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# A Three-Level Extension for Fast and Robust Overlapping Schwarz (FROSch) Preconditioners with Reduced Dimensional Coarse Space

Alexander Heinlein, Axel Klawonn, Oliver Rheinbach, and Friederike Röver

## 1 Fast and Robust Overlapping Schwarz preconditioners

The Fast and Robust Overlapping Schwarz framework [7, 8], which is part of the Trilinos Software library [18], contains a parallel implementation of the *generalized Dryja–Smith–Widlund* (GDSW) preconditioner. The GDSW preconditioner is a two-level overlapping Schwarz domain decomposition preconditioner [17] with an energy minimizing coarse space [3, 4]. It is constructed based on a domain decomposition of the computational domain  $\Omega$  into  $N$  nonoverlapping subdomains  $\{\Omega_i\}_{i=1,\dots,N}$ . These are then extended by  $k$  layers of elements, resulting in a corresponding overlapping domain decomposition  $\{\Omega'_i\}_{i=1,\dots,N}$ . The two-level GDSW preconditioner can then be written as

$$M_{\text{GDSW}}^{-1} = \underbrace{\Phi K_0^{-1} \Phi^T}_{\text{coarse level}} + \underbrace{\sum_{i=1}^N R_i^T K_i^{-1} R_i}_{\text{first level}}, \quad (1)$$

where  $\Phi$  contains the coarse basis functions. Contrary to the classical approach, where the coarse basis functions are chosen as nodal finite element functions on

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a coarse triangulation, for the GDSW preconditioner, these are chosen as discrete harmonic extensions of certain interface functions  $\Phi_\Gamma$  to the interior of each subdomain. In particular, the functions  $\Phi_\Gamma$  are restrictions of the null space of the global Neumann matrix to the vertices, edges, and faces, which form a nonoverlapping decomposition of the domain decomposition interface. The matrix  $K_0 = \Phi^T K \Phi$  is the coarse matrix and the matrices  $K_i = R_i K R_i^T$ ,  $i = 1, \dots, N$ , correspond to the overlapping subdomain problems on the first level. The local subspaces corresponding to the overlapping subdomains are denoted as  $V_1, \dots, V_N$ , and the GDSW coarse space is denoted by  $V_0$ . For scalar elliptic problems, the condition number is bounded by

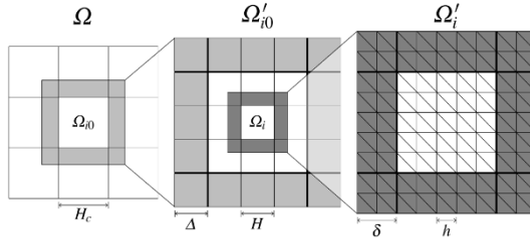
$$\kappa(M_{\text{GDSW}}^{-1}K) \leq C \left(1 + \frac{H}{\delta}\right) \left(1 + \log\left(\frac{H}{h}\right)\right)^2, \quad (2)$$

where  $C$  is a constant independent of the finite element size  $h$ , the size  $H$  of the non-overlapping subdomains, and the width of the overlap  $\delta = kh$ ; see [3]. The GDSW coarse space can be constructed in an algebraic fashion, i.e., without geometric information. For a further reduction of the coarse space, the FROSch framework provides an implementation of a reduced dimensional coarse space (RGDSW) [11]. For the reduced dimensional GDSW coarse space, the basis functions are constructed from nodal interface functions. Two options are currently available in FROSch: a fully algebraic version (*Option 1*) [5, 11], where the interface values are defined through the number of adjacent vertices, or the less algebraic version (*Option 2.2*) [5, 11], where the interface values are defined through the distance to the adjacent vertices; cf. [5, 11]. In general, the two options result in different partitions of unity. The interior values of each subdomain are determined as in the classical GDSW approach.

## 2 Three-level extension

For a large number of subdomains, the coarse problem of the two-level (R)GDSW preconditioners may become too large to be solved by a sparse direct solver. As in the three-level BDDC methods [19], we can resolve this by applying the GDSW preconditioner recursively to the coarse problem [9, 10]. This technique can be extended to a multi-level version, as in multi-level BDDC [1, 16] (which compete with inexact FETI-DP methods [13]), multilevel Schwarz methods [14, 15], or multigrid methods. We only discuss the three-level extension in this paper.

To apply the (R)GDSW preconditioner to the coarse problem, we need to define an additional layer of decomposition. We therefore decompose the domain into non-overlapping subregions  $\Omega_{i0}$  of diameter  $H_c$ , whereas each subregion is a union of subdomains. To obtain overlapping subregions  $\Omega'_{i0}$ , we extend each subregion by recursively adding layers of subdomains, as we do with finite elements on the subdomain level; see Figure 1. We denote the subregion overlap by  $\Delta$ . The notation on the subdomain level is kept consistent with the two-level method.



**Fig. 1:** Structured decomposition of an exemplary two-dimensional computational domain  $\Omega$  into nonoverlapping subregions  $\Omega_{i0}$ (left), a zoom into one overlapping subregion  $\Omega'_{i0}$  consisting of subdomains  $\Omega_i$  (middle), and a zoom into one overlapping subdomain  $\Omega'_i$  (right). Each level of zoom corresponds to one level of the preconditioner; image from [9].

We define the three-level GDSW preconditioner [9, 10] by

$$M_{\text{GDSW-3L}}^{-1} = \Phi \left( \overbrace{\Phi_0 K_{00}^{-1} \Phi_0^T}^{\text{third level}} + \sum_{i=1}^{N_0} \overbrace{R_{i0}^T K_{i0}^{-1} R_{i0}}^{\text{second level}} \right) \Phi^T + \sum_{j=1}^N \overbrace{R_j^T K_j^{-1} R_j}^{\text{first level}}, \quad (3)$$

where the first level and the matrices  $\Phi$  are defined as in the two-level method and where  $K_{00} = \Phi_0^T K_0 \Phi_0$  and  $K_{i0} = R_{i0}^T K_0 R_{i0}$ . The restriction operators, restricting to the overlapping subregions  $\Omega'_{i0}$ , are defined as  $R_{i0} : V^0 \rightarrow V_i^0 := V^0(\Omega'_{i0})$  for  $i = 1, \dots, N_0$ . The respective coarse space is denoted as  $V_{00}$  and spanned by the coarse basis functions  $\Phi_0$ .

### 3 Implementation

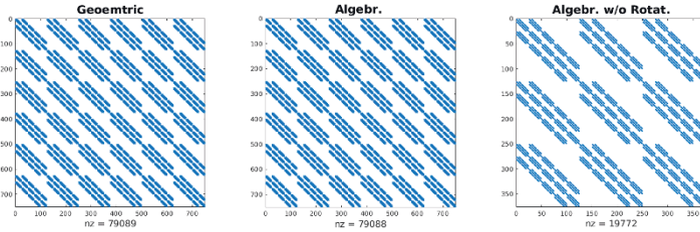
The Fast and Robust Overlapping Schwarz (FROSch) framework [7, 8] is part of the package ShyLU from the Trilinos software library [18]. It contains parallel implementations of the GDSW and RGDSW preconditioners based on the Trilinos linear algebra interface Xpetra; it enables the use of both Trilinos linear packages Epetra and Tpetra. To test the three-level extension to the FROSch implementation, we considered a linear elasticity model problem on the unit cube  $[0, 1]^3$  with homogenous Dirichlet boundary condition on  $\partial\Omega$ . We use piecewise trilinear finite elements and a structured decomposition of the computational domain. To assemble the stiffness matrix we apply the Trilinos package Galeri. Here, each process owns the same number of rows of stiffness matrix resulting in different subdomain sizes. We use a generic right-hand side vector in which each entry is set to one. If the coarse space is constructed as described in Section 1, the columns of the matrix  $\Phi$  will be a generating set of the coarse space. However, for our model problem, the columns will not be linearly independent and, hence, not form a basis of the coarse space. This is because the restriction of the six-dimensional null space, consisting of

translations and linearized rotations, to an interface component may yield linearly dependent vectors. For instance, the restriction of the null space to a single vertex yields only a three dimensional space. In order to make sure that the coarse matrix  $K_0$  is invertible, we have to deal with this in our implementation. In particular, before building  $K_0$ , we replace linearly dependent coarse functions by null vectors until all other basis functions are linearly independent; in order to identify linear dependencies, we perform local orthonormalization using LAPACK's SGEQRF routine for computing a QR factorization using Householder transformations. This procedure yields zero rows and columns in  $K_0$ . Therefore, in order to make  $K_0$  invertible, we finally replace those rows and columns by the corresponding unit vectors, leaving a one on the diagonal and zeros otherwise. This also has the nice side effect that the size of the coarse matrix is always the number of interface components times the dimension of the null space. The coarse level is decomposed into subregions in an unstructured way using the Parallel Hypergraph and Graph Partitioning (PGH) from the Trilinos package Zoltan2 [20]; see also [12]. As a Krylov iteration method, we apply the preconditioned conjugate gradient method (PCG) provided by the Trilinos package Belos (BelosPseudoBlockCG). The implementation offers a condition number estimate using the tridiagonal matrix constructed in the Lanczos process. We use the relative stopping criterion  $\|r^k\|_2/\|r^0\|_2 \leq 10^{-6}$ , where  $r^k$  is the residual in the  $k$ -th iteration step and  $r^0$  is the initial residual. For all tests, we chose  $20^3 * 3$  rows of the stiffness matrix for each process and approximately  $8^3$  subdomains per nonoverlapping subregion. The overlap is obtained by extending each subdomain by one layer of elements and by extending each subregion by one layer of subdomains. We performed all numerical tests on the GCS supercomputer SuperMUC-NG. The INTEL 19.0 compiler is used. The sparse linear subproblems arising in the preconditioner are solved using the sparse direct linear solver PardisoMKL [2].

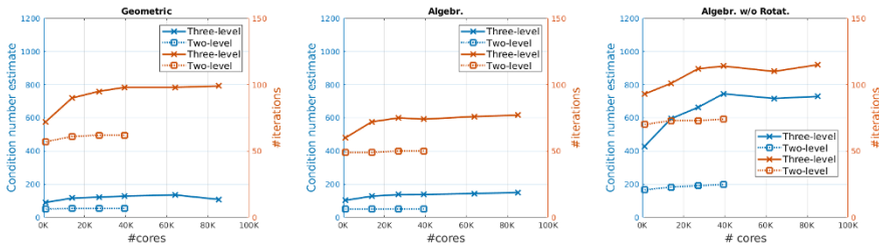
## 4 Weak parallel scalability results for the three-Level extension

In this section, we focus on weak parallel scalability results for the three-level GDSW preconditioner with a reduced dimensional coarse space. We always use *Option 1* to construct the coarse basis functions. In Trilinos the data is distributed among the processes via the map object. We use a repeatedly decomposed map to determine the interface  $\Gamma$ . This map can be passed as an input to the FROSch framework.

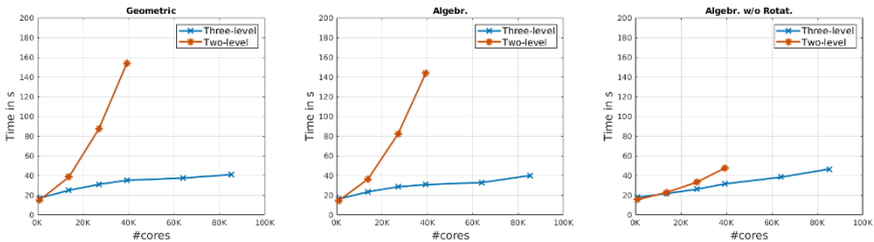
For our weak parallel scalability tests, we consider three different setups to determine the interface  $\Gamma$ , which result in different sizes and sparsity patterns for the coarse problem; see Figure 2. We either use the *Geometric Map*, which is constructed from the structured non-overlapping domain decomposition on the first level, or the *Algebraic Map* [6], which is built algebraically from the uniquely decomposed row map of the input matrix. In particular, the interfaces and hence the vertices, edges, and faces may differ slightly for the two different maps; this effect may be more pronounced for unstructured domain decompositions.



**Fig. 2:** Sparsity of the coarse matrix  $K_0$  for our linear elasticity model problem in three dimensions with 216 subdomains; using the *Geometric Map* (left) and the *Algebraic Map*, with rotations (middle) and without linearized rotations (right). The subdomain size is chosen such that each process of the uniquely decomposed map owns  $30^3$  nodes.



**Fig. 3:** Weak numerical scalability for the three- and two-level method with a reduced dimensional coarse space; see Table 1 for the data; using the *Geometric Map* and the *Algebraic Map* with and without rotations.



**Fig. 4:** Weak parallel scalability for the three- and two-level method with a reduced dimensional coarse space; see Table 1 for the data.

When using the *Algebraic Map*, we also consider the case where the rotations are neglected (*Algebr. w/o Rotat.*). In Figure 2, we only see minor differences in the sparsity pattern of  $K_0$  using the *Geometric Map* and the *Algebraic Map*. For higher numbers of subdomains, the differences between these two approaches will be more visible: for our largest test case with 85 184 subdomains, we have 539 460 as a maximum nonzero entries per core in  $K_0$  for the *Geometric Map*; this compares to 578 340 maximum nonzeros per core for the *Algebraic Map*. For all input maps, the

# Subd.	# Subr.	# Dofs	Map	Two-level			Three-level		
				$\kappa(M^{-1}K)$	Iter	Solver Time	$\kappa(M^{-1}K)$	iter	Solver Time
1 000	4	$2.4 \cdot 10^7$	Geom.	51.45	57	15.11s	90.46	72	16.99s
			Algebr.	50.73	49	14.54s	103.02	60	16.12s
			Algebr. w/o Rotat.	166.68	70	15.48s	429.05	93	17.91s
13 824	27	$3.3 \cdot 10^8$	Geom.	53.61	61	38.40s	116.19	90	24.89s
			Algebr.	51.08	49	36.39s	127.91	72	23.38s
			Algebr. w/o Rotat.	182.46	73	22.75s	594.97	101	21.58s
27 000	64	$6.5 \cdot 10^8$	Geom.	53.77	62	87.28s	122.18	95	30.87s
			Algebr.	51.12	50	82.01s	137.42	75	28.46s
			Algebr. w/o Rotat.	191.12	73	33.17s	663.44	112	26.02s
39 304	125	$9.4 \cdot 10^8$	Geom.	53.82	62	153.88s	128.39	98	35.12s
			Algebr.	51.12	50	144.01s	137.96	74	30.63s
			Algebr. w/o Rotat.	198.05	74	47.48s	745.26	114	31.40s
64 000	216	$1.5 \cdot 10^9$	Geom.	-	-	-	135.58	98	37.29s
			Algebr.	-	-	-	143.87	76	32.81s
			Algebr. w/o Rotat.	-	-	-	717.06	110	38.24s
85 184	275	$2.0 \cdot 10^9$	Geom.	-	-	-	108.49	99	40.80s
			Algebr.	-	-	-	150.37	77	39.87s
			Algebr. w/o Rotat.	-	-	-	729.14	115	46.45s

**Table 1:** Data corresponding to Figure 3 and 4. By *Iter*, we denote the number of PCG iterations, and  $\kappa$  is the condition number of the preconditioned operator. *Solver Time* is the time to build the preconditioner and to perform the Krylov iterations; see also Figure 3 and 4. The subdomain size is chosen such that each process of the uniquely decomposed map owns  $20^3$  nodes. We have  $H_c/H \approx 8$ . One layer of finite elements respectively one layer of subdomains is chosen as the overlap for each level.

two- and the three-level method are numerically scalable, whereas the *Algebraic Map without Rotations* yields the highest iteration counts and condition number estimates; cf. Figure 3 and Table 1. Replacing the direct solver for the coarse problem (used in the two-level method) by the application of the RGDSW preconditioner for the three-level method generally results in higher condition number estimates and iteration counts.

However, the three-level extension of the FROSch framework shows a better parallel weak scalability than the two-level method; cf. Figure 4 and Table 1. The *Solver Time* is the time to build the preconditioner and to perform the Krylov iterations. The time includes the factorization and forward backward substitution for the sparse direct solvers. For the three-level method the time for the unstructured decomposition of the coarse problem is also included. For all test settings, the three-level method is faster for 13 824 and more cores. Moreover, at 39 304 cores the three-level method is faster by more than a factor of four: Using the three-level method, we obtain a *Solver Time* of 35.12 s using the *Geometric Map* and 30.63 s using *Algebraic Map*. This compares to a *Solver Time* of 153.88 s for the *Geometric Map* and 144.01 s for the *Algebraic Map* in the two-level method. Using *Algebraic Map without Rotation* results in a smaller coarse problem, making the two-level methods more competitive. Here, the three-level method (*Solver Time* 31.40 s) is still faster by a factor of 1.5 than the two-level method (*Solver Time* 47.48 s). As the results are clear, we did not perform tests beyond the 39 304 cores for the two-level method. To illustrate the strong influence of the size of the coarse problem on the

preconditioner time, we consider the test case of 39 304 cores in Table 2. For this test case, the solution of coarse problem  $K_0$  in the *Geometric Map* setup takes 78% (120.19 s) of the total *Solver Time* (153.88 s).

# Subd.	# Subr.	# Dofs	Map	Two-level method		Three-level method	
				Size $K_0$	$K_0$ Solve Time	Size $K_{00}$	$K_{00}$ Solve Time
1 000	4	$2.4 \cdot 10^7$	Geom.	4 374	0.24s	6	<1e-5s
			Algebr.	4 374	0.22s	6	<1e-5s
			Algebr. w/o Rotat.	2 184	0.08s	3	<1e-5s
13 824	27	$3.3 \cdot 10^8$	Geom.	73 002	12.03s	366	0.01s
			Algebr.	73 002	12.04s	366	0.01s
			Algebr. w/o Rotat.	36 501	2.02s	174	0.003s
27 000	64	$6.5 \cdot 10^8$	Geom.	146 334	59.38s	1 056	0.08s
			Algebr.	146 334	45.75s	1 116	0.08s
			Algebr. w/o Rotat.	73 167	11.69s	546	0.04s
39 304	125	$9.4 \cdot 10^8$	Geom.	215 622	120.19s	2 508	0.29s
			Algebr.	215 622	114.06s	2 556	0.25s
			Algebr. w/o Rotat.	107 811	22.14s	1 290	0.11s
64 000	216	$1.5 \cdot 10^9$	Geom.	355 914	-	4 980	0.81s
			Algebr.	355 914	-	4 938	0.63s
			Algebr. w/o Rotat.	177 957	-	2 319	0.21s
85 184	275	$2.0 \cdot 10^9$	Geom.	477 042	-	6 432	0.63s
			Algebr.	477 042	-	6 660	0.72s
			Algebr. w/o Rotat.	238 521	-	3 222	0.16s

**Table 2:** Cost for solving the problem on the coarsest level. *Solve Coarse Problem Time* includes the time of the factorization of the problem as well as the forward and backward substitution in the Krylov iterations.

For this test case, the solution of coarse problem  $K_0$  in the *Geometric Map* setup takes 78% (120.19 s) of the total *Solver Time* (153.88 s). This time compares to less than a second (0.29 s) to solve the coarse problem corresponding to  $K_{00}$  in the three-level method. Similar results are obtained for the *Algebraic Map* where 114.06 s for the two-level method compare with 0.25 s for the three-level method. For the *Algebraic Map without Rotations* the size of the coarse problem is reduced by a factor of two; cf. Table 2. Therefore, the cost of the coarse problem reduces to 22.14 s for the two-level method, which compares to 0.11 s for the three-level method. Although the *Algebraic Map* has the largest coarse problem size (see Table 2) this is consistently the fastest setup of the three-level method. The stronger connectivity given by this coarse problem improves the iteration count and therefore decreases the *Solver Time*. Resulting in the highest number of iterations (cf. Table 1) the *Algebraic Map without rotations* is the slowest test case.

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