estimate of setup cost, and differ from wall-clock times by a small constant.

As with many AMG methods, RN AMG has several parameters. The optimal SOC measure is problem dependent, although the evolution measure tends to be more robust for problems with strong anisotropy. Still, the evolution measure does result in higher setup costs, as observed in Section 5.1. Another parameter is that of the degree  $d$  of sparsity pattern for  $P$  (see line 7 in Algorithm 3). Generally, more difficult problems such as anisotropic diffusion require  $d$  to be as large as 4–5. The number of smoothing iterations applied to  $P$  is generally set to  $\lceil 1.5d \rceil$ , as a larger sparsity pattern requires more iterations to minimize the energy of the columns. For SPD problems, the CG-based variant of energy minimization is the least costly, while GMRES is used for non-symmetric problems (GMRES does not offer improved convergence for SPD problems; see Remark 9). Filtering  $P$  also requires user-level decisions (lines 9 and 17 in Algorithm 3). Based on experimentation, a drop tolerance of  $\theta$  is most effective (in contrast to a fixed number of elements per row  $k$ ) and is used in the following tests. Optimal values depend on the connectivity of matrix and are beyond the scope of the numerical tests presented, however  $\theta \in [0.05, 0.25]$  is used in most scenarios.

In each setup phase below, a maximum coarse-grid size of 20 is used, and candidate vectors are improved (line 13, Algorithm 3) with four sweeps of relaxation. For cycling, each test uses an accelerated V-cycle to iterate to a relative residual tolerance of  $10^{-8}$ . Details of the specific relaxation scheme, acceleration type, and other details are specified on a problem-by-problem basis below.

<sup>4</sup>See `root_node_solver`

<sup>5</sup>See `smoothed_aggregation_solver`

### 5.1. 3D-diffusion.

**5.1.1. Filtering and complexity.** The first example demonstrates the effectiveness of the proposed filtering strategy in reducing the cycle and setup complexity of a RN solver. Although filtering is a key component of RN AMG on nearly all problems, it is especially applicable in 3D, where there is high connectivity between nodes, resulting in relatively dense operators. Consider the anisotropic diffusion problem

$$(24) \quad u_{xx} + u_{yy} + 0.001u_{zz} = f.$$

Linear finite elements are used to discretize (24) on an unstructured tetrahedral mesh of the unit cube, with homogeneous, Dirichlet boundaries, yielding a matrix with approximately 2.65M DOFs. While the anisotropy is aligned with the coordinate axis, the unstructured mesh yields a variety of non-grid-aligned anisotropies, known to be more difficult for AMG than the grid-aligned case (see Section 5.2). Table 1 shows complexities and average convergence factors ( $\rho$ ) for solving (24) using various combinations of pre- and post-filtering on  $P$ . A V(1,1)-cycle with symmetric Gauss-Seidel relaxation and CG acceleration is used. The evolution SOC measure is used, with a drop tolerance of 4.0, and CG energy minimization with  $d = 4$ .

Pre-filter $\theta$	–	0.1	0.2	–	–	0.1	0.2
Post-filter $\theta$	–	–	–	0.1	0.2	0.1	0.2
SC	1157.9	161.7	123.2	581.5	432.9	156.2	129.8
OC	3.0	1.4	1.3	1.8	1.5	1.4	1.3
CC	18.9	8.0	7.1	10.1	8.0	7.7	6.9
$\rho$	0.52	0.60	0.62	0.51	0.53	0.59	0.61

Table 1: Impact of filtering for the 3D-anisotropic diffusion problem.

Table 1 shows up to an order-of-magnitude reduction in SC and more than a 50% reduction in CC by using filtering. While filtering does not guarantee a reduction in complexity, significant savings are often observed with only a marginal impact on convergence. In some situations, filtering has been observed to not only reduce cost, but also improve convergence (see Section 5.4).

The SC in Table 1 is broken down into four main categories in Table 2. “Aggregation” is the cost of computing the strength matrix  $S$  and forming aggregates  $\mathcal{A}$ , most of which is due to using the evolution measure. “Candidates” refers to relaxing candidate set  $B$  and restricting  $B$  to a coarse level. Column “ $P$ ” refers to the cost of forming the tentative interpolation operator and applying energy-minimization smoothing iterations to construct  $P$ , while column “ $RAP$ ” represents a measure of the triple-matrix product. Each column gives the total cost for the given processes over all levels, measured in WUs.

Filtering  $P$  has a direct impact on all setup components that use  $R$  and  $P$  matrix operations (see “ $RAP$ ” and “Candidates” in Table 2). Consequently, this reduces the cost of restricting the residual and the coarse-grid correction as measured in the CC, along with the cost of smoothing  $P$ , which is the focus of pre-filtering. Table 2 also highlights the high cost of the strength measure in cases when coarse-grid complexity is not contained — i.e. filtering is not used, demonstrating the additional benefit of reduced costs on all subsequent grids through a sparser coarse-grid operator,  $A_c = RAP$ .

Pre-filter $\theta$	Post-filter $\theta$	Aggregation	Candidates	$P$	$RAP$	Total SC
–	–	478.7	24.1	469.7	271.1	1243.6
0.2	–	48.5	10.4	55.1	9.2	123.2
–	0.2	64.1	11.7	338.7	18.5	432.9
0.2	0.2	46.3	10.1	65.4	7.9	129.8

Table 2: Break down of setup cost in WUs for 3D-anisotropic diffusion.

*Remark 9.* Although GMRES energy-minimization targets the SAP and CG the WAP, when applied to SPD problems, there is not a notable difference in convergence. For instance if GMRES is used in Table 1, then the convergence rates change by no more than 0.005 and operator complexities remain essentially the same.

**5.1.2. Modified evolution SOC.** One contribution of this paper are the detailed SC estimates. Precise SC estimates are important when comparing AMG methods — e.g., components such as energy-minimization and the evolution SOC measure may add significantly to the overall setup costs. Moreover, these estimates help identify areas that are opportune for cost reduction.

In considering Table 2, one possible area for cost reduction is the aggregation phase, where the cost of the SOC computation dominates. Traditional SOC measures require only a few work units, usually 2 or 3 times the operator complexity. However, Table 2 shows that evolution-based SOC is a substantial part of the setup phase, which is attributed to the global spectral radius estimate used in weighted Jacobi [39]. This estimate is calculated with an Arnoldi/Lanczos process and costs roughly 15 matrix-vector multiplies on each level. However, alternative methods [4] use  $\ell_1$ -Jacobi relaxation to provide an inexpensive local row-wise weight. This alternative is explored here for the evolution SOC measure.

Table 3 depicts detailed SC results for using the modified  $\ell_1$ -Jacobi evolution measure (cf. Table 2). This change results in similar operator and cycle complexities and nearly identical convergence rates as the original evolution measure. When comparing Tables 2 and 3, it is apparent that the “Aggregation” phase has been significantly reduced in cost. Additionally, when examining the most efficient solvers, where pre-filtering uses  $\theta = 0.2$ , the overall savings in the setup complexity are roughly 20%.

Pre-filter $\theta$	Post-filter $\theta$	Aggregation	Candidates	$P$	$RAP$	Total SC
–	–	449.1	24.4	473.0	249.5	1195.9
0.2	–	25.5	10.4	53.6	7.6	97.1
–	0.2	43.2	11.7	326.7	16.9	398.6
0.2	0.2	22.7	10.2	63.3	6.9	103.1

Table 3: Break down of setup cost in WUs for 3D-anisotropic diffusion.

**5.2. Totally anisotropic diffusion.** Diffusion-like operators are prototypical AMG problems, as they are elliptic and SPD. However, with strong anisotropy, these problems still pose a challenge to multilevel solvers. A 3D example is used in

Section 5.1 to demonstrate filtering. In this example, consider a 2D rotated anisotropic diffusion problem on the domain  $\Omega$  of the form

$$\begin{aligned} -\nabla \cdot Q^T D Q \nabla u &= f \quad \text{for } \Omega = [0, 1]^2, \\ u &= 0 \quad \text{on } \partial\Omega, \end{aligned}$$

where

$$Q = \begin{bmatrix} \cos \psi & -\sin \psi \\ \sin \psi & \cos \psi \end{bmatrix}, \quad D = \begin{bmatrix} 1 & 0 \\ 0 & \epsilon \end{bmatrix}.$$

Here,  $\epsilon$  represents anisotropy and  $\psi$  the angle of rotation from the coordinate axis, which both contribute to the difficulty of this problem [41, 39, 27, 22, 9, 36, 10, 13, 12, 23, 33, 45, 21, 15, 9]. To see this, consider the non-rotated case of  $\psi = 0$ , which has a spectrum of the form

$$\begin{aligned} -\nabla \cdot D \nabla u_{jk} &= \lambda_{jk} u_{jk}, \quad \text{where} \\ u_{jk} &= \sin(j\pi x) \sin(k\pi y), \\ \lambda_{jk} &= \pi^2(j^2 + \epsilon k^2), \end{aligned}$$

for  $j, k \in \mathbb{Z}^+$ . If  $\epsilon = 1$ , then the lowest energy mode is the lowest Fourier mode:  $u_{11} = \sin(\pi x) \sin(\pi y)$ , which is locally representative of all low-energy modes. However, if  $\epsilon \approx 0$ , for small  $j$  there are high-frequency eigenfunctions in the  $y$ -direction ( $k \gg 0$ ) with relatively small eigenvalues, that are no longer represented locally by the lowest Fourier mode. As a result, relaxation schemes are unable to capture these modes, while coarse-grid correction is not equipped to handle such *hidden* high frequency error.

For angles  $\psi$  aligned with the mesh, line relaxation or semi-coarsening along the direction of anisotropy can be used [44]. However, for strong anisotropies,  $\epsilon \approx 0$ , with angles that are not aligned with the mesh, efficient and effective multigrid solvers remain elusive. This section considers a finite element discretization of strongly and *totally* anisotropic diffusion,  $\epsilon = 0.001$  and  $\epsilon = 0$ , respectively, on a unit square with Dirichlet boundary conditions. Totally anisotropic diffusion is particularly challenging as the problem is effectively reduced to a sequence of 1D problems on a 2D domain.

Figure 6 compares RN AMG, CF AMG, and SA AMG applied to totally anisotropic diffusion for various angles. All solves use a symmetric V(1,1)-cycle with symmetric Gauss-Seidel relaxation and CG acceleration. Grid-aligned anisotropies are generally easier to solve than non-grid aligned, hence the excellent convergence factors at  $\theta = \pi/4$  on the structured mesh. In the case of an unstructured mesh, all angles are effectively non-grid aligned resulting in consistent performance across angles.

Classical AMG uses a classical strength measure with drop tolerance of 0.5, and standard CF-splitting and interpolation [42]. Smoothed aggregation uses a symmetric strength measure (with a drop tolerance of 0.0 — i.e. the strength matrix is given by  $A$  and normalized as in (7)) and two iterations of weighted Jacobi interpolation smoothing [48]. The root-node solver in this case uses two steps of an evolution strength measure (with a drop tolerance of 4.0), along with six iterations of CG energy-minimization smoothing of  $P$ . For the energy minimization, a degree  $d = 4$  sparsity pattern is used with filtering,  $\theta_{\text{pre}} = \theta_{\text{post}} = 0.1$ . Two and three Jacobi smoothing iterations were applied to the SA AMG solver in an attempt to mimic the expanded sparsity pattern used in RN AMG, but the convergence with respect to CC did not improve.

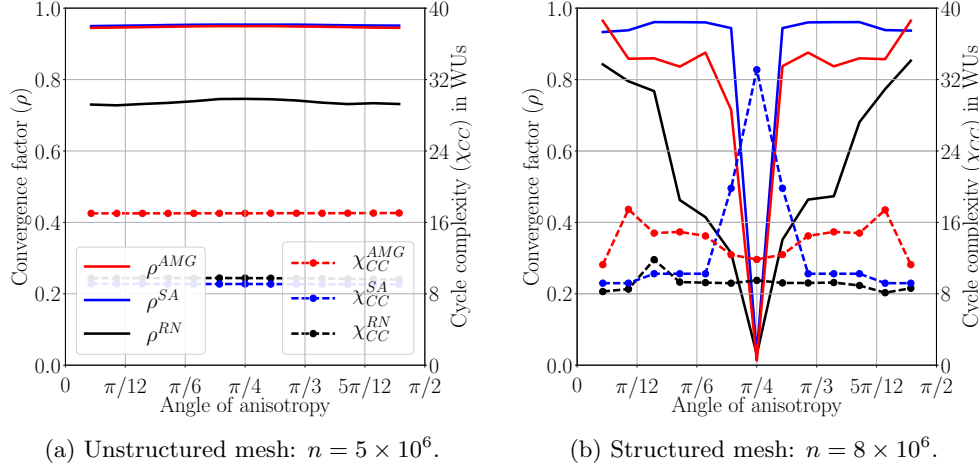


Fig. 6: Results for totally anisotropic diffusion ( $\epsilon = 0$ ), with angles in  $(0, \pi/2)$ . A structured grid of size  $2000 \times 2000$  and an unstructured mesh with resolution  $h \approx 1/2000$  are used. Legend entries in (a) apply to (b) as well. RN AMG, shown in black, outperforms SA AMG and CF AMG in all cases.

Figure 6 highlights the effectiveness of RN AMG over all angles with moderate operator and cycle complexities. For time-to-solution with respect to floating point operations and wall-clock time, RN AMG achieves between  $3\text{--}30\times$  speed-up in comparison to SA AMG and CF AMG on a structured mesh and unstructured mesh, with moderate cycle complexities in all cases. It should be noted that performance of SA AMG improves when using the modified evolution SOC measure introduced in Section 5.1.2, but RN AMG still performs  $2\text{--}3\times$  faster with respect to time and complexity. This is a notable achievement in performance for this problem, as anisotropic diffusion remains a significant challenge to most solvers.

Figure 7 demonstrates the scaling of convergence factors as problem size increases for  $V(1,1)$ - and  $W(1,1)$ -cycles. In the case of  $\epsilon = 0.001$ , V-cycle convergence factors asymptote and scale perfectly, independent of  $h$ , on structured and unstructured meshes, up to 25 million unknowns; the SC and CC also scale, but are not shown. However, in the case of  $\epsilon = 0$  there is a slow growth in the convergence factor as the problem size increases for both V- and W-cycles. To analyze, consider a convergence factor,  $\rho(h)$ , dependent on spatial step size  $h$ :

$$(25) \quad \rho = \bar{\rho}(1 - ah^q),$$

where  $q = 1$  in the case of linear finite elements,  $a$  is some constant, and  $\bar{\rho}$  is the asymptotic convergence factor,  $\lim_{h \rightarrow 0} \rho = \bar{\rho}$ . A log of (25) and an expansion yields  $\log(1 - ah) = -ah + O(h^2)$  or  $-\log(\rho) = -\log(\bar{\rho}) + ah$ . A linear fit on the three smallest step sizes in Figure 7 for  $\epsilon = 0$  leads to the following asymptotic convergence factors

$$\text{structured: } \rho \rightarrow 0.82, \quad \text{and unstructured: } \rho \rightarrow 0.88.$$

Although an asymptotic convergence factor of approximately 0.88 is relatively slow,

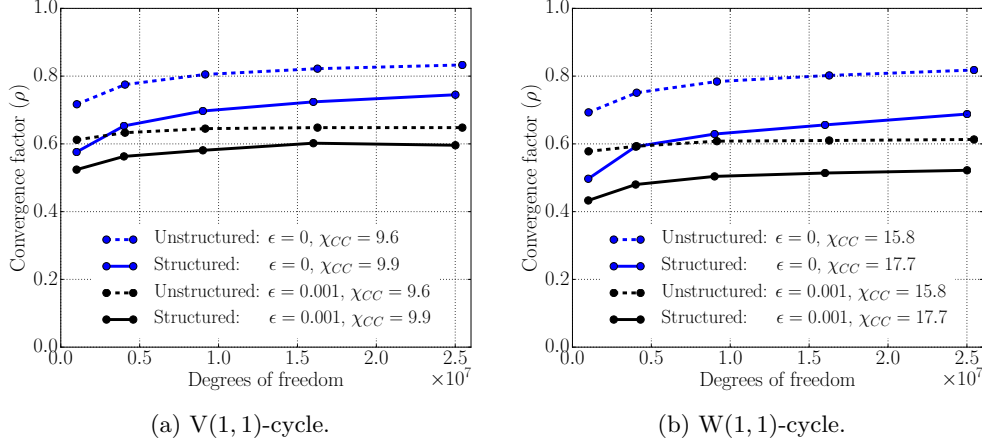


Fig. 7: Scaling results for RN AMG for anisotropic diffusion with  $\epsilon = 0.001$ ,  $\epsilon = 0$ , and  $\theta = 3\pi/16$ . Cycle complexity,  $\chi_{CC}$ , is shown and is constant for all problem sizes.

*scalable* convergence of totally anisotropic diffusion on unstructured meshes has not been achieved by other AMG methods.

In practice, using W-cycles over V-cycles should increase the accuracy of the coarse-grid correction. However, Figure 7 reveals W-cycles offer only minor improvements. This indicates that the algebraically smooth error is not well represented by the coarse grid, and that improved strength measures and coarsening routines [39, 22, 9, 36, 13, 12] may improve the coarse grid, thereby improving convergence for this problem.

**5.3. Recirculating flow (non-symmetric).** One of the benefits of the root-node approach is the ability to handle a variety of problems, including non-symmetric problems and systems, without redesign of methodology and implementation. In this section, a standard recirculating flow example known as the *double-glazing problem* is used [24], which models temperature distribution over a domain when an external wall is hot. The governing PDE is given by

$$(26) \quad -\epsilon \nabla \cdot \nabla u + \mathbf{b}(\mathbf{x}) \cdot \nabla u = f,$$

where  $\epsilon = 0.005$  and wind is given by  $\mathbf{b}(\mathbf{x}) = [2x_1(1 - x_0^2), -2x_0(1 - x_1^2)]$ . Dirichlet boundaries are imposed on the domain  $[0, 1] \times [0, 1]$ , where  $u = 0$  on the north, south and west sides of the domain, and  $u = 1$  on the east, leading to boundary layers near corners with discontinuities.

A standard Galerkin finite element method (GFEM) based on a regular triangular mesh is used, resulting in a non-symmetric discrete linear system. Multigrid theory for non-symmetric problems is less developed in comparison to the symmetric case; however, SA AMG has been extended to non-symmetric problems [43] and is used in this section (see Algorithm 3).

Each example uses a V(1,1)-cycle of weighted Jacobi with GMRES acceleration. While Jacobi relaxation is not guaranteed to converge for a non-symmetric problem, it remains around half the cost of using relaxation on the normal equations. In Section 5.4 an upwind discretization is considered which requires relaxation on the



normal equations for effective convergence. For SA AMG, a classical strength measure (with drop tolerance 0.25) is used along with one and three steps of Jacobi smoothing steps applied to  $P$ , labeled  $SA_1$  and  $SA_3$ , respectively. The symmetric traditional SA strength measure is not used due to the non-symmetry of the problem. Two steps of the evolution measure (with drop tolerance 3.0) is used for RN AMG along with two iterations of GMRES energy minimization for  $P$  with  $d = 1$  (labeled  $RN_1$ ), and five iterations of GMRES energy minimization for  $P$  with  $d = 3$  (labeled  $RN_3$ ).

Table 4 demonstrates optimal results are achieved for this example with no filtering and a small sparsity pattern ( $d = 1$ ). This agrees with practical experience: generally it is effective to increase the filtering tolerance as the degree of the sparsity pattern for  $P$  increases, or as the connectivity of matrix  $A$  increases. This is observed in the 3D-anisotropic diffusion problem, where high connectivity and a  $d = 4$  sparsity pattern allows for a large  $\theta = 0.2$ . If the sparsity pattern increases in distance from the root-node, or the matrix is highly connected, then it is likely there are entries that are not critical to performance and are candidates for removal.

$\theta$	$d$	2000 $\times$ 2000				3000 $\times$ 3000				4000 $\times$ 4000			
		SC	OC	CC	$\rho$	SC	OC	CC	$\rho$	SC	OC	CC	$\rho$
–	$SA_1$	72	1.4	5.2	0.74	71	1.4	5.2	0.82	71	1.4	5.2	0.88
	$SA_3$	229	2.3	11.2	0.96	230	2.3	11.2	0.96	227	2.3	11.2	0.93
–	$RN_1$	98	1.4	5.1	0.46	96	1.4	5.0	0.50	95	1.4	4.9	0.45
	$RN_3$	403	2.5	9.7	0.68	407	2.4	9.4	0.63	405	2.3	9.2	0.76
0.1	$RN_1$	126	1.4	5.1	0.52	125	1.4	5.0	0.56	124	1.4	4.9	0.56
	$RN_3$	284	1.8	6.7	0.53	287	1.8	6.7	0.52	285	1.8	6.6	0.63

Table 4: Non-symmetric SA AMG and RN AMG for the recirculating flow problem.

Classical CF AMG is not designed for non-symmetric problems, making RN AMG the clear choice for a problem such as this. Root-node AMG achieves more than a  $6\times$  speed-up over SA AMG for the largest problem size considered, *with a lower CC*, and only slightly larger SC. Furthermore, RN AMG convergence factors with degree  $d = 1$  appear to have reached an asymptote, while SA AMG is still demonstrating a steady increase with problem size.

*Remark 10.* With the recirculating flow, as the grid-size ( $h$ ) approaches zero, there are two competing factors that contribute to the numerical difficulty of the problem. As  $h \rightarrow 0$ , the diffusive part of the problem,  $-\epsilon \nabla \cdot \nabla$ , becomes increasingly dominant, as the discretization scales like  $\frac{1}{h^2}$ , while  $\mathbf{b}(\mathbf{x}) \cdot \nabla$  scales like  $\frac{1}{h}$ . The resulting linear system is more symmetric and diffusion-like, which is preferable for AMG. However, as  $h \rightarrow 0$ , convergence factors often increase to an asymptotic value (see (25) and Figures 7a and 7b), due to smaller eigenvalues and an increasing number of levels in the AMG hierarchy. Together, these factors correspond to the so-called *half-grid Reynolds number* or *cell Reynolds number*,  $R_h = \frac{|\mathbf{b}|h}{2\epsilon}$ , where convergence factors are expected to be consistent for a fixed  $R_h$ , and degrade for  $R_h \gg 1$ . Figure 8 demonstrates this phenomenon. Convergence factors asymptote for each half-grid Reynolds number as  $h \rightarrow 0$ . In this case, convergence degrades by a factor of 10 when increasing from  $R_h = 1.25$  to  $R_h = 2.5$ ; thus it is faster to solve a refined problem with  $R_h = 1.25$  (and several times as many DOFs), rather than a system with  $R_h = 2.5$ .

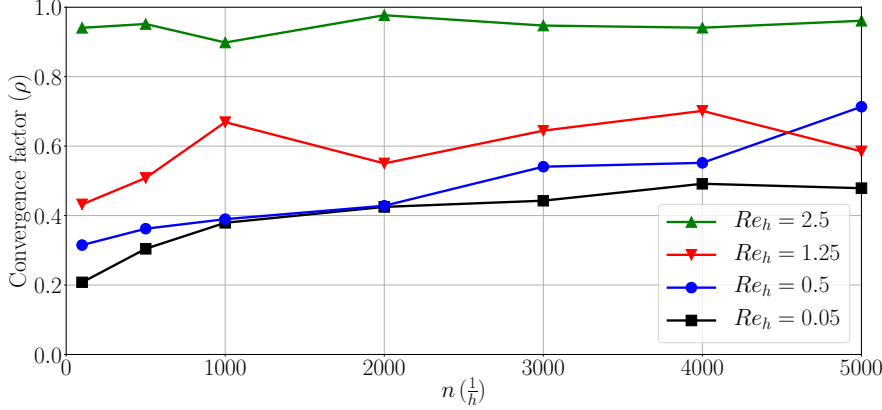


Fig. 8: RN AMG convergence factor as a function of  $h$ , for fixed half-grid Reynolds numbers,  $Re_h \in \{0.05, 0.5, 1.25, 2.5\}$ .

Based on the scalability of RN AMG performance for fixed  $Re_h$ , with the appropriate grid-size, most convection-diffusion problems as in (26) should exhibit scalability in convergence. Of course, this relies on the diffusion operator dominating the discrete system. As a limiting case, RN AMG is applied to the steady-state transport equation in the following section.

**5.4. Upwind transport (non-symmetric).** In this section, a highly non-symmetric upwind discretization of the steady-state transport equation is considered. Define the domain as  $\Omega = (0, 1) \times (0, 1)$ , with constrained inflow boundaries  $\Gamma_{\text{in}} = \{(x, y) : x = 0 \text{ or } y = 0\}$ , and free outflow boundaries  $\Gamma_{\text{out}} = \{(x, y) : x = 1 \text{ or } y = 1\}$ . The steady-state transport equation is then given as

$$(27) \quad \begin{aligned} \mathbf{b}(x, y) \cdot \nabla u + c(x, y)u &= f(x, y) & \Omega, \\ u &= g(x, y) & \Gamma_{\text{in}}. \end{aligned}$$

Root-node AMG performance is demonstrated for variations in  $\mathbf{b}(x, y)$  and  $c(x, y)$ , using an upwind lumped bilinear discontinuous (LBLD) finite element discretization [34]. A discontinuous discretization is used to allow for discontinuities in material coefficient  $c(x, y)$ , and to account for the fronts that develop in hyperbolic-type PDEs. The transport problem, (27), results in a highly non-symmetric (nearly lower-triangular in the proper ordering) matrix  $A$ , for which traditional relaxation schemes such as Jacobi and Gauss-Seidel diverge, even for small problems. Thus two sweeps of Gauss-Seidel on the normal equations are chosen as the relaxation scheme, which contributes to a larger CC.

One anomalous feature of the LBLD discretization applied to (27) is that the best convergence rates are obtained with aggressive filtering,  $\theta_{\text{pre}} = \theta_{\text{post}} = 0.4 - 0.5$ . The rationale is that a large degree sparsity pattern with aggressive filtering results in a sparsity pattern that is long and narrow, following the direction of flow. For a hyperbolic PDE with the solution following characteristic curves, effective sparsity patterns for a given column of  $P$  will align with the characteristic associated with the root-node. However, in practice this requires either *a priori* knowledge of the problem to motivate aggressive filtering, or experimentation. For most problems,

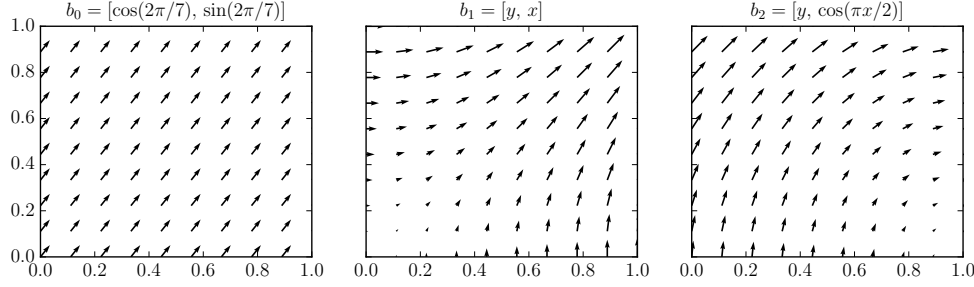


Fig. 9: Vector fields of flow functions,  $\mathbf{b}(x, y)$  considered in (27), with inflow boundaries  $\Gamma_{in} = \{(x, y) : x = 0 \text{ or } y = 0\}$ .

filtering values of  $\theta_{pre} = \theta_{post} \in [0.05, 0.15]$  are effective choices, increasing to  $\theta_{pre} = \theta_{post} \in [0.1, 0.25]$  for problems with high connectivity — e.g. 3d-diffusion as discussed in Section 5.1. However, the LBLD results demonstrate that further work is needed to algebraically determine the optimal interpolation sparsity pattern for a given problem.

Root-node AMG is applied to the LBLD discretization of (27) for three different material coefficients  $c(x, y)$ , as shown in (28), and three directional functions as shown in Figure 9.

$$\begin{aligned}
 \text{Constant: } c_1(x, y) &= 1 \\
 \text{Square in square (SnS): } c_2(x, y) &= \begin{cases} 10^4 & x, y \in [0.25, 0.75] \\ 10^{-4} & x, y \notin [0.25, 0.75] \end{cases} \\
 \text{Split: } c_3(x, y) &= \begin{cases} 10^{-4} & x < 0.5 \\ 10^4 & x \geq 0.5 \end{cases}
 \end{aligned}
 \tag{28}$$

Results are shown in Table 5 for a discretization with 4,000,000 DOFs, sufficiently large to observe asymptotic convergence factors. A classical SOC with drop tolerance  $\theta = 0.35$  is used, and GMRES energy-minimization applied to a degree four sparsity pattern for interpolation and restriction operators in RN AMG, with filtering  $\theta_{pre} = \theta_{post} = 0.45$ . The multigrid solver is used as a preconditioner for GMRES, and problems solved to  $10^{-8}$  residual tolerance. Candidate vectors are taken as the constant and are *not* improved for this problem, as convergence tended to degrade.

$\mathbf{b}(x, y)$		$[\cos(2\pi/7), \sin(2\pi/7)]$			$[y, x]$			$[y, \cos(\pi x/2)]$		
$c(x, y)$		1	SnS	Split	1	SnS	Split	1	SnS	Split
RN	SC	164	147	130	125	105	107	126	112	109
	OC	1.88	1.76	1.66	1.92	1.83	1.78	1.91	1.83	1.77
	CC	17.9	16.7	15.8	18.2	17.4	17.0	18.2	17.4	16.8
	$\rho$	0.79	0.74	0.75	0.86	0.84	0.85	0.87	0.87	0.86

Table 5: Non-symmetric RN AMG for variations on the 2d steady-state transport equation (27). Matrices have 4,000,000 DOFs and 15,986,004 nonzeros.

Although convergence factors in the case of non-constant flow are higher than

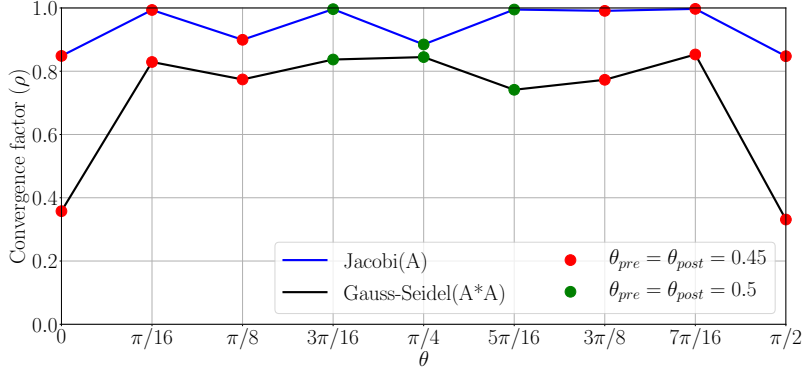


Fig. 10: Convergence factors for GMRES-accelerated RN AMG applied to the steady-state transport equation, as a function of the direction of flow  $\mathbf{b}(x, y) = (\cos(\theta), \sin(\theta))$ . Results are shown for Gauss-Seidel on the normal equations (GSNE) relaxation as well as Jacobi, to demonstrate why normal-equation relaxation is necessary for highly non-symmetric problems. Other solver parameters are fixed for all tests, with the exception of filtering, which is modified slightly for improved convergence on angles close to  $\pi/4$ . Convergence with GSNE is in the range  $[0.75, 0.85]$  for interior angles, with significant improvement close to the boundaries.

desired (worst case  $\rho \approx 0.87$  for RN AMG), it is encouraging that AMG methods are able to solve upwind discretizations of a hyperbolic PDE with discontinuous and non-constant coefficient functions. Convergence factors of SA AMG quickly approached one, and did not converge in 500 iterations.

Often when considering transport-type problems, “sweeps” are performed, where the problem is discretized in angle and a linear solve performed for each angle. Transport sweeps are an important part of the DSA algorithm for models of neutral particle transport [2]. For this reason, it is of interest to demonstrate RN AMG’s capability on the entire spectrum of angles, similar to anisotropic diffusion as considered in Section 5.2. Figure 10 shows convergence factors of RN AMG as a function of direction of flow,  $\theta \in [0, \pi/2]$ , as applied to the LBLD discretization of (27).

**5.5. Elasticity example (systems problem).** Algebraic multigrid for systems of PDEs is an important topic due to the vast number of problems formulated this way. Although there are some problem-specific extensions to CF AMG for systems, such as elasticity [5], where rigid body modes are incorporated into a modified method, CF AMG is not applicable to systems in a general setting. In contrast, SA AMG specifically targets systems-based problems such as elasticity, and performs well in many cases. Here, the suitability of RN AMG for systems is illustrated through a comparison with SA AMG for a standard elasticity model problem, a problem for which SA AMG was designed.

The test problem is isotropic linearized elasticity defined by

$$(29) \quad -\operatorname{div}(\lambda \operatorname{tr}((\nabla \mathbf{u} + \nabla \mathbf{u}^T)/2) \mathbf{I} + \mu(\nabla \mathbf{u} + \nabla \mathbf{u}^T)) = f,$$

where  $\lambda$  and  $\mu$  are the Lamé parameters,  $\mathbf{I}$  is the identity matrix, and  $\operatorname{tr}()$  is the

trace function. Example two from the MFEM package [1] is used to discretize (29) using a regular mesh of a beam that is eight times longer than it is wide (`data/beam-tri.mesh`). Three material choices, of increasing difficulty for AMG, are considered: a steel beam, with Young’s modulus and Poisson ratio of  $E = 180\text{E}9$  and  $\nu = 0.30$ ; a rubber beam, with  $E = 0.1\text{E}9$  and  $\nu = 0.499$ ; and a more difficult rubber beam, where  $\nu = 0.4999$ . In general, the closer  $\nu$  is to 0.5, the more difficult the system is for AMG to solve. The east side of the beam is fixed to a wall with  $\mathbf{u} = \mathbf{0}$  Dirichlet boundary conditions. Neumann conditions are applied on other boundaries, with  $\mathbf{u} \cdot \mathbf{n} = f$ , where  $f = 0$  for the north and south sides, and  $f = -0.01$  on the west side, representing a downward force. Standard linear triangular finite elements are used to produce an SPD system with two variables at each spatial location and the corresponding block sparse row matrix with  $2 \times 2$  blocks.

Each example uses a  $V(1,1)$ -cycle of symmetric block Gauss-Seidel with CG acceleration. Both RN AMG and SA AMG use the classical strength measure with a drop tolerance of 0.5. Root-node AMG does not use the evolution measure here because the classic measure performed well and lowered the setup complexity.<sup>6</sup> The terms  $\text{SA}_k$  and  $\text{RN}_k$  again refer to the degree of interpolation smoothing. For  $\text{RN}_4$ , 6 iterations of CG energy minimization are used, while for  $\text{RN}_1$ , 2 iterations are used. Experimentation indicates that only pre-filtering is needed, so  $\theta$  values only refer to pre-filtering. The candidate vectors  $B_0$  are the three rigid-body-modes in 2D, two translations and one rotation.

Table 6 gives the numerical results for three problem sizes, from 256K to 4M DOFs. Various filtering values produce qualitatively similar results, so a single result is reported. All methods converge well for the steel beam, with the 0.5 convergence rate for  $\text{RN}_1$  being offset by the low operator complexity of 1.2. The rubber beams are considerably harder, with the most difficult beam problem yielding convergence rates well above 0.9.

Comparing the methods with each other, the results reveal that  $\text{RN}_1$  converges slowly in comparison to  $\text{SA}_1$  or  $\text{RN}_4$ . This is because the three candidate vectors provided as interpolation constraints require a richer sparsity pattern than  $d = 1$  in order to satisfy the constraints.

Another important point is that RN AMG with  $d = 4$  does achieve a better convergence rate and smaller operator complexity than SA AMG but at the cost of a larger SC. Taking the largest problem size for test 2, with 4M DOFs (a grid size  $4096 \times 512$ ), the difference in convergence rate yields 87 iterations for  $\text{RN}_4$  versus 107 iterations for  $\text{SA}_1$ . However even with this faster convergence, SA AMG uses the fewest overall work units.<sup>7</sup> SA AMG similarly provides slightly better overall work units for setup and solve with test 3. In conclusion, this section shows that RN AMG is a viable approach to solving systems, and is competitive with SA AMG for the considered elasticity problem, for which SA AMG was originally designed. Applying RN AMG to systems is still an active research topic, focusing on issues such as reducing the setup cost and considering more complicated examples.

<sup>6</sup>If the three elasticity test cases are re-run for  $\text{RN}_4$  using the evolution strength measure, then the convergence rate degrades by about 0.01–0.02 and the operator complexities remain similar, i.e., RN AMG using the evolution measure converges similarly to SA using the classic measure. While SOC for systems is an active topic of AMG research and not well-understood, one possible reason for this small difference between the two measures is that the “anisotropy” in the beam is grid-aligned, and the classical measure is known to handle grid-aligned anisotropies well.

<sup>7</sup>It is possible, given the faster creep in convergence rate suffered by SA AMG, that RN AMG could outperform overall as the problem size increases.

Test	Grid size:		$1024 \times 128$				$2048 \times 256$				$4096 \times 512$			
	$\theta$	$d$	SC	OC	CC	$\rho$	SC	OC	CC	$\rho$	SC	OC	CC	$\rho$
1	0.1	SA <sub>1</sub>	105	1.6	9.8	0.23	106	1.7	9.8	0.26	105	1.7	9.8	0.26
		RN <sub>1</sub>	84	1.2	6.9	0.50	84	1.2	6.9	0.49	85	1.2	6.9	0.51
		RN <sub>4</sub>	226	1.4	8.2	0.19	226	1.4	8.2	0.21	225	1.4	8.2	0.19
2	0.1	SA <sub>1</sub>	104	1.6	9.6	0.83	106	1.6	9.8	0.84	104	1.7	9.8	0.84
		RN <sub>1</sub>	77	1.2	6.7	0.92	77	1.2	6.7	0.94	78	1.2	6.7	0.93
		RN <sub>4</sub>	221	1.4	8.2	0.80	223	1.4	8.2	0.80	223	1.4	8.3	0.81
3	0.1	SA <sub>1</sub>	104	1.6	9.7	0.93	105	1.6	9.8	0.94	105	1.7	9.8	0.93
		RN <sub>1</sub>	82	1.2	6.7	0.97	82	1.2	6.7	0.97	81	1.2	6.7	0.97
		RN <sub>4</sub>	228	1.4	8.2	0.92	229	1.4	8.3	0.92	229	1.4	8.3	0.92

Table 6: Application of SA AMG and RN AMG to three linearized elasticity problems. The tests are progressively more difficult, with Test 1 being a steel bar with Poisson ratio  $\nu = 0.30$ , Test 2 being a rubber bar with  $\nu = 0.499$ , and Test 3 being a rubber bar with  $\nu = 0.4999$ .

*Remark 11.* It is important to note that the ideal  $P$  in equation (20) is only ideal for the specific two-grid setting outlined in [25]. This ideal  $P$  is not guaranteed to be ideal for multilevel methods, or even for all two-grid methods. However, the ideal  $P$  is still a useful target for interpolation as evidenced by the experiments in this work and by the fact this it is ideal under the assumptions in [25]. Along with approximating  $P_{\text{ideal}}$ , RN AMG combines other proven approaches to constructing quality interpolation operators, interpolation smoothing and guaranteed interpolation of known algebraically smooth modes.

As an example of how this combination of strategies makes RN AMG more robust than using  $P_{\text{ideal}}$ , two-grid experiments for the three test cases from Table 6 are run on a small mesh yielding 4.5K DOFs. For the steel beam (Test 1), the convergence factors are 0.19, 0.17 and 0.18, for SA<sub>1</sub>, RN<sub>4</sub> and ideal  $P$ , respectively. Moving to the rubber beam (Test 2), the convergence factors are 0.80, 0.79 and 0.97. Finally for the harder rubber beam (Test 3), the convergence factors are 0.92, 0.91, 0.98. Experimentally, it appears that as the Poisson ratio approaches 0.5, convergence  $P_{\text{ideal}}$  degrades significantly faster than the interpolation operators produced by SA AMG and RN AMG.

**6. Conclusions.** The presented RN AMG methodology has proven successful on a wide variety of problems. The proposed filtering of the interpolation sparsity patterns before and after energy-minimization smoothing iterations greatly reduces setup and cycle complexity, and at times improves convergence, making RN AMG a robust solver for difficult SPD problems, non-symmetric problems, and systems-related problems. Total complexity remains reasonable for all of the examples tested, and tends to scale with the size of the problem. Providing such total complexity estimates, including the setup phase, is a specific contribution of this work.

One particularly difficult problem is strongly anisotropic diffusion, which RN AMG is able to solve effectively in comparison to other AMG methods. Because the focus here is on a general root-node methodology, coupled with energy-minimization smoothing of  $P$ , minimal testing is done in coupling RN AMG with other advance-



ments in AMG. In the case of strongly anisotropic diffusion, many other works consider modified SOC and advanced coarsening schemes in order to better capture the anisotropy [39, 22, 9, 36, 13, 12, 9], thereby improving convergence rates. Future work involves coupling RN AMG with a larger variety of SOC measures, aggregation routines, and sparsity patterns for  $P$ , to further improve convergence on anisotropic problems. Root-node is also shown effective at solving convection-diffusion, and a discontinuous, upwind discretization of a hyperbolic PDE (steady-state transport), demonstrating that AMG need not be limited to elliptic problems.

Parallelization of RN AMG is straightforward. The basic computational kernels are typically available in AMG codes (e.g., matrix-matrix multiply), are light-weight (e.g., filtering entries), or purely local in computation (e.g., the row-wise projection operation (8)).

New theoretical motivation for coupling energy minimization with RN AMG is provided in the symmetric and non-symmetric setting. Proving Conjecture 7 would complete the two-grid convergence proof for ideal operators in the non-symmetric case. However, the stability constraint is of limited use in practice, as it is not directly approximated like the weak and strong approximation properties. Further work on developing energy-based, non-symmetric convergence theory is important and ongoing work in understanding how to construct AMG solvers for non-symmetric systems.

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